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WATTS' DICTIONARY OF CHEMISTRY

VOL. IV.

WATTS'

DICTIONARY OF CHEMISTRY

REVISED AND ENTIRELY REWRITTEN (1894)

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ASSISTED BY EMINENT CONTRIBUTORS

IN FOUR VOLUMES

VOL. IV.

WITH ADDENDA

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INTRODUCTION

TO THE ARTICLES ON INORGANIC CHEMISTRY.

IT has been thought advisable to include in an Addenda brief accounts of the chief work done in descriptive inorganic chemistry since the publication of Vols. I., II., and III., and the printing off the final proofs of Vol. IV.

Nothing bearing on organic chemistry has been included in the ADDENDA, as to give an account of what has been done in this department since the various volumes were published would occupy many hundred pages.

Dates are attached to the references made to original memoirs in the ADDENDA; and references are frequently made to abstracts of the memoirs in the Journal of the Chemical Society.

M. M. PATTISON MUIR.

INTRODUCTION

TO THE ARTICLES RELATING TO ORGANIC CHEMISTRY.

THE names used to denote ring formulæ are given below for convenience of reference.

Since the publication of the last volume I have been assisted in the work of reading and making abstracts of original memoirs by Drs. T. Cooksey, T. A. Lawson, Samuel Rideal, Messrs. J. Wilkie, G. N. Huntly, and J. T. Norman. I have also been assisted by Mr. Arthur G. Green and Mr. Cecil W. Cunnington in the work of revising the proof-sheets. I have great pleasure in thanking these gentlemen for the energetic and efficient manner in which they have carried out their share of the work.

H. FORSTER MORLEY.

Pentamethenyl hydride.

CH CH CH Tetramethenyl.

Nomenclature of Ring Formulæ.

Hydrocarbons.

$C_{e}H_{e} < CH_{2} > CH$ Indonaphthene.
compounds.
N≪N=CH>CH Pyridazine.
N≪CH.CH≫N Pyrasins.
N < CH:N CH Pyrimidine.
N <ch: n="">CH Triazoline.</ch:>
N≪N:N CH Osotetrasole.
$C_{\mathfrak{s}}H_{\mathfrak{s}} < \stackrel{CH:CH}{\underset{N=CH}{\leftarrow}} Quinoline.$
$C_6H_4 < \stackrel{CH:CH}{\underset{CH:N}{CH:N}}$ Isoquinoline.
$C_{\mathfrak{o}}H_{\mathfrak{o}} < N : CH $ Quinovaline.
$0_{e}H_{i} < \stackrel{CH:N}{\underset{N=CH}{CH}}$ Quinasoline.
$C_{\bullet}H_{\bullet} < CH:CH \\ N=N$ Cinnoline.
C.H. CH:N Phontriasine.
C.H. C(NH).CH:O.NPh C.H. Rosinduline
C ₆ H ₄ C _{O(NH).OH:O.NPh} C ₁₆ H ₆ { Naphthore induline.

N CH:CH CH Pyridine.

Oxygen compounds.

Sulphur compounds.

INITIALS OF SPECIAL CONTRIBUTORS.

- S. A. SVANTE ARRHENIUS, Ph.D., Professor of Chemistry in Stockholm. Contributes
 SOLUTIONS I.
- J. W. C. J. W. CAPSTICK, D.Sc., M.A., Fellow of Trinity College, and Demonstrator in Physics in the University, Cambridge. Contributes Capillarity, Methods Based on; and Viscosity of Liquids.
- G. G. GEORGE GLADSTONE, Esq. Contributes Optical methods, section Refraction and Dispersion.
- W. D. H. W. D. HALLIBURTON, M.D., F.R.S., Professor of Physiology at King's College, London. Contributes Proteïds.
- W. N. H. W. N. HARTLEY, F.R.S., Professor of Chemistry in the Royal College of Science Dublin. Contributes Optical methods, section Spectroscopic methods.
- E. A. L. | E. A. LETTS, Ph.D., Professor of Chemistry at Queen's College, Belfast. Contributes Phosphines.
- R. M. RAPHAEL MELDOLA, F.R.S., Professor of Chemistry in the Finsbury Technical College. Contributes Photographic Chemistry.
- W. O. WILHELM OSTWALD, Ph.D., Professor of Chemistry in the University of Leipzig.

 Contributes Electrical methods.
- C. O'S. | CORNELIUS O'SULLIVAN, F.R.S., Burton-on-Trent. Contributes STARCH and SUGARS.
- S. U. P. SPENCER PICKERING, M.A., F.R.S. Contributes Solutions II.
- S. R. SAMUEL RIDEAL, D.Sc. Contributes Tannin.
- W. A. T. WILLIAM A. TILDEN, D.Sc., F.R.S., Professor of Chemistry at the Royal College of Science, South Kensington. Contributes Terpenes.
- T. E. T. T. E. THORPE, D.Sc., Ph.D., F.R.S., Head of the Government Laboratories, Somerest House. Contributes Specific volumes.
 - Articles by Mr. MUIR are initialed M. M. P. M.
 - Unscened Astroles are by Dr. MORLEY, except those in the Addenda, which are by Mt. Mt. R.

ABBREVIATIONS

I. JOURNALS AND BOOKS.

When an author has been mentioned in an article, he is usually referred to thereafter in that article by his initial only.

```
Liebig's Annalen der Chemie.
A. A.
            Annales de la Sociedad Cientifica Argentina.
A. Ch.
             Annales de Chimie et de Physique.
P. Am. A.
            Proceedings of the American Academy of Arts and Sciences.
Am.
             American Chemical Journal.
Ann. M..
            Annales des Mines.
Am. S. .
A. C. J. .
             American Journal of Science.
            Journal of the American Chemical Society.
             American Chemist.
Am. Ch.
Am. J.
             American Journal of Pharmacy.
  Pharm.
             The Analyst.
An.
A. Ph. S.
            Proceedings of the American Philosophical Society.
Ar. N. .
             Archives néerlandaises-The Hague.
            Mémoires de l'Académie des Sciences.
Acad.
Ar. Ph. .
            Archiv der Pharmacie.
Ar. Sc. .
            Archives des Sciences phys. et nat.
В.,
            Berichte der deutschen chemischen Gesellschaft.
            Reports of the British Association.
B. A.
Bl. .
            Bulletin de la Société chimique de Paris.
        •
B. B.
            Berliner Akademie-Berichte.
B. C.
            Biedermann's Centralblatt für Agricultur-Chemie.
            Berzelius' Jahresberichte.
B. J. .
B. M.
            Berliner Monatsberichte.
C.S. Mem.
            Memoirs of the Chemical Society of London.
            Journal of the Chemical Society of London.
C. J. .
C.J. Proc.
            Proceedings of the Chemical Society of London.
C. N.
            Chemical News.
C. R.
            Comptes-rendus hebdomadaires des Séances de l'Académie des Sciences-
                 Paris.
c. c..
            Chemisches Central-Blatt.
D. P. J.
            Dingler's polytechnisches Journal.
Fresenius' Zeitschrift für analytische Chemie.
Fr. . .
G..
            Gazzetta chimica italiana.
G. A..
            Gilbert's Annalen der Physik und Chemie.
H. .
            Hoppe-Seyler's Zeitschrift für physiologische Chemie.
        ٠
T..
            Proceedings of the Royal Irish Academy.
     •
            Jahresbericht über die Fortschritte der Chemie und verwandter Theile
                 anderer Wissenschaften.
            Jahresbericht für Chemische Technologie.
J. C. T. .
J. M.
            Jahrbuch für Mineralogie.
J. de Ph.
J. Ph.
            Journal de Physique et des Sciences accessoires.
            Journal de Pharmacie et de Chimie.
J. pr.
J. Th.
            Journal für praktische Chemie.
Jahresbericht über Thierchemie.
        4
            Journal of the Russian Chemical Society.
Jenaische Zeitschrift für Medicin und Naturwissenschaft.
J. R. .
J. Z. .
L. V.
            Landwirthschaftliche Versuchs-Stationen.
            Monatshefte für Chemie und verwandte Theile anderer Wissenschaften.
М.
M.S.
            Le Moniteur Scientifique.
Mém. S.
            Mémoires de la Société d'Arcueil.
  \mathcal{X}A.
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Mémoires couronnés par l'Académie de Bruxelles.

Mém. B.

Vol. IV.

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Nature.
N.Ed.P.J.
            New Edinburgh Philosophical Journal.
N. J. P.
            Neuer Jahresbericht der Pharmacie.
N. R. P.
            Neues Repertorium für die Pharmacie.
N. J. T. .
            Neues Journal von Trommsdorff.
N. Z. R.
P. M.
            Neue Zeitschrift für Rübenzuckerindustrie.
            Philosophical Magazine.
P. .
P. B. .
            Poggendorff's Annalen der Physik und Chemie.
            Beiblätter zu den Annalen der Physik und Chemie.
Pf.
Pr. E.
            Pflüger's Archiv für Physiologie.
            Proceedings of the Royal Society of Edinburgh.
Ph.
            Pharmaceutical Journal and Transactions.
Ph. C.
Ph. C.
            Pharmaceutisches Central-Blatt.
        ٠
Pr. P. R. L.
            Proceedings of the Boyal Society.
            Proceedings of the Royal Institution of Great Britain.
P. Z.
            Pharmaceutische Zeitschrift für Russland.
R. T. C..
            Recueil des travaux chimiques des Pays-Bas.
R. P.
Q. J. S. .
            Repertorium für die Pharmacie.
            Quarterly Journal of Science.
Š. .
            Schweigger's Journal der Physik.
Scher. J.
            Scherer's Journal der Chemio.
8. C. I. .
            Journal of the Society of Chemical Industry.
Sits.W. .
            Sitzungsberichte der K. Akademie zu Wien.
T. or Tr.
T. E. . .
            Transactions of the Royal Society.
            Transactions of the Royal Society of Edinburgh.
W.
            Wiedemann's Annalen der Physik und Chemie.
w. J.
            Wagner's Jahresbericht.
z.
            Zeitschrift für Chemie.
Zeit.ang.
            Zeitschrift für angewandte Chemie.
  Oh.
Z. B.
            Zeitschrift für Biologie.
Z. f. d. g.
Natur-
            Zeitschrift für die gesammten Naturwissenschaften.
  wiss.
Z. K..
            Zeitschrift für Krystallographie und Mineralogie.
Z. P. C.
Z. V.
            Zeitschrift für physikalische Chemie.
Zeitschrift des Vereins für die Rübenzuckerindustrie des deutschen
                Reiches.
Bn.
            Handbuch der organischen Chemie: von F. Beilstein, 2te Auflage.
Bn. .
E. P.
            English Patent.
G. P.
            German Patent.
Gm.
            Gmelin's Handbook of Chemistry—English Edition.
            Gmelin-Kraut: Handbuch der anorganischen Chemie.
Gm.-K.
            Traité de Chimie organique: par Charles Gerhardt.
Gerh.
K..
            Lehrbuch der organischen Chemie: von Aug. Kekulé.
G. O.
            Graham-Otto: Lehrbuch der anorganischen Chemie [5th Ed.]
Stas,
            Stas' Recherches, &c.
  Rech.
                                              Aronstein's German translation is re-
Stas,
            Stas' Nouvelles Recherches, &c.
                                                   ferred to as Chem. Proport.
 Nouv. R.
Th. . . Thomsen's Thermochemische Untersuchungen.
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II. TERMS AND QUANTITIES, &c., FREQUENTLY USED.

```
Water; e.g. NaOHAq means an aqueous solution of caustic soda.
                         18 parts by weight of water.
                        Residues of mono-, di-, and tri-basic acids. Thus, in describing the salts of a monobasic acid NaA', CaA'<sub>2</sub>, AlA'<sub>3</sub> may be written, HA' standing for the acid. For a dibasic acid we should write Na<sub>2</sub>A'', CaA'', Al<sub>4</sub>A''<sub>3</sub> &c. Stand for bases of the ammonia type, in describing their salts. Thus the hydrochloride would be B'HCl or B''2HCl, according as the base is
A"
Δ′′′
B' B" &c.
                                  monacid or discid, &c.
                         Concentrated.
sone.
ML.
                        Dilute.
                        gram.
                        milligram.
mgm.
mm. .
                        millimetre.
mol. .
                        molecule
```

```
oil.
            liquid, nearly, or quite, insoluble in water.
            precipitate.
pp.
to ppt.
            to precipitate.
            precipitating.
ppg. .
            precipitated.
ppd. .
sol.
             soluble in.
insol..
            insoluble in.
w. e. sol. .
            very easily
v. sol.
             Very
                             soluble in.
m. sol.
             moderately
sl. sol.
             slightly
v. sl. sol.
             very slightly
v. .
             see.
cf..
             compare.
about.
(°)
             a melting-point.
             a boiling-point.
н.:
             Hardness (of minerals).
At. w.
             Atomic weight.
Mol. w. or
             Molecular weight.
  M. w.
D.. .
             Density.
cor.
             corrected.
uncor.
             uncorrected.
i.V. .
             in vapour.
v.D. .
             vapour-density, i.e. density of a gas compared with hydrogen or air.
S.G. .
             Specific gravity compared with water.
S.G. &
                              at 10^{\circ} compared with water at 0^{\circ}.
S.G. 13
S.G. 13
                        ,,
                 ,,
                               ,, 12°; compared with water of which the temperature is
                not given.
S.H. .
             Specific heat.
S.H.v.
                            of a gas at constant volume.
S.H.p.
                                                  pressure.
             Quantity of heat, in gram-units, produced during the complete combustion of the mass of a solid or liquid body represented by its
H.C. .
                  formula, taken in grams.
H.C.v.
             Heat of combustion in gram-units of a gram-molecule of an element or
                  compound, when gaseous, under constant volume.
             The same, under constant pressure.
H.C.p.
             Quantity of heat, in gram-units, produced during the formation of the
H.F.
                  mass of a solid or liquid body represented by its formula, taken in
                  grams, from the masses of its constituent elements expressed by
                  their formulæ, taken in grams.
H.F.v.
             Heat of formation of a gram-molecule of a gaseous compound from the
                  gram-molecules of its elements under constant volume.
H.F.p.
             The same, under constant pressure.
H.V.
             Heat of vaporisation of a liquid, i.e. gram-units of heat required to change
                  a gram-molecule of the liquid compound at B.P. into gas at same
                  temperature and pressure.
T.C. .
             Thermal conductivity (unit to be stated).
S.V. .
             Specific volume; or the molecular weight of a gaseous compound divided
by the S.G. of the liquid compound at its boiling-point compared with
                  water at 4°.
S.V.S.
             Specific volume of a solid; or the mass of the solid expressed by its
                  formula, taken in grams, divided by its S.G.
E.C. . . . C.E. (10°
             Electrical conductivity (the unit is stated in each case).
             Coefficient of expansion (between 10° and 20°).
  to 20°)
                                     of a gas - volume dissolved by 1 volume of water.
                                    of a liquid or solid = number of grms. dissolved by
S. (alco-
             Solubility in water
                        " alcohol
                                    100 grms. of water. In both cases the temperature
  hol)
                                     is stated.
             Index of refraction for hydrogen line \beta.
\mu_{B} .
μ<sub>D</sub>, &c.
                                     sodium
                                                     D, &c.
            Molecular refraction for sodium light, i.e. index of refraction for line
R. V.
                  minus one, multiplied by molecular weight, and divided by S.G. at 15"
                  compared with water at 0°.
             The same; S.G. being determined at 15°-20° and referred to water at 4°. The same for line of infinite wave-length, index being determined by
                  Cauchy's formula (Brühl's B.).
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| Specific rotation for sodium light.
[a]_{D} .
                                                         [\alpha] = \frac{100}{n} \times \frac{\alpha}{d}.
                                   " neutral tint.
                                                                          a - observed rotation for
[a]<sub>1</sub>
                    100 mm. of liquid. d = S.G. of liquid. p = no. of grammes of active
                    substance in 100 grammes of liquid.
M. M
              Molecular magnetic rotatory power = \frac{m \times w}{d \times \alpha' \times m'}, where m = molecular
                    weight of the body of S.G. = d, a = angle of rotation under magnetic
                    influence, a' = angle of rotation of water under same influence, and
                    m' = molecular weight of water (18).
               Acetyl C<sub>2</sub>H<sub>3</sub>O.
Ac
              Benzoyl C.H.O.
Cyanogen CN.
\mathbf{Bz}
         .
Cy
Εt
              Ethyl C2H5.
Мe
              Methyl CH,.
Ph
              Phenyl C.H.
                                                       in formulæ.
              Normal Propyl CH<sub>2</sub>. CH<sub>2</sub>. CH<sub>3</sub>. Isopropyl CH(CH<sub>3</sub>)<sub>2</sub>.
\mathbf{Pr}
Pr
R, R' &c.
              Alcohol radicles or alkyls.
prim.
              primary.
8ec
              secondary.
              tertiary.
tert
n.
              normal.
m, o, p
              meta-ortho-para.
C
              consecutive.
  .
      .
          ٠
              irregular.
              symmetrical.
8
u
              unsymmetrical.
  .
      ٠
ψ
              pseudo.
              attached to nitrogen.
ν
              Employed to denote that the substituent is attached to a carbon atom
                    which is next, next but one, or next but two, respectively, to the terminal carbon atom. The end to be reckoned from is determined by the nature of the compound. Thus CH_cHBr.CO<sub>2</sub>H is a-bromo-
α
ß
   .
                    propionic acid.
               denotes that the element or radicle which follows it is attached to a ter-
                    minal carbon atom.
               indicate position in an open chain, only.
\alpha, \beta, \gamma, &c.
              indicate position in a ring only.
1,2,3,&c.
               Used when \alpha, \beta, &c. are employed in a sense different from the above,
(\alpha), (\beta),
              e.g. (a)-di-bromo-camphor.
Baeyer's Nomenclature:
   &o.
 (B.)
                    benzene ring.
                    pyridine ring
(Py.) .
                        Thus (B.1:3) dichloroquinoline, means a meta-dichloroquinoline in
                    which the chlorine atoms are both in the benzene ring.
                        While (Py. 1:3) dichloroquinoline, means a similar body, only the
                    chlorine atoms are in the pyridine ring. The numbers are counted from two carbon atoms which are in different rings, but both united
                    to the same carbon atom.
               denotes the central ring in the molecule of anthracene, acridines, and
(A.) .
                    azines.
              means that the element or radicle it precedes is in a closed ring.
680-
         .
                                                                         not in a benzene ring.
exo-
               denotes isomerism that is not indicated by ordinary formulæ; thus maleic
allo- .
                    acid may be called allo-fumaric acid.
              denotes displacement of oxygen by sulphur.
,, the group SO,H, except in the word sulphocyanide.
thio- .
sulpho- .
                        the group SH.
sulphydro-
               Tribromonitrobenzene sulphonic acid [1:2:3:4:5] means that the three
                    bromines occupy positions 1, 2, and 3; the nitro- group the position 4,
                    and the sulpho-group the position 5.
    Denotes that the formula to which it is affixed has not been determined by
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analysis. But it by no means follows that formulæ without this mark are those of analysed compounds.

All ten peratures are given in degrees Centigrade unless when specially stated otherwise.

Wave-lengths are given in 10-7 mm.

Formulæ, when used instead of names of substances; have a qualitative meaning Thomsen's notation is used in thermochemical data.

DICTIONARY OF CHEMISTRY.

PHENYL - AZOXAZOLE CH: N O. [30°].

Formed by the action of CO2 on an alkaline solution of the oxim of phenyl-glyoxal (Russano, B. 24, 3503). Needles, insol. water.

PHENYL - BENZAMIDINE C13H12N C₆H₅C(NH).NHPh or C₆H₅.C(NPh).NH₂ [112°]. Formed from benzonitrile and aniline hydrochloride at 230° (Bernthsen, A. 184, 348; 192, 33). Formed also from benzimido-ether and aniline, and from CPhCl:NPh and NH, (Kobbert, A. 265, 138). Nodules (from alcohol), sl. sol. water. Forms a crystalline nitrate.

Reactions.—1. Split up by heat into aniline and benzonitrile.—2. Dry H₂S at 130° forms C₄H₂.CS.NHPh and some thio-benzamide.— 3. CS, at 100° forms thio-benzanilide and phenylbenzamidine sulphocyanide.-4. Reduced by sodium-amalgam to C₆H₃.CH(NH₂).NHPh. — 5. Nitrous acid forms benzanilide (Mieran, A. 265, 141).

The derivatives C.H.C(N.SO.Ph).NHPh [139°] and C,H,(SO,N:C(NHPh).C,H,), [196°] have been prepared (Wallach, A. 214, 214; Jackson, Am. 9, 346).

s - Di - phenyl - benzamidine C19H16N2 i.e.

C₄H₅.C(NPh).NHPh. [144°].

Formation.- 1. From be zanilide by treatment with PCl_s and aniline (Gerhardt, A. 108, 219; Hofmann, Z. 1866, 161; Wallach, A. 184, 83).—2. From Ph.CCl_s and aniline with or without ZnCl2 (Limpricht, A. 135, 82; Döbner, B. 15, 233).—3. A product of the action of aniline hydrochloride at 250° on benzonitrile, thiobenzamide, or phenyl-benzamidine (Bernthsen, A. 184, 352).-4. By heating benzanilide ith phenyl cyanate at 190° (Kühn, B. 18, 1476).-5. By heating the hydrochloride of benzimido-ether in alcohol with aniline for a

long time (Kobbert, A. 265, 155). Properties .- Needles (from alcohol), m. sol. benzene. Its alcoholic solution is neutral to litmus. Not affected by nitrous acid. Split up into aniline and benzanilide when boiled for a

long time with alcohol.

Reactions.-1. Dry H2S at 160° forms thiobenzanilide. - 2. CS, at 140° forms PhCS.NHPh and NPhCS (Bernthsen, A. 192, 84).-3. Conc. HClAq at 150° yields aniline and benzoic acid.

Salts.-B'HCl, v. sl. sol. water.-B', H2PtCl,

-B'C_cH₂N₃O₇: yellow needles. u-Di-phenyl-benzamidine

C₆H_a.C(NH).NPh₂. [112°]. Formed by heating diphenylamine hydrochloride with b nzonitrile at 180° for 5 days (Bernthsen, A. 192, 5). Trimetric tablets, v. sol. alcohol and benzene. Voz. IV.

Its alcoholic solution turns litmus blue. Conc. H₂SO, gives a violet-blue colour on warming.

Reactions .- 1. Dilute HClAq at 180° forms NPh2Bz.-2. The free base and its hydrochloride yield benzonitrile and diphenylamine on heating.—3. Nitrous acid forms NPh,Bz (Klobbert, A. 265, 157).—4. H₂S at 130° forms Ph.CS.NH2, diphenylamine, PhCS.NPh2, and NH3.-5. CS2 at 140° forms PhCS.NPh, and HNCS.

Salts. — B'HCl. [c. 223°]. crystals, a:b:c = .529:1:.507; $\beta = .85°52'$. V. sol. water, sol. alcohol, insol. ether.—B'2H2PtCl8.— B'HNCS. [203°]. Yellow prism PHENYL-BENZAMIDOXIM Yellow prisms.

C_eH_s.C(NOH).NHPh. [136°]. Formed by boiling C.H. CS.NHPh with an alcoholic solution of hydroxylamine (Müller, B. 19, 1669). Needles (from water). Yields B'HCl. ClCO₂Et forms C₁₄H₁₆N₂O₂ [167°].

DI-PHENYL-BENZAZIDINE C₁₉H₁₉N₄ i.e.

C₆H₅C(N.NHPh).NH.NHPh. [179°]. Formed by the action of phenyl-hydrazine on PhCCl, or on the hydrochloride of benz-imido-ether dissolved in absolute alcohol (Pinner, B. 17, 182; Marckwald, C. C. 1888, 1410). Dark-red needles.

PHENYL-BENZENE v. DIPHENYL.

p-Di-phenyl-bensene C₁₈H₁₄ i.e. C_eH₄Ph₂.

Mol. w. 230. [207°]. (c. 405°).

Formation.—1. A product of the action of sodium on an ethereal solution of p-di-bromobenzene, or, better, of a mixture of p-di-bromo-benzene with bromine (Riese, Z. [2] 6, 192, 735; A. 164, 168).—2. Together with its isomeride, and other products, by passing the vapour of benzene, or of a mixture of benzene and toluene, through a red-hot tube (G. Schultz, B. 6, 415; A. 174, 230; 203, 118; Carnelley, C. J. 37, 712).—3. Together with the isomeride and other products, by the action of AlCl, on a mixture of MeCl and diphenyl (Adam, Bl. [2] 49, 97; A. Ch. [6] 15, 241).

Properties.-Needles (from benzene), or iridescent laminæ (by sublimation); insol. alcohol, v. sl. sol. HOAc. Its solution in benzene shows blue fluorescence. Conc. H₂SO₄ gives a dirtygreen solution changing to purple. CrO, in HOAc oxidises it to diphenyl p-carboxylic [216°] and terephthalic acids. Does not form a compound with picric acid. Not volatile with steam. Yields C₁₈Cl₁₄ on exhaustive chlorination with SbCl₅ (Merz a. Weith, B. 16, 2884).

References. - Brono - and Tri-chloro - Di-

PHENYL-BENZENES.

Iso-di-phenyl-benzene C₁₈H₁₄. [85°]. (c. 370°). Formed as above. Slender prisms, v. sol. alcohol.

CrO₃.

s-Tri-phenyl-benzene C₂₁H₁₈ i.e. C₂H₃Ph₃.

Formed from Mol. w. 306. [171°]. (above 360°). Formed from acetophenone by treatment with P2O3, with alcoholic NH₃, or with dry HCl (Engler, B. 6, 638; 7, 1123). Formed also by heating C₆H₃(C₆H₄.CO₂H)₃ with KOH (Gabriel a. Michael, B. 11, 1007). It is also a by-product in the action of acetophenone on dimethylaniline in presence of ZnCl₂ (Döbner a. Petschoff, A. 242, 336). Trimetric crystals; a:b:c = .566:1:.766, sl. sol. alcohol. Yields benzoic acid on oxidation by CrO, in HOAc (Mellin, B. 23, 2533). By exhaustive chlorination with SbCl, it yields per-chloro-triphenyl-benzene (Merz a. Weith, B. 16, 2883). On heating with I and red P at 275° it yields the crystalline dodeca-hydride $C_{24}H_{39}$ and finally oily $C_{24}H_{38}$ (Mellin, B. 23, 2534). Fuming H_2SO_4 at $C_{24}H_{38}$ (Mellin, B. 23, 2534). Furning H_2SO_4 at 100° forms a disulphonic acid, which yields the Ba salt C, H, S, O, Ba crystallising in needles.

PHENYL-BENZENE-SULPHAZIDE Phenyl-hydrazide of BENZENE SULPHONIC ACID. DĬ-PHENYL-BENZENYL-DI-UREA

NHPh.CO.NH.CPh:N.CO.NHPh. Formed from benzamidine hydrochloride, phenyl cyanate, and NaOHAq (Pinner, B. 22, 1607). Needles, insol. water, v. sl. sol. hot alcohol.

PHÉNYL-BENZIMIDO-ETHER C15H15NO i.e. CPh(NPh).OEt. A product of the action of aniline on benzimido-ether (Lossen, A. 265, 138). Oil. Decomposed by conc. HClAq into EtCl and benzanilide.

PHENYL BENZOATE v. Benzoyl derivative of PHENOL

PHENYL-BENZOIC ACID v. DIPHENYL-CARB-OXYLIC ACID.

PHENYL-BENZOPHENONE C19H14O i.e. [1:4] C₆H₅.C₈H₄.CO.C₆H₅. [104°]. V.D.9.0. Formed by oxidation of p-benzyl-diphenyl with chromic acid (Goldschmiedt, M. 2, 437; Köller, M. 12, Formed also from diphenyl, BzCl, and Yields benzophenone-p-carboxylic acid on further oxidation by CrO...
Oxim. [194°].

Needles. When HCl is passed into its solution in HOAc containing Ac₂O and the mixture heated in sealed tubes at 100° , it is converted into the isomeric C_6H_5 . C_6H_4 . CO.NHC₆H₅ [224°]. The oxim is reduced by sodium-amalgam to the corresponding amine C_6H_5 . C_6H_4 . $CH(NH_2)$. C_6H_5 [77°].

Benzoyl derivative of the Oxim [193°]. Needles (from alcohol).

Phenyl-hydrazide. [144°]. Yellow needles (from dilute alcohol).

Di - phenyl - benzophenone $(C_6H_5.C_8H_4)_2CO.$ [229°]. Formed by the action of COCL on diphenyl in presence of AlCl, (Adam, Bl. [2] 47, 688; A. Ch. [6] 15, 259). Got also by oxidising $\mathrm{CH}_2(\mathrm{C_4H_4Ph})_2(\mathrm{Weiler}, B.7, 1188)$. White needles, sol. acetone, v. sl. sol. alcohol. Not attacked by fuming HNO, or by H2SO, and HNO, at 100°. By adding Na to its solution in alcohol-benzene it is reduced to (C,H,C,H,)2CH.OH, [151°], S. (ether) 5; S. (alcohol) 1.25 at 15°. Potash-fusion gives diphenyl-p-carboxylic acid [218°].
PHENYL-BENZOPHENONE CARBOXYLIC

ACID C₈H₄Ph.CO.C₆H₄.CO₂H. [220°]. Formed from diphenyl, phthalic anhydride, and AlCl. (Kaiser. A. 257. 96). Needles. Yields an oxim

Yields benzoic acid on oxidation by [140°], a phenyl-hydrazide [194°], and a methyl ether MeA' [85°-90°]

PHENYL-BENZOYL is BENZOPHENONE.

PHENYL-BENZOYL-ACETIC AC'D

CHBzPh.CO₂H. Methyl ether MeA'. Formed from de-oxybenzoïn, NaOEt, and ClCO, Me (Rattner, B. 21, 1316). Oil, decomposed by distillation into C2H2Ph2, CO2, and HOBz.

PHENYL-BENZOYL-BUTYRIC ACID

CHBzPh.CH₂.CH₂.CO₂H. (β)-Desyl-propionic acid. [136°]. Formed from deoxybenzoin, NaOEt, and \$\beta\$-iodo-propionic ether (Knoevenagel, \$B. 21, 1344). Needles. Yields MeA' [64°] and EtA' [34°].

The isomeric acid CHBzPh.CHMe.CO.H [215°], formed from β -bromo-propionic acid, is less sol. alcohol and ether.

PHENYL-BENZOYL-CARBINOL v. BENZOÏN. PHENYL-BENZOYL-PROPIONIC ACID

CHBzPh.CH₂.CO.H. [156°]. Formed from de-oxybenzoïn, NaOEt, and chloro-acetic ether (Meyer a. Oelkers, B. 21, 1295; Knoevenagel, B. 21, 1344). Small tables (from alcohol).

Di-phenyl-benzoyl-propionic acid CH₂Bz.CPh₂CO₂H. [183°]. Got by the action of alcoholic potash on the lactone of oxy-tri-phenylcrotonic acid (Japp a. Klingemann, B. 22, 2882). Yields a phenyl-hydrazide $C_{28}H_{22}N_2O$ [185°], and an oxim C22H27NO2 [152°

PHENYL-BENZYL ALCOHOL C₁₃H₁₂O i.c. C₈H₄Ph.CH₂OH. Got from C₈H₄Ph.CH₃ by treatment with Br and alcoholic potash; the resulting syrupy C.H.Ph.CH2OEt being treated with HI (Adam, Bl. [2] 49, 97). Syrup. PHENYL-BENZYL-AMIDO-DI-PHENYL-

METHANE C₂₈H₂₃N *i.e.* CH₂Ph.C₆H₄NPh.C₇H₇. Formed by heating NPh₂H (1 mol.) with benzyl chloride (2 mols.) and ZnCl₂ at 210° (Meldola, C. J. 41, 200). Solid; v. sol. ether, insol. alcohol and HOAc.

PHENYL-BENZYL-AMINE v. BENZYL-ANIL-

Nitrosamine C,H,NPh.NO. [58°]. Needles (Antrick, A. 227, 360). Yields benzyl-aniline and benzylidene-aniline on treatment with alcoholic HCl (O. Fischer, A. 241, 828).

Phenyl-di-benzyl-amine v. DI-BENZYL-ANIL

Di-phenyl-benzyl-amine NPh2.CH2Ph. [87°]. Got from C₈H₅.CSNPh₂, zinc-dust, and HClAq (Bernthsen a. Trompetter, B. 11, 1761; cf. Wilm a. Girard, B. 8, 1196). Needles, sl. sol. can alcohol. Yields a green dye on heating with HClAq and arsenic acid (Meldola, B. 14, 1385).

DI-PHENYL-DI-BENZYL-TETRAZONE NPh(CH₂Ph).N:N.NPh(CH₂Ph). [109°]. Formed from (a)-phenyl-benzyl-hydrazine, CHCl₃, and HgO (Michaelis a. Philips, A. 252, 290). Needles.

PHENYL-BENZYL-CARBINOL C, H14O i.e. CH_Ph.CH(OH).C.H.. Toluylene hydrate. [42°]. Formed by reducing deoxybenzoïn, and by the action of alcoholic potash or the same body (Limpricht a. Schwanert, A. 155, 62; Goldenberg, A. 174, 332; Zagoumenny, A. 184, 163; Anschütz, A. 261, 298). Formed also [62°] by the action of nitrous acid on CH2Ph.CHPh.NH2 (Leuckart, B. 22, 1410). Long slender needles from alcohol), insol. water. Reduced by HIAq o dibenzy. Yields a liquid acetyl derivative. o dibenzyı.

PHENYL BENZYL ETHER v. Phenyl ether

of BENZYL ATGOROL.

PHENYL-BENZYL-ETHYL-THIO-UREA C₁₈H₁₈N₂S. [91°]. Formed from ethyl-thiocarbimide and benzyl-aniline in alcohol (Dixon, C. J. 59, 566). Rectangular prisms, insol. cold

water, v. e. sol. boiling alcohol.

An isomeride [91° uncor.] formed from benzyl-thiocarbimide and ethyl-aniline crystallises in oblique prisms grouped in rosettes.

lises in oblique prisms grouped in rosettes.

PHENYL-BENZYL-HYDRAZINE C₁₃H₁₄N₂
i.e. CH₂Ph.NPh.NH₂. [26°]. Formed from sodium phenyl-hydrazine and benzyl chloride, and got also by reduction of phenyl-benzyl-nitrosamine (Antrick, A. 227, 361; Michaelis a. Philips, A. 252, 286). Needles (containing aq), decomposed by heat. Benzoic aldehyde forms CH₂Ph.NPh.N:CHPh. [111°]. In benzene solution it gives with SOCl₂ the thionyl compound C₂H₂NPh.N:SO [65°] (Michaelis a. Ruhl, A. 270, 12Δ). C₃H₃PCl₂ forms C₂H₃NPh.N:PC₄H₃ [141°], crystallising from ether in needles.—B'HCl. [167°]: colourless needles. Benzene phosphinate B'C₃H₃PH₂O₂ [108°] (Michaelis, A. 270, 135).

Acetyl derivative. [121°]. Scales. Bensylo-chloride (C,H,)2NPhCl.NH2.

[154°].

PHENYL-BENZYLIDENE-ALLYL-HYDRA-ZINE NPh(C₃H₃).N:CHPh. [52°]. Formed from phenyl-allyl-hydrazine and benzoic aldehyde (Michaelis a. Claesson, B. 22, 2237). Needles, v. sol. ether and hot alcohol.

PHENYL-BENZYLIDENE-AMINE v. BENZ-

YLIDENE-ANILINE.

Phenyl-benzylidene-diamine

 C_8H_8 .CH(NH₂)(NHPh). [115°]. Formed by reduction of phenyl-benzamidine with zinc and HCl (Bernthsen a. Szymanski, B. 13, 917). Small crystals.—B'HCl. [224°]. Thick prisms.—B'₂H₂PtCl₆: spikes or plates.

DI-PHENYL-BENZYLIDENE - ETHYLENE.

DIAMINE CHPh NPh C₂H₄. [137°]. Formed

from benzoic aldehyde and di-phenyl-ethylene-

diamine (Moos, B. 20, 732). Needles, split up

by dilute HCl into the parent bodies.

PHENYL - BENZYLIDENE - ETHYL-HYDRAZINE PhNEt.N:CHPh. [59°]. Formed from benzoic aldehyde and phenyl-ethylhydrazine (Michaelis a. Philips, A. 252, 272).

PHENYL-BENZYLIDENÉ-HYDRAZINÉ v. Phenyl-hydrazide of Benzoic Aldehyde.

DI-PHENYL-BENZYLIDENE-DI-INDOLE CHPh($C_{14}H_{10}N$)₂. [263° uncor.]. Formed from benzoic aldehyde and $C_{6}H_{4} < \begin{array}{c} CH \\ NH \end{array} > CPh$ (E. Fischer a. Schmidt, B. 21, 1074). Slender leaflets, v. sl. sol. hot alcohol.

PHENYL-BENZYL-INDOLE C₂₁H₁₇N i.e. C₃H₄ C(C₇H₁₇) CPh. [101°]. Formed by heating the phenyl-hydrazide of di-benzyl ketone with alcoholic HCl (Trenkler, A. 248, 112). Hexagonal prisms (from hot ligroïn). Does not give the pine, wood reaction.

Does not give the pine-wood reaction.

PHENYL BENZYL KETONE C₁₄H₁₂O i.e.

C₆H₃.CH₂.CO.C₆H₅. Desoxybenzoin. Deoxybenzoin. Mol. w. 196. [60°]. (322° cor.). (177.6°

at 12 mm.) (Anschütz a. Berns, B. °0, 1392).

Formation.—1. By the action of zinc and HClAq on benzoin (Zinin, A. 119, 179; 126, 218; 149, 375; Z. [2] 4, 718; V. Meyer,

B. 21, 1296; Wachter, B. 25, 1728).—2. By passing benzoïn over heated zinc-dust (Limpricht a. Schwanert, A. 155, 59).—3. By heating bromo-di-phenyl-ethylene with water at 190° (L. a. S.).—4. By distilling a mixture of calcium benzoate and calcium phenyl-acetate (Radziszewski, B. 6, 489; 8, 756).—5. By the action of AlCl, on a mixture of phenyl-acetyl chloride and benzene (Graebe a. Bungener, B. 12, 1079).—6. From phenyl-acetic acid, benzene, and P₂O₅ (Zincke, B. 9, 1771).—7. By the action of alcoholic ammonium sulphide (or KHS) on benzil (Zinin, J. pr. 33, 35; Jena, A. 155, 87).—8. By dissolving s-di-phenyl-acetylene in H₂SO₄ and adding water (Béhal, Bl. [2] 49, 337).

Properties.—White plates (from alcohol); sl. sol. hot water, volatile with steam.

Reactions .- 1. On heating with alcoholic potash it yields phenyl-benzyl-carbinol and 'diethyl carbobenzonic 'acid $C_{18}H_{18}O_2$ [100°], which gives EtA' (209° at 11 mm.) (Zagoumenny, A. 184, 163; Anschütz, A. 261, 298). The acid $C_{18}H_{18}O_2$ is converted by P and HI into an isomeric is converted by F and III med an assume a cid [134°], by HNO, (S.G. 1·18) into $C_{18}H_{10}O_3$ [120°], by conc. HNO₃ into $C_{18}H_{16}(N_{2})_{2}O_{3}$ [156°], and by potash-fusion into β -benzylisobutyric acid.—2. KOH in propyl alcohol at 150° forms $C_{20}H_{22}O_2$ [90°] and also an isomeride [139°] which yields $C_{20}H_{20}(NO_2)_2O_2$ [176°].—3. KOH in isobutyl alcohol at 140° forms diisobutyl-carbobenzonic 'acid C22H28O2 [148°], S. (alcohol) 5. The homologous isoamyl compound $C_{24}H_{30}O_{2}[160]$ may be got in like manner.—4. Reduced by HI to di-phenyl-ethylene and di-phenylethane .- 5. Sodium-amalgam reduces it to C28H26O2 and finally to phenyl-benzyl-carbinot .--6. Nitric acid (S.G. 1.2) yields benzil, nitro-benzil, and p-nitro-benzoic acid. Conc. HNO, (S.G. 1.475) at 0° forms $C_{14}H_{11}$ (NO₂)O [142°], whence $C_{14}H_{11}$ (NH₂)O [95°]. The nitro- and amidocompounds yield oxims melting at 107° and 141° respectively. Fuming HNO₃ (S.G. 1.51) at 0° forms three isomeric di-nitro- derivatives [116°], [126°] and [155°] (Golubeff, B. 13, 2403; J. R. 13, 23).—7. Bromine forms C₁₄H₁₁BrO [55°] (Knoevenagel, B. 21, 1355) and C₂H₂.CO.CBr₂.C₂H₃ [112°].—8. PCl₃ yields CHPh.COCBP.—9. Alceholic motion and baryonic CHPh:CClPh.—9. Alcoholic potash and benzoic aldehyde form benzamarone $C_{70}H_{86}O_{4}$ [215°] (Japp a. Klingemann, B. 21, 2934; cf. Zinin, Z. 1871, 127) .- 10. NaOEt and nitrous acid yield the mono- oxim of benzil.—11. NaOEt and CSCl₂ yield golden (C₆H₃.C(CS).CO.C₆H₃), [286°] (Bergreen, B. 21, 350).—12. NaOEt and phenyl-ethyl MeI vield phenyl C₆H₅.CO.CHMePh. Other alkyl iodides act in like manner (V. Meyer, B. 21, 1295). -13. Sodium acting on a benzene solution in absence of air forms C_sH_s.CO.CHNa.C_sH_s, a very hygroscopic yellow substance converted by CO₄ into an acid, whence hydroxylamine forms CHPh.CO O [160°] (Beckmann a. Paul, 4. 266, 22).—14. CSCl₂ or CS₂ forms desaurin C₂₀H₂₀N₂O₂ which is fiery yellow, sparingly soluble, and jields a violet solution in \hat{H}_2SO_4 (Bergreen a. Meyer, B. 21, 353; Meyer a. Wege, B. 24, 3535; Wachter, B. 25, 1727). HNO, and H.SO, convert desaurin into C₁,H.SN₂O₂? [60°]. Furning HNO, forms C₁,H.SN₂O₁₀ [60°] and m-nitro-benzoic acid. Desaurin when heated for a long time with

B 2

aniline yields phenyl benzyl ketone, tri-phenylguanidine, and H2S

Oxim C₆H₅.CH₂.C(NOH).C₆H₅. [98°].

Phenyl-hydrazide

CH₂Ph.C(N₂HPh).Ph. [106°]. Needles (from

alcohol) (Ney, B. 21, 2447).

PHENYL BENZYL KÉTONE-o-CARBOXY-LIC ACID C₁₅H₁₂O₃ i.e. C₆H₅.CO.CH₂.C₆H₄.CO₂H. Deoxy-benzoin-carboxylic acid. [163]. Formed by heating isobenzylidene-phthalide with NaOH (Gabriel, B. 18, 2446). Needles, v. sol. alcohol. Reduced by sodium-amalgam to C₆H₅.CH(OH).CH₂.C₆H₄.CO₂H. Ammonia forms isobenzylidene-phthalimidine. Hydroxylamine hydrochloride in alcoholic solution at 100° forms C_eH₄ < CH₂, CPh N [139°]. AgA': pp.

Anhydride v. Isobenzylidene-phthalide.

Methylamide CH₂Bz.C₆H₄.CO.NHMe. [144°]. Formed from isobenzylidene-phthalide and methylamine in alcohol at 100° (Gabriel, B. 20, 2866). White needles.

Phenyl benzyl ketone o-carboxylic acid

C.H., CH., CO.C., H., CO., H. Deoxybenzoin carboxylic acid. [75°]. Formed by boiling benzylidene-phthalide with KOHAq (Gabriel a. Michael, B. 11, 1018). Prisms (containing aq). -AgA': crystalline pp.

Anhydride v. Benzylidene-phthalide.

Amide CH2Ph.CO.C6H4.CO.NH2. Formed from benzylidene-phthalide and alcoholic NH, at 100° (Gabriel, B. 18, 2434). Needles, sol. hot water, converted into benzylidene-phthalimidine by boiling with HOAc.

Ethylamide CH.Ph.CO.C,H.CO.NHEt. [140°]. Formed from benzylidene-phthalide and alcoholic NEtH₂ at 100° (Gabriel, B. 18, 1258, 2434). Converted by boiling HOAc into C₆H₄ C(CHPh) NEt. [77°]. Hydroxylamine

forms C₆H₄<CO—ON [177°], which is also

formed by the action of hydroxylamine on phenyl benzyl ketone carboxylic acid.

Phenyl benzyl ketone dicarboxylic acid $C_6H_4(CO_2H).CO.CH_2.C_6H_4.CO_2H.$ [239°]. Formed together with an isomeride [250°] by heating phenyl-acetic-o-carboxylic acid with phthalic anhydride and NaOAc at 190° (Ephraim, B. 24, 2821). Needles. Gaseous HCl acting on its solution alcoholic forms the anhydride $C_6H_4 < CH_2.CO > C_8H_4.$ [260°]. Hydroxyl-

yields C₄H₄<CH₂ CO₂N C.C₄H₄.CO₂H [230°] crystallising in needles. Alcoholic NH,

forms crystalline C₁₆H₁₁NO₂.

PHENYL-BENZYL-METHYL-AMINE

C_{1.}H₁₈N *i.e.* NMePh.CH₂Ph. M. S. [8] 13, 39). [306°] (Nölting,

Methylo-chloride B'MeClaq. Formed from di-methyl-aniline and benzyl chloride (Michler, B. 10, 2079). Tables, v. sol. water and alcohol. Converted by successive Tables, v. sol. treatment with Ag₂SO₄ and baryta into a syrupy hydroxide, which is split up on distillation into benzył alcohol and di-methyl-aniline.

PHENYL-BENZYL-METHYL-THIO-UREA i.e. NPhMe.CS.NHC,H,. C15H16N2S [850 Formed from benzyl-thiocarbimide and methylaniline (Dixon, C. J. 59, 563). Prisms.

Isomeride NHMe.CS.NPhC,H, Formed by boiling methyl-thiocarbimide with benzyl-aniline in alcoholic solution (D.). Tufts of white prisms, v. sl. sol. hot water.

PHENYL-BENZYL-METHYL-UREA

CH₂Ph.NH.CO.NPhMe. [84°]. Formed from CH2Ph.NH.COCl and methyl-aniline (Kühn a. Riesenfeld, B. 24, 3817). V. sol. alcohol.

BENZYL-PHÉNYL-NITROSAMINE

C,H,NPh(NO). Nitrosamine of benzyl-aniline. Formed from benzyl-aniline (10 g.), alcohol (125 c.c.), H₂SO₄ (6 c.c.), and NaNO₂ in the cold, the product being poured into water (500 c.c.) (Antrick, A. 227, 360). Pale-yellowish needles, v. sol. alcohol and ether.

PHENYL-BENZYL-OXIDE v. Phenyl ether

of BENZYL ALCOHOL.

PHENYL-BENZYL-PHOSPHINE?

CH₂Ph.PHPh or C₂₃H₂₄P₂. [171°]. Formed y heating C₆H₃PCl₂ with benzyl chloride and zinc (Michaelis a. Gleichmann, B. 15, 1961). Needles, converted by successive treatment with chlorine and alkalis into $C_{12}H_{13}PO$ or $C_{23}H_{24}P_{2}O_{2}$ [155°].

Di-phenyl-benzyl-phosphine dichloride C,H,PCl₂(C,H₅)₂. [187°]. Formed from (C₆H₅)₂PCl and benzyl chloride at 180° (Dörken, B. 21, 1506; cf. Michaelis, B. 18, 2117). Prisms.

Di-phenyl-benzyl-phosphine oxide C₁H,PÔ(C₆H₅)₂. [196°]. Formed by decomposing the preceding body with water. Yields a

tri-nitro- derivative [206°].
DI-PHENYL-DI-BENZYL-SUCCINIC ACID. Nitrile. CN.CPh(CH,Ph).CPh(CH,Ph).CN. [235°]. Formed from CN.CNaPh(CH2Ph) and I (Chalanay a. Knoevenagel, B. 25, 290). crystalline powder, sl. sol. alcohol

PHENYL BENZYL SULPHONE C13H12SO2 i.e. C₆H₅.SO₂.CH₂Ph. [148°]. Formed from C.H.SO,Na and benzyl chloride (Knoevenagel. B. 21, 1344). Sl. sol. ether, m. sol. alcohol.

DI-PHENYL-BENZYL-THIOSEMICARBAZ-IDE NPh(C,H,).NH.CS.NHPh. [150°]. Formed from phenyl-benzyl-hydrazine in alcohol and phenyl-thiocarbimide (Michaelis a. Philips, A.

252, 289). Crystals, v. sol. hot alcohol. PHENYL-BENZYL-THIO-UREA C., H., N. S. i.e. NHPh.CS.NHCH,Ph. [154°]. Formed by mixing alcoholic solutions of benzylamine and phenyl-thiocarbimide (Dixon, C. J. 55, 300). Prisms (from alcohol), sl. sol. CS.

Phenyl-di-benzyl-thio-urea $C_{21}H_{20}N_2S$ i.e. NPh(C,H₁),CS.NH.C,H_r. [103°]. Formed from phenyl-thiocarbimide and benzyl-aniline (D.).

White prisms, v. sl. sol. hot water.

PHENYL-BENZYL-p-TOLYL-BIURET $C_{22}H_{21}N_3O_2$. [95°-104°]. Needles (from dilute alcohol) (Kühn a. Henschel, B. 21, 504).

PHENYL-BENZYL-UREA C,4H,4N,0 i.e. NHPh.CO.NHCH,Ph. [168°]. Formed by mixing benzyl cyanate with aniline (Letts, C. J. 25, 448). Needles, v. sol. alcohol.

Phenyl-di-benzyl-urea C21H20N2O. [128°]. Got from (C₇H₇)₂N.COCl and aniline (Hammerich, B. 25, 1819). Silky needles (from alcohol)

PHENYL-BISMUTHINE v. vol. i. p. 517.

PHENYL-BIURET C.H., N.O. i.e.
NHPh.CO.NH.CO.NH., Formed from phenylurea and PCl. (Weith, B. 10, 1744). Crystals.
s-Di-phenyl-biuret NH(CO.NHPh). [210°].

Formed by boiling allophanic ether, biuret, or

sthyl-allophanic ether, with aniline (Hofmann, B. 4, 250; Leuckhart, J. pr. [2] 21, 27). Formed also by the action of phenyl cyanate on phenylurea (Kühn a. Henschel, B. 21, 504). Needles.

u-Di-phenyl-biuret NH₂.CO.NPh.CO.NHPh. [165°]. Got from di-phenyl dicyanate and alcoholic NH₃ (Hofmann). Prisms, sol. alcohol.

s-Tri-phenyl-biuret NPh(CO.NHPh)₂. [148°]. Formed from di-phenyl dicyanate and aniline (H.) and by heating phenyl cyanate with diphenyl-ures at 150°. Prisms (from alcohol). An isomeride [105°] was got by Schiff (B. 3, 651) by distilling phenyl-carbamic ether.

References. - DI-BROMO-DI- and OXY- PHENYL-

BIURET.

PHENYL-BORATE C₆H₃BO₂. Formed by heating phenol (3 pts.) with B₂O₃ (2 pts.), or, better, by heating tri-phenyl borate with alcohol at 150° (Schiff, A. Suppl. 5, 202). Sticky mass.

Tri-phenyl borate (C₈H₅), B₃O₅. Formed by boiling phenol with B₂O₃. Glassy mass, decom-

posed by hot water.

Tetra-phenyl diborate (C₆H₅)₄B₂O₅. S.G. 2
1·124. Formed, with the preceding body, by heating C₆H₅BC₂ at 350°. Thick oil, quickly de-

composed by water.

Phenyl-boric acid C₂H₅.B(OH)₂. [204°]. Formed from C₅H₅BCl₂ and water (Michaelis a. Becker, B. 15, 181). Needles, sol. alcohol, ether, and hot water. Powerful antiseptic, with but slight physiological action. With HgCl₂ it gives a pp. of PhHgCl. Reduces ammoniacal AgNO₃, forming a mirror. On heating, it yields the oxide C₆H₅BO [190°] (above 360°), which forms crystals, sol. alcohol.

Salts.—NaA": dimetric tables.—CaH2A"2:

crystals.—AgHA": yellow pp.

Ethyl ether Et.A". (176°). Oil.

Chloride v. vol. i. p. 531.

PHENYL BROMIDÊ v. BROMO-BENZENE.

PHENYL BROMO-ALLYL OXIDE v. Bromoallyl derivative of PHENOL.

PHENYL BROMO-BENZYL KETONE

 C_6H_5 .CO.CHBrPh. [55°]. Got by brominating phenyl benzyl ketone (Knoevenagel, B.21,1355).

PHENYL BROMO-BUTYL KETONE

CaH.CO.CH.CH.CH.CH.Br. [61°]. Formed from phenyl oxy-butyl ketone anhydride or its carboxylic acid and conc. HBrAq (W. H. Perkin, jun., C. J. 51, 732; B. 19, 2559). Six-sided plates, v. sol. alcohol. Converted by warm alcoholic potash into the parent

 $CH_2 < CH_2 \cdot CH_2 \cdot CH_2 > 0.$

Phenyl di-bromo-butyl ketone

C_sH_s.CO.CH_x.CH_x.CHBr.CH₂Br. Formed from allyl-acetophenone and Br (Perkin, C. J. 45, 188). Oil. Bromine yields C₁₁H₁₁Br_sO [122°] crystallising from dilute alcohol in prisms.

PHENYL BROMO-ETHYL KETONE
C_aH_a.CO.C_aH_aBr. Formed from phenyl ethyl
ketone and Br in CS_a (Pampel a. Schmidt, B.
19, 2897). Oil with pungent odour.

PHENYL BROMO-ETHYL OXIDE v. Bromo-

ethyl derivative of Phenol

PHENYL BROMO-ETHYL SULPHONE C₄H₃.SO₂.CHMeBr. [50°]. Formed by boiling an aqueous solution of the Na salt of the acid C₄H₃.SO₂.CMeBr.CO₂H [134°], which is got by bromination of C₄H₃.SO₂.CMeH.CO₄H (Otto, J. pr. [2] 40, 550). Rectangular tables.

PHENYL BROMO-IMESATIN v. ISATIN.

PHENYL-BROMO-METHENYL-DI-ETHYL-TRI-SULPHONE C₄H₃·SO₂·CBr(SO₂·C₂H₃)₂· [135°]. Formed by bromination of the sulphone CH(SO₂Ph)(SO₂Et)₂ (Laves, B. 25, 364). Plates (from alcohol) or needles (from Aq).

TRI-PHENYL-BROMO-METHENYL TRI-SULPHONE CBr(SO₂Ph)₃. [255°]. Got by brominating CH(SO₂Ph)₃ (Laves, B. 25, 351). Amorphous insoluble pp.

PHENYL BROMO-METHYL KETONE υ. ω-Βromo-acetophenone.

PHENYL BROMO-METHYL SULPHONE C_sH₂,SO₂.CH₂Br. [48°]. Formed, together with C_sH₃,SO₂.CHBr₂ [76°], from C_sH₃,SO₂.CH_{.c}CO₂H and Br (Otto, *J. pr.* [2] 40,542). Both compounds form monoclinic tables, v. sol. hot alcohol.

PHENYL BROMO · (α)-NAPHTHYL KETONE C_αH₅.CO.C₁₉H₆Br. [98°]. Formed by bromination of phenyl (α)-naphthyl ketone (Elbs a. Steinike, B. 19, 1966). Yields a crystalline dinitro-derivative decomposing at about 90°. Yields C₁₇H₁₆Br(SO₃H) [116°] on sulphonation.

PHENYL-BROMO-NITRO-METHANE v.

Bromo-nitro-toluene.

PHENYL - BROMO - DI - NITRO - PHENYL - AMINE v. Bromo-di-nitro-di-phenyl-amine.

PHENYL - BROMO - NITRO - PHENYL-HYDRAZINE C_bH₂,N₂H₂,C_bH₃Br(NO₂). [3:1:4]. [165°]. Formed from phenyl-hydrazine and C₂H₃Br(NO₂)₂ (Willgerodt, J. pr. [2] 37, 453). Red needles (from alcohol).

PHENYL-p-BROMO-PHENYL-HYDRAZINE C₆H₅,N₂H₂,C₆H₄Br. [115°]. Formed by reduction of C₆H₅,N₂,C₆H₄Br by alcoholic ammonium sulphide (Janovsky, B. 20, 364). Tables.

PHENYL BROMO-PHENYL KETONE v.

BROMO-BENZOPHENONE.

DI-PHENYL BROMO-PROPYLENE DISUL-PHONE C,H,Br(SO,Ph),. [160°]. Got from CH,Br,CHBr,CH,Br and NaSPh in alcohol (Stuffer, B. 23, 1411). Needles.

PHENYL BROMO-PROPYL KETONE

C_aH_s.CO.CH₂.CH₂.CH₂Br. [39°]. Formed from benzoyl-trimethylene carboxylic' acid and fuming HBr (W. H. Perkin, jun., C. J. 47, 844). Crystalline mass, v. sol. alcohol.

PHENYL-BUTANE v. BUTYL-BENZENE.

Di-phenyl-butane CH,Ph.CH,CH,CH,CH,Ph. [52°]. Formed by heating di-phenyl-butylene with HI and P at 250° (Freund a. Immerwahr, B. 23, 2858). Crystals, v. sol. alcohol.

Di-phényl-butane CHMePh.CHMePh. [123.5°]. Formed by the action of zinc-dust or Na on CHMePhBr (Radziszewski, B. 7, 142; Freder R. 7, 1127). Needles (from ether)

Engler, B. 7, 1127). Needles (from ether).

Di-phenyl-butane CPh, MeEt. [128°].

Formed from CPh, Me.CO.CH, phosphorus, and
HI (Zincke a. Thörner, B. 11, 1990). Tables or

prisms, m. sol. alcohol.

Di-phenyl-butane CH₄.CH(CH,Ph),. (300°).

Formed by heating acetophenone with HIAq

and P at 180° (Graebe, B. 7, 1627). Oil.

References.—Amido-, Tri-chloro-, and Trichloro-di-nitro- and Oxy- phenyl-butanes.

PHENYL-BUTANE TRICARBOXYLIC ACID. Ethyl ether C.H., CH., C(CO., Et), CHMe.CO., Et. (338° cor.). S.G. ^{3p} 1·1003. µD 1·4850 at 20°. Formed from sodium-propane tri-carboxylic ether and benzyl chloride (Bischoff s. Mints.)

B. 23, 654). The free acid gives benzyl-

methyl-succinic acid on heating

Di-phenyl-butane dicarboxylic acid $C_0H_4(CO_2H).CH_2.CH_2.CH_2.CH_2.C_0H_4.CO_2H.$ [198°]. Formed by heating C,H,(CO.C,H,.CO,H), with HIAq and P at 160° (Gabriel a. Michael, B. 10, 2208). Nodules, insol. water.—Ag₂A": pp.

PHENYL-BUTENYL ALCOHOL C, H12O i.e. C₆H₈.C₂H(OH).C₂H₈. (225°). S.G. 19 985. Got from its acetyl derivative (223°-230°), formed from CPh:CEt by successive treatment with HBr and AgOAc (Morgan, C. J. 29, 162). Liquid. PHENYL-BUTENYL KETONE

CH2:CH.CH2.CH2.CO.Ph (285°-288°) at 710 mm. Obtained from allyl-benzoyl-acetic acid and boiling dilute alcoholic potash (W. H. Perkin, jun., C. J. 45, 187). Oil, smelling like camphor. Insol. water. Combines with Br. Excess of Br added to a glacial acetic acid solution forms bromo-phenyl-butenyl ketone dibromide, [122°].

PHENYL-BUTINENE C₁₀H₁₀ i.e. CPh:CEt. Phenyl-ethyl-acetylene. (202°). S.G. ²¹ 923.

Formed from sodium phenyl-acetylene and EtI (Morgan, C. J. 29, 162). Liquid.

Phenyl-butinene C₁₀H₁₀. (185°-190°). Got from CH_Ph_CH_2CHBr.CH_Br and alcoholic potash at 180° (Aronheim, A. 171, 231). Liquid.

Di-phenyl-butinene CHPh:CH.CH:CHPh. (320°-340°). Obtained by heating CHPh:CH.CH:CPh.CO2H (Rebuffat, G. 15, 107; 20, 154). Micaceous plates.

References.—AMIDO and Oxy - PHENYL -

BUTINENES

. DICARBOXYLIC PHENYL - BUTINENE ACID CHPh:CH.CH:C(CO₂H)₂. [208°]. Formed by heating cinnamic aldehyde with malonic acid and HOAc at 100° (Stuart, C. J. 49, 366). Needles; gives off CO2 when melted.

DI-PHENYL-DI-BUTINYL KETONE CO(CH:CH.CH:CHPh)₂. [142°]. Formed from cinnamic aldehyde, acetone, and NaOHAq (Diehl a. Einhorn, B. 18, 2325). Needles.

PHENYLBUTINYL METHYL KETONE CH₃.CO.CH:CH.CH:CHPh. [68°]. Formed from cinnamic aldehyde, acetone, and NaOHAq (Diehl a. Einhorn, B. 18, 2321). Plates (from ether). Yields a dibromide [174°] and a phenylbydrazide [180°] crystallising in plates.
DI-PHENYL-BUTONENE C₁₆H₁₂.

Phenyl naphthalene? [101°]. (346°). Formed from di-oxy-ethyl-benzene (styrolene alcohol) by (Zincke a. Breuer, A. 226, 23). Plates, v. sol. alcohol. Oxidised by K₂Cr₂O₇, and HOAc to C₁₆H₁₀O₂ [110], which crystallises in golden needles, m. sol. alcohol, and is readily polymerised by exposure of its solution to light, yielding two polymerides [207°] and [225°-229°]. The quinone $C_{16}H_{10}O_2$ is reduced by SnCl₂ to the hydroquinone $C_{16}H_{10}(OH)_2$ [93°] which gives $C_{16}H_{10}(OAc)_2$ [152°]. The quinone $C_{16}H_{10}O_2$ is $C_{16}H_{10}(OAc)_2$ [152]. The quinone $C_{16}H_{10}O_2$ is reduced by aqueous SO₂ at 120° to the quin-hydrone $C_{22}H_{22}O_4$ [183°]. On heating the quinone with NaOHAq it is converted into the oxy-quinone $C_{16}H_{6}(OH)O_2$ [144°], which gives $C_{16}H_{6}(OAc)O_2$ [111°], and may be reduced to the oxy-quinhydrone [155°], and the oxy-hydroquinone [78°]. The oxyquinone is a sideline $A(AC)O_2$ [110] and $A(AC)O_2$ [110] and $A(AC)O_2$ [111°], and $A(AC)O_2$ [110°] and $A(AC)O_2$ [111°]. converts the quinone into C_{1e}H₁₁NO₂ [174°] while ethylamine, aniline, o- and p- toluidine, and naphthylamine form compounds melting at [130°], [158°], [108°], [155°], and [148°] respectively. The compound $C_{16}H_{11}NO_2$ forms an acetyl derivative [201°], and on treatment with aqueous SO_2 at 140° gives the oxyquinone and a substance $C_{18}H_{22}O_5$ [187°].

PHENYL-sec-BUTYL-ALCOHOL C,0H,1O i.e. C₆H₅.CH₂.CH₂.CH(OH).CH₅. [68°]. Formed by reducing styryl methyl ketone with sodium-

amalgam (Engler a. Leist, B. 6, 255). Crystals. Phenyl-tert-butyl alcohol C.H.O i.e. C.H., CH., CMe., OH. [22°]. (220°-236°). Got from phenyl-acetyl chloride and ZnMe. followed by water (Popoff, B. 8, 768). Needles.

Tri-phenyl-tert-butyl-alcohol CPh,.CMe,OH. (above 260°). Formed from CCl, CMe, OH, benzene, and AlCl, (Willgerodt a. Genieser,

J. pr. [2] 37, 368).

DI-PHENYL-DI-ISOBUTYL-TETRAZONE C,H,NPh.N:N.NPh.C,H,. [107°]. Got from phenyl-isobutyl-hydrazine in ether by treatment with HgO (Michaelis a. Philips, A. 252, 284).

PHENYL-BUTYLENE C₁₀H₁₂ i.e. C₂H₃.CH₂.CH₂.CH₃.CH₄.CH₂.C chloride, allyl iodide, and Na (Aronheim, B. 5, 1068; A. 171, 219). The same hydrocarbon appears to be formed by distilling the lactonic appears to be formed by distinct the acid of y-oxy-y-phenyl-s-di-methyl-succinic acid (Penfield, A. 216, 125). Oil, yielding a liquid dibromide. KNO₂ and HOAc form C₁₀H₁₂N₁O₃, which gives on reduction a base $C_{10}H_{15}NO$, whence B'HCl and B'₂H₂PtCl₈ (Tönnies, B. 11,

(a)-Phenyl-butylene C₆H₆.CH:CH.CH₂.CH₃. (186°). Formed by distilling C₁₀H₁₁Br obtained by bromination of n-butyl-benzene (Radziszewski, B. 9, 260). Forms a dibromide $C_{10}H_{12}Br_2$ [71°] crystallising in white needles. Probably identical with the phenyl-butylene (187°) obtained by Perkin (C. J. 32, 667; 35, 140) from phenylangelic acid by successive treatment with HBr and Na CO Aq or with HI and KOH. Perkin's phenyl-butylene forms a crystalline dibromide [67°] converted by alcoholic potash into oily C₁₀H₁₁Br, which forms oily C₁₀H₁₁Br₂.

(3)-Phenyl-butylene C₆H₅CH:CMe₂. (185°).

Formed by heating benzoic aldehyde with sodium isobutyrate and isobutyric anhydride for 3 hours with inverted condenser (Perkin, C. J. 35, 138,, and by distilling \$\beta\$-oxy-phenyl-valeric acid (Fittig a. Jayne, \$A\$. 216, 118). Oil. Forms an oily dibromide converted by alcoholic potash into C₁₀H₁₁Br, which forms crystalline C₁₀H₁₁Br, [63.5°]. Yields benzoic and acetic acids on oxidation. Nitrous acid forms C₁₀H₁₂N₂O₃ [112°]

(Angeli, B. 25, 1962).

acrylic acid (Freund a. Immerwahr, B. 23, 2858). Crystals, v. sol. alcohol. Yields a dibromide [83°].

DI-PHENYL-BUTYLENE-DIAMINE C₄H₈(NHPh)₂. Formed from isobutylene dihydroquinone [73°]. The oxyquinone is oxidised by alkaline KMnO, to an acid C₂H₂O₃ [177°-197°] whence BaA"2aq, K₂A", Cu₂(OH)₂A"6aq, and Ag₂A". Alcoholic NH₂ [122°]. S. 10 at 15°; 20 at 100°.

DI-PHENYL-ISOBUTYL-GLYOXALINE

CPh.NH CPh.N >C.CH₂₽r. C, H20N2 i.e.

Formed from benzil, isovaleric aldehyde, and NH, Aq (Japp a. Wynne, C. J. 49, 468).— B"H,PtCla: crystalline.

PHENYL-ISOBUTYL-HYDRAZINE

C₄H₂NPh.NH₂. (240°-245°). Formed from isobutyl bromide and sodium phenyl-hydrazine (Michaelis a. Philips, B. 20, 2485; A. 252, 282; 270, 122). Liquid, which reduces hot Fehling's solution. Converted by thionyl-aniline into oily C.H. NPh.N:SO-B'H.SO.: plates.

Acetyl derivative C.H. NPh.NHAc.

[114°

PHENYL BUTYL KETONE C.H. CO.CH.Pr. Mol. w. 162. (237°) at 720 mm. Got by boiling propyl-benzoyl-acetic ether with alcoholic pctash (Perkin a. Calman, C. J. 49, 162). Oil.

Phenyl isobutyl ketone C.H. CO.CH.Pr. (228°) at 720 mm. S.G. 175 .993. Formed from isopropyl-benzoyl-acetic ether (P. a. C.) and also by distilling a mixture of calcium benzoate and calcium isovalerate (Popoff, A. 162, 153). Liquid. Does not combine with NaHSO₃. Yields benzoic, isobutyric, and acetic acids on oxidation.

PHENYL BUTYL KETONE CARBOXYLIC ACID C.H. CO.CH2. CHEt.CO2H. Ethyl-benzoylpropionic acid. [83°]. Formed by heating the dicarboxylic acid. Small needles.—CaA'₂ aq.

Ethyl ether EtA'. Oil.

Phenyl butyl ketone carboxylic ether C_aH_a.CO.CHPr.CO₂Et. Propyl - benzoyl - acetic acid (251° at 300 mm.). Formed from benzoylacetic ether, NaOEt, and PrI at 100° (Perkin a. C₆H₅.CO.CHPr.CO₂Et. Calman, C. J. 49, 160). Liquid.

Phenyl isobutyl ketone carboxylic ether C₈H₈.CO.CHPr.CO₂Et. (237° at 225 mm.). Formed

in like manner, using isopropyl iodide.

Phenyl butyl ketone dicarboxylic acid C₄H₅.CO.CH₂.CEt(CO₂H)₂. Benzoyl-ethyl-isosuc-cinic ether. Got by saponifying its ether, which is formed from sodium-malonic ether and wbromo-acetophenone (Dittrich a. Paal, B. 21, 3453). Crystalline mass.— $(NH_4)_2A''$: amorphous.— K_2A'' : pearly plates, v. sol. water.—CaA'' aq.— Ag₂A": plates. Hydrazine salt. [103°].

Phenyl hydrazide C., H., N.O. [132°].
PHENYL BUTYL METHYLENE DIKETONE C₆H₅.CO.CH₂.CO.CH₂Pr. Valeryl-acetophenone. (184° at 30 mm.). Oil. Got from acetophenone, sovaleric ether, and NaOEt (Stylos, B. 20, 2181).

PHENYL - BUTYL MÈTHYL KETONE CARBOXYLIC ETHER

CH₂.CO.CEt(CH,Ph).CO.Et. (c. 297°). Formed from sodium benzyl-acetoacetate and EtI (Conrad, B. 11, 1057).

PHENYL BUTYL OXIDE v. Butyl derivative

of PHENOL

PHENYL-ISOBUTYLPHENYL-THIO-UREA C.H.,NH.CS.C.H.,CH.Pr. [152°]. Formed from C.H. (NH₂).CH₂Pr and phenyl-thiocarbimide (Mainzer, B. 16, 2023). Plates, sol. alcohol.

DI-PHENYL-ISOBUTYL-QUINOXALINE

 $C_sH_s(CH_2Pr) < N:CPh N:CPh$ [144°]. Formed from benzil and isobutylphenylene-diamine (Gelzer, B. 20, 8257). Needles, v. sol. alcohol.—B'2HCl. PHENYL-BUTYL-THIO-ALLOPHANIC ACID

NHPh.CS.N(C,H,).CN. [139°]. Formed from sodium cyanamide, phenyl thiocarbimide, and

isobutyl iodide (Hecht, B. 25, 822). Slender needles, v. sl. sol. hot water.

DI-PHENYL-ISOBUTYL-THIO-SEMI-CARB-NHPh.CS.NH.NPh(CH₂Pr). Formed from phenyl-isobutyl-hydrazine and phenyl-thiocarbimide (Michaelis a Philips, A. 252, 284). White crystals.

PHENYL-ISOBUTYL-THIO-UREA

NHPh.CS.NHC₄H₉ [82°]. Formed from phenylthiocarbimide and isobutylamine (Hecht, B. 25, 815). Needles, m. sol. hot water.

PHENYL-BUTYL-THIOHYDANTOIN

CS<NPh.CO NH.CH(C,H,): [179° uncor.]. Formed by fusing phenyl-thiocarbimide with leucine (Aschan, B. 17, 426). Minute colourless prisms.

PHENYL-BUTYRIC ACID C₁₀H₁₂O₂ i.e. CH₃CH₂.CHPhCO₂H. [42°]. (272°). Formed by saponifying the nitrile which is made by the action of EtI and solid NaOH on benzyl cyanide (Neure, A. 250, 153). Crystalline mass.—CaA'₂ 2aq; groups of needles.—AgA'.

Methylether MeA'. (228°).

Nitrile PhCHEt.CN. $(245^{\circ}).$

 γ -Phenyl-n-butyric acid

CH₂Ph.CH₂.CH₂.CO₂H. [47·5°]. (290°). Formed from oxy-phenyl-butyric acid and HI (Burcker, A. Ch. [5] 26, 459) and from phenyl-isocrotonic acid by protracted treatment with sodium-amalgam (Jayne, A. 216, 107). Long flat plates (from water).—CaA'₂: amorphous mass.
Phenyl-isobutyric acid CH₂Ph.CHMe.CO₂H.

Benzyl-methyl-acetic acid. [37°]. (272°). S.

·31 at 15°.

Formation.-1. By heating benzyl-methylmalonic acid (Conrad, B. 13, 598).—2. By heating benzyl-methyl-acetoacetic ether with conc. KOHAq .- 3. By reduction of the phenylcrotonic acid obtained from benzoic aldehyde, sodium propionate, and propionic anhydride.-4. By reducing phenyl-angelic (methyl-cin-namic) acid with sodium-amalgam (von Miller, B. 23, 1888).

Properties .- Plates. Converted by H2SO, at

150° into oxy-methyl-indonaphthene.

Salt .-- AgA'. S. 248 at 20°. Ethylether EtA'. (287°). S.G. ‡ Oil (Conrad a. Bischoff, A. 204, 177). Benzylether C,H,A'. (320°-325°). S.G. 23 1.05.

165 1.046. A product of the action of sodium on benzyl propionate (Conrad a. Hodgkinson, A. 193, 312).

[109°]. Amide CH₂Ph.CHMe.CONH₂. Formed by heating the NH, salt (Edeleano, B. 20, 618). Needles, v. sol. alcohol and ether.

Di-phenyl-butyric acid

[126°]. CH,Ph.CMePh.CO,H. Got from its nitrile, which is obtained from a-phenyl-propionitrile and benzyl chloride (Janssen, A. 250, Needles, v. sol. ether and alcohol. NaA' 7aq: long thin needles.—CaA'₂.—BaA'₋.—CuA₂'. [73°].—AgA': white insoluble powder.

Nitrile C₁₅H₁₅.CN. (337°). Oil.

Isomeride v. DI-BENZYL-ACETIC ACID. References .- Bromo-, NITRO-, NITRO AMIDO-, and Oxy- PHENYL-BUTYRIC ACID.

PHENYL-ISOBUTYRIC ALDEHYDE

CH₃.CH(CH₂Ph).CHO. (227°). Formed by distilling the corresponding lime salt with calcium formato (Miller a. Rohde, B. 23, 1080). Oil.

γ-PHENYL-BUTYBIC-o-CARBOXYLIC ACID C₁₁H₁₂O₄ i.e. C₆H₄(CO₂H).CH₂.CH₂.CH₂.CO₂H. [189°]. Formed by heating at 180°-190° the double-lactone of benzoyl-propionic-o-carboxylic acid $C_0H_4 < \frac{CO \cdot O}{CO \cdot CO} > C + \frac{CH_2}{O \cdot CO} > CH_2$ with HI and P (Roser, B. 18, 3118). Small plates. V. sol. alcohol, sl. sol. cold water.—A"Ba: very soluble in water.

PHENYL-BUTYRO-LACTONE v. Lactone of

OXYPHENYL BUTYRIC ACID.

PHENYL-CACODYL v. vol. i. p. 320.

PHENYL-CAMPHORAMIC ACID v. CAM-PHORIC ACID.

PHENYL-CAMPHYL-THIO-UREA v. CAM-

PHYL-THIO-UREA.

PHENYL CARBAMATE NH2.CO2Ph. [143°]. Formed by the action, in ethereal solution, of NH₂ on ClCO₂Ph or of NH₂.COCl on phenol (Kempf, B. 2, 740; Gattermann, A. 244, 43). Formed also, together with Et₂CO₃, by heating phenyl ethyl carbonate at 300° (Bender, B. 19, 2268). Needles (from water), sol. alcohol and ether. Decomposed by NaOHAq into NH3, phenol, and Na.CO. NH.Aq at 150° forms phenol and urea. PHENYL-CARBAMIC ACID NHPh.CO.H.

Carbanilic acid. The ethers of this acid are got by the action of aniline on the chloroformic ethers Cl.CO.OR, and also by the action of phenyl cyanate on alcohols. They are decomposed by

potash into CO2, aniline, and alcohols.

Acetyl derivative NPhAc.CO.H. The Na salt is formed by passing CO₂ over sodium acetanilide in the cold (Seitert, B. 18, 1358). It is decomposed by water into acetanilide and NaHCO3. By heating at 140° under pressure it is converted

into NPhH.CO.CH2.CO2Na.

Methylether NHPh.CO.OMe. [47°] (Hentschel, B. 18, 978). Prisms. Converted by HNO, (S.G. 1·48) into C₆H₂(NO₂)₂NH.CO₂Me [127°] and [6:4:2:1]C₆H₂(NO₂)₃NH.CO₂Me [192°] (Van Romburgh, E. T. C. 10, 135). H₂SO₄ forms SO, H.C, H, NH.CO, Me which is converted by bromine into C.H.3Br.NH.CO.Me [96.5°] (Hentschel, J. pr. [2] 34, 423). Distillation with lime at 260° gives aniline, methyl-aniline, di-methyl-aniline, and di-phenyl-urea (Nölting, B. 21, 3154).

Ethyl ether NHPh.CO₂Et. [52°]. (238°) (Wilm a. Wischin, C. J. 21, 192). Formed as above and also by boiling the product of the action of phenyl-urea on aceto-acetic ether with conc. HCl (Behrend, A. 233, 6) and by the action of sodium acetanilide on chloro-formic ether (Paal a. Otten, B. 23, 2590). Needles (from water). Not affected by boiling conc. HClAq, but decomposed by HClAq at 150° into CO2 aniline, and EtCl. Boiling alcoholic KOH gives aniline and K₂CO₂. Yields a bromoderivative [81°], a di-nitro-derivative [110°], and a tri-nitro- derivative [144°]. When distilled with NaOPh at 220° it gives phenol and diphenyl-urea [235] (Hentschel, J. pr. [2] 27, 498).— NKPh.CO₂Et. Formed by dissolving phenyl-carbamic ether in alcoholic potash. Hygrorsopic needles, decomposed by water.

Chloro-ethyl ether

NHPh.OO₂CH₂CH₂Cl. [51°]. Formed from aniline and Cl.CO.OC.H₄Cl (Nemirowsky, J. pr. [2] 31, 174; Otto, J. pr. [2] 44, 15). Needles, sl. sol. hot water. When boiled alone and after-

wards with conc. KOH it yields NPh CO2>CH, [124°], which is converted by HClAq at 170° into chloro-ethyl-aniline.

Ethylene ether C.H.A". [158°]. Formed from ethylene glycol and phenyl cyanate (Snape, B. 18, 2430; C. J. 47, 773). Prisms. Propyl ether PrA'. [59°]. Slender needles,

v. sol. alcohol (Römer, B. 6, 1101).

Isopropyl ether PrA'. [90°] (Gumpert, J. pr. [2] 81, 119; 32, 278); [43°] (Spica, G. 17, 165). Needles (from dilute alcohol).

Di-chloro-propyl ether (Otto, J. pr. [2] 44, 22) CH_Cl.CHCl.CH_A'. [74°]. Prisms.

Isobutyl ether C_H_A'. [80°]. (216°).

Needles, v. sol. alcohol (Mylius, B. 5, 972).

Heptenyl ether C.H.A'. [85°]. Formed by the action of phenyl cyanate on the heptenyl alcohol got by reducing suberone (Markownike f., C. R. 110, 466). Prisms (from alcohol).

Glyceryl ether v. GLYCERYL-TRI-PHENYL-

TRI-CARBAMATE.

Phenyl ether NHPh.CO.Ph. [126°]. Formed from phenyl cyanate and phenol (Gumpert; cf. Hofmann, B. 4, 245). Needles (from benzene). With NH, Aq it forms phenyl-urea.

Phenylene ethers. The o [165°], m [164°], and p [207°] compounds are got by heating pyrocatechin, resorcin, and hydroquinone respectively with phenyl cyanate (Snape, B. 18, 2428).

Phenylphenyl ether C.H.C.H.A' [110°]. Formed from p-amido-diphenyl and

ClCO₂Et (Zimmermann, B. 13, 1965).

(a)-Naphthyl ether C10H2A'. Slender needles (Leuchart a. Schmidt, B. 18, 2340; Snape, C. J. 47, 776). Decomposed by heat into (a)-naphthol and phenyl cyanate.

(β) - Naphthyl ether C₁₀H,A'. [230°]. (L. a. S.); [155°] (S.). Thick prisms. The tetrahydride C₁₀H₁₁O.CONPhH [98·5°] is formed from (B)-naphthol ac-tetrahydride and phenyl cyanate.

Di-phenyl-carbamic acid.

Chloride NPh2.COCl. [85°]. Formed from COCl₂ and diphenylamine in CHCl₃ (Michler,

B. 9, 396). White scales (from alcohol).

Ethyl ether NPh₂CO₂Et. [72°]. (above 360°). Formed by heating diphenylamine with ClCO₂Et (Merz a. Werth, B. 6, 1511; Hager, B. 18, 2573). Large prisms (from benzene)

Phenyl ether PhA'. [104°]. Formed from the chloride and KOPh (Lellmann a. Bor höffer, B. 20, 2122). The compound C.H. (NO2)A' [114°], made in like manner from o-nitrophenol, may be reduced to $C_6H_4(NH_2)A'$ [191°]. The mand p-nitro-phenyl ethers [90°] and [116°] yield m- and p- amido-phenyl ethers [133°] and [146°] (Lellmann a. Benz, B. 24, 2111)

p-Tolyl-ether C,H,A'. [81°]. (L. a. B.). References.—Amido-, Bromo-, DI-BROMO-NITEO-, NITEO-, NITEO-AMIDO-, NITEO-OXY-, OXY-, OXY-AMIDO-, and NITRO- PHENYL-CARBAMIC ACID

and ETHER.

PHENYL-CARBAMINE C.H.NC. Mol. w. 103. (167°). S.V. 121.6 (Lossen, A. 254, 73). Formed by distilling aniline with chloroform and alcoholic potash (Hofmann, A. 144, 117). Stinking liquid, quickly decomposed by acids into aniline and formic soid. Changes into the isomeric benzonitrile by heating at 210° (Weith, B. 6, 210). Yields phenyl-thiocarbimide on heating with sulphur. Mixed with propionitrile and ether, it yields, by successive treatment with sodium and water, the compound CPh(NH).CHMe.CN [97°] (Von Meyer, J. pr.

[2] 39, 189).

Chloride NPh.CCl₂. (212°). Formed from phenyl-thiocarbimide and Cl (Sell a. Zierold, B. 7, 1228). Pungent liquid, with nasty smell, yielding s-di-phenyl-urea when heated with water at 100°

PHENYL-CARBAZIC ACID. The crystalline salt N2H2Ph.CO.ON2H4Ph is formed by the action of CO2 on phenyl-hydrazine and water. It is deliquescent, and sl. sol. water and ether (Fischer, A. 190, 124).

Methyl ether NHPh.NH.CO, Me. [117°].

Short prisms (Heller, A. 263, 281).

Ethyl ether NHPh.NH.CO, Et. [87°] (F.); [82°] (N.). Formed from phenyl-hydrazine and ClCO₂Et (E. Fischer, B. 22, 1936), or AcC(CO₂Et)₂ (Nef, A. 266, 107). Needles. Ac₂O forms C₆H₂, N₂HAc.CO₂Et [103°] (H.).

PHENYL-semi-CARBAZIDE C₂H₂N₃O i.e.

NHPh.NH.CO.NH₂. [172°]. Formed from potassium cyanute and phenyl-hydrazine hydrochloride (E. Fischer, A. 190, 113; Freund, B. 21, 2463). Formed also by heating phenylhydrazine hydrochloride (1 mol.) with urea (2 mols.) at 160° for 4 hours (Pinner, B. 20, 2358; 21, 2329) and by heating di-phenyl-carbazide with urea (Skinner a. Ruhemann, C. J. 53, 550; B. 20, 3373). Prisms, v. sol. hot water. Yields di-oxy-phenyl-triazole [263°] on heating with urea. By heating at 160° it is converted into di-phenyl-urazine $C_{14}H_{12}N_4O_2$ [264°] and other products. COCl₂ forms $C_8H_7N_3O_2$ [167°].

Bensoyl derivative NPhBz.NH.CO.NH2. [203°]. Formed from benzoyl-phenyl-hydrazine cyanate (Michaelis a. Schmidt, A. 252, 317).

Di-phenyl-semi-carbazide

[173°]. C₄H₅NH.CO.N₂HPh. Formed from phenyl-hydrazine by combination with phenyl cyanate; and also by heating phenyl-hydrazine with phenyl-urea (Kühn, B. 17, 2884; Skinner, C. J. 53, 552). Needles or plates (from alcohol or benzene), sl. sol. water. COCl₂ in benzene forms C₁₄H₁₁N₂O₂ [173°] (Freund, B. 21, 2465). Di-phenyl-carbazide CO(N₂H₂Ph)₂. [151°]

(S. a. K.); [164°] (E. Fischer, B. 22, 1930). Formed by heating phenyl-hydrazine (2 mols.) with carbamic ether (1 mol.) (Skinner a. Ruherann, C. J. 53, 550; B. 20, 3372). Formed also from phenyl-hydrazine and COCl₂ (Heller, A. 263, 277). Crystalline. Forms with HgCl₂ a crystalline compound B'HgCl₂ not melted at 136°. Alcoholic potash forms a red solution containing di-phenyl-carbazone N2H2Ph.CO.N:NPh crystallising in orange needles [157°]. Benzene and CSCl₂ form CS CNPh.N C.N:NPh [170]

(Freund a. Kuh, B. 23, 2833). PHENYL-CARBIMIDE v. Phenyl isocyanate,

vol. ii. p. 315. DI-PHENYL-DI-CARBIMIDO-TETRA-p-AMIDO-TETRA-TOLYL-DI-o-SULPHIDE

PhN:C NH.C,H.,S.C,H.,NH C:NPh. Di-thiop-tolyl-di-phenyl-di-guanidine. [c. 119°]. Formed by heating di-thiocarbonyl-tetra-amido-tetratolyl-di-sulphide with aniline and HgO (Truhlar, B. 20, 674). Amorphous; v. sol. alcohol.

DI-PHENYL-CARBINOL C13H12O i.e.

Ph₂.CH(OH). Benshydrol. Mol. w. 184. [68°]. (298°). S. 05 at 20°. Initial velocity of etherification 22 (Menschutkin, J. R. 1882, 162). Formed by reducing benzophenone with sodiumamalgam (Linnemann, A. 133, 6; Beckmann, B. 22, 915); or by heating benzophenone with zinc and alcoholic KOH (Zagumenny, A. 184, 174). Slender silky needles, v. e. sol. alcohol and ether. Converted into benzophenone by oxidation with chromic acid. Bromine forms a dibromo-benzhydrol [163°]. Zn and HOAc form benzpinacone. Zn and HCl in acetic acid solution reduce it to tetra-phenyl-ethane. Distillation partly resolves benzhydrol into water and benzhydrolic ether. P₂S₃ forms C₂₅H₂₂(SH)₂ [151°] and oily Ph₂CH(SH) (Engler, B. 11, 922).

Ethyl derivative C18H1, OEt. S.G. 20 1.03. From benzhydrol, alcohol, and H₂SO₄, or from Ph₂CHBr and alcoholic KOH (Friedel a. Balsohn, Bl. [2] 33, 339). Liquid.

Isoamyl derivative C13H11OC3H11. (310°). Chloride Ph₂CHCl. [14°]. From benzhydrol and HCl (Engler a. Bethge, B. 7, 1128).

Bromide Ph₂CHBr. [45°]. From di-phenylmethane and bromine at 150° (F. a. B.). Water at 150° decomposes it into benzhydrol and benzhydrolic ether. Conc. NH₂Aq forms monophydrolic ether. and di-benzhydryl-amine (Friedel a. Balsohn, Bl. [2] 33, 587).

Acetyl derivative Ph2CH.OAc. [42°]. (302°). S.G. 22 1.49. Prisms, v. sol. alcohol, acetic acid, and ether (Vincent, Bl. [2] 35, 304).

Benzoyl derivative Ph₂CH.OBz. [89° Non-volatile. Trimetric, a:b:c=1:477:668.

Succinyl derivative

 $(Ph_2CH.O)_2$: \tilde{C}_2O_2 : C_2H_4 . [142°]. Gives, on distillation, succinic acid and benzhydrolene C. H. [210°].

Anhydride $C_{26}H_{20}O$ i.e. $Ph_2C-CPh_2(?)$

[111°]. (315°) at 745 mm. From benzhydrol by long boiling with water or by treatment with PCl₃, BzCl, or diluted H₂SO₄. Monoclinic crystals (from benzene). Reduced by zinc and HCl in HOAc to tetra-phenyl-methane (Zagumenny, J. R. 12, 431). Zincke a. Thörner (B. 11, 1398) obtained this anhydride by heating benzpinacone, $C_{23}H_{22}O_2$, assume the formula $C_{20}H_{20}O$. and consequently

References .- DI-AMIDO-, DI-BROMO-, and OXY-DI-PHENYL-CARBINOL.

Tri-phenyl-carbinol C, H, O i.e. CPh, OH.

Mol. w. 260. [159°]. (above 360°).

Formation. — 1. By boiling tri-phenylmethane with chromic acid mixture (Hemilian, B. 7, 1203).—2. From CPh, Br and water.—3. By the action of water on CPh,Cl got from AlCl₃, benzene, and CCl₄, or CCl₃.COCl (Friede a. Crafts, A. Ch. [6] 1, 499; Hentschel, J. pr. [2] 36, 311).—4. By heating its dicarboxylic acid with baryta (Hemilian, B. 19, 3073).

Properties. - Six-sided plates, v. sol. alcohol. Acetyl derivative CPh. OAc. [99°]. Acetyl aerward. Prisms (Allen a. Kölliker, A. 227, 116). Prisms (Allen a. Kölliker, A. 227, 116). Laminæ.

Ethyl ether CPh.OEt. [79°] (F. a. C.); [83°] (A. a. K.). Chloride v. Chlobo tri-phenyl-methans.

DI-PHENYL-CARBINOL CARBOXYLIC | ACID v. OXY-BENZYL-BENZOIC ACID.

Di-phenyl-carbinol dicarboxylic acid

CH(OĤ)(C_eH₄.CO_eH)₂. Formed by heating benzil di-o-carboxylic (diphthalylic) acid with KOHAq (50 p.c.) at 130° for 5 minutes (Graebe a. Juillard, A. 242, 238). The acid splits up at the moment of liberation into water and anhydride.-BaA" aq.

Anhydride C₁₈H₁₀O₄. [203°]. S. ·022 at 23°. Formed as above, and also by reducing benzophenone di-o-carboxylic acid. Monoclinic crystals (from alcohol). On heating, it yields a white sublimate [172°] reconverted by treatment with NaOHAq and HCl into the original anhydride. Phenyl-hydrazine gives C₂₁H₁₆N₂O₃. H₆SO₄ and HNO₂ give C₁₅H₆(NO₂)₂O₄ [c. 275°], which forms EtA' [148°].—Ba(C₁₅H₉O₄)₂ 2½aq.—CuA'₂ 8aq.—AgA'.—C₁₅H₉MeA'. [155°].—EtA'. [100°].—C₁₅H₉O₃(NH₂). [160°]. Needles.

Isomeride v. OXY-BENZYL-ISOPHTHALIC ACID.

Di-phonyl-carbinol tri-carboxylic acid

C(OH)(CO₂H)(C₀H₄.CO₂H)₂.

An hy dride C_{1e}H_{1e}O_e. [170°]. Formed by heating benzil di-o-carboxylic acid with NaOHAq (4 p.c.) at 112° for 3 minutes (G. a. J.). Minute crystals, sl. sol. cold water. Gives off CO2 on heating, yielding the anhydride of di-phenyl-carbinol dicarboxylic acid.

Ethers of the anhydride Me2A". [148°]. -Et₂A". [108°]. Prisms, v. sol. alcohol.

Tri-phenyl-carbinol o-carboxylic acid

*CPh2(OH).C.H4.CO2H. The Na salt is formed by heating di-phenyl-phthalide with NaOHAq, but the acid splits up on liberation into water and its anhydride, di-phenyl-phthalide (Baeyer, A. 202, 50).

Tri - phenyl - carbinol m - carboxylic acid. [162°]. Formed by oxidation of di-phenyl-mtolyl-methane with CrO, and HOAc (Hemilian, B. 16, 2369). Trimetric plates.

Tri-phenyl-carbinol p-carboxylic acid (?) [187°]. Formed by oxidation of CHPh. C.H. CHO (Oppenheim, B. 19, 2028) or of di-phenyl-tolylmethane (Hemilian, B. 7, 1210). Needles.—BaA', 7aq. Needles, v. sl. sol. cold water.

Tri-phenyl-carbinol dicarboxylic acid. An-

hydride $CPh_2 < \frac{C_0H_2(CO_2H)}{CO} > \frac{C_0H_2(CO_2H)}{CO}$

Di-phenyl-phthalide carboxylic acid. [246°]. Formed by oxidation of di-phenyl-y-xylyl-methane, di-phenyl-tolyl-methane o-carboxylic acid, di-phenyl-tolyl-carbinol m-carboxylic acid, or di-phenyl-methyl-phthalide (Hemilian, B. 16, 2372). Crystals, v. sol. alcohol and HOAc. Yields benzophenone and terephthalic acid on fusion with potash. Zinc-dust and NaOHAq reduce it to tri-phenyl-methane dicarboxylic acid.

Tri-phenyl-carbinol dicarboxylic acid C₂₁H₁₈O₅ i.e. CPh₂(OH).C₅H₃(CO₂H)₂ [1:3:4]. [180°]. Formed by oxidation of di-phenyl-xylylmethane with $K_1Gr_2O_2$ and H_2SO_4Aq (Hemilian, B. 19, 3072). Needles (from water), v. e. sol. alcohol. On fusion it yields an amorphous anhyuride C_1 $H_{14}O_4$. On fusion with Ba(OH), it forms tri-phenyl-carbinol.—Ag2A": pp.

Tri-phenyl-carbinol dicarboxylic acid CPh₂(OH).O₄H₂(CO₂H)₂ [1:2:4]. Anhydride C₂₁H₁₄O₄. Di-phenyl-phthalide carboxylic acid. [223°]. Formed by oxidation of the corresponding di-phenyl-xylyl-methane (H.). Tables (con-

taining EtOH).—CaA', Saq.—AgA': needles.
DI-PHENYL-CARBINYLAMINE C, H, N oc. CHPh2.NH2. Benzhydrylamine. Exo-amidodi-phenyl-methane. (289°). Formed, together with tetra-phenyl-di-carbinyl-amine, by leaving benzhydryl bromide, CHPh2Br, with conc. NH, Aq for 48 hours (Friedel a. Balsohn, Bl. [2] 33, 587). Got also by reducing benzophenone. oxim in alcoholic solution by sodium-amalgam and acetic acid (Goldschmidt, B. 19, 8283).

Properties.—Alkaline liquid, absorbing CO.

from the air.

Salts.—B'HCl: [270°]; long needles, sl. sol. cold water.—B'2H2PtCl22aq: lancet-like needles (L. s. B.); B'H,PtCl, aq (G.).—*B',H,CO, [91°].
Formyl derivative Ph,CH.NH.CHO.

[182°]. (860°). Formed, almost quantitatively, by heating benzophenone with ammonium formate at 200°-220° (Leuchart a. Bach, B. 19, 2129).

Urea Ph.CH.NH.CO.NH2. [143°] (L. a. B.). Tetra-phenyl-di-carbinyl-amine (Ph.CH), NH. [136°]. Prepared as above, crystallises from alcohol in slender needles, not affected by MeI

or AcCl at 100° (F. a. B.).

Tri-phenyl-carbinyl-amine CPh₃.NH₂. [103°]. Formed by passing dry NH, through a solution of CPh.Br in benzene (Hemilian a. Silberstein, Bl. [2] 43, 118; B. 17, 741; cf. Nauen, B. 17, 442). Needles, sl. sol. cold alcohol. Yields B'HCl, B',H,PtCl, 7 aq, CPh,NHMe [73°], CPh,NMe, [97°], CPh,NHPh [145°], CPh,NHCH,Ph [110°], and the acetyl derivative CPh, NHAc [208°]. Forms a crystalline oxalate [253°] (Elbs, B. 17, 701), and the compounds C19H17NBr2, B'I2, and

PHENYL-CARBINYL CHLORIDE v. CHLORO-

TRI-PHENYL-METHANE.

TRI-PHENYL - CARBINYL - MALONIC ETHER CPh_s.CH(CO₂Et)₂. [133.5°]. Got from sodium malonic ether and CPh.Br (Henderson, B. 20, 1014). Needles (from alcohol

TRI-PHENYL-CARBINYL SULPHOCYAN-IDE CPh. NCS. [137°]. Formed from CPh. Br and ammonium sulphocyanide (Elbs, B. 17, 700).

TRI - PHENYL - CARBINYL - TOLUIDINE CPh. NHC, H₁. The o [142°] and p [177°] compounds are formed from CPh.Br and o and p toluidine respectively (Elbs, B. 17, 706). The p-compound yields a crystalline nitrosamine [145°_148°]

PHENYL-CARBIZINE. Formyl derivative C_eH_aNN.CHO. [72°]. (256°). Formed by the action of COCl2 in benzene on the phenyl-hydrazide of formic aldehyde (Freund, B. 21, 1240, 2458). Needles (from CS₂), v. sol. alcohol and ether. Decomposed by boiling Na₂CO₂Aq into CO₂, and the phenyl-hydrazide of formic aldehyde.

Acetyl derivative CO NPh. (280°). Formed in like manner from acetic aldehyde. Monoclinic prisms; a:b:c=1.210:1:1.557; $\beta = 73^{\circ} 23'$.

Propionyl derivative. [63°]. Needles. Benzoyl derivative CO NBz [114°].

Needles (from alcohol) or plates (from HOAc). PHENYL CARBONATES.

Phenyl-carbonic acid *C.H.O.CO.H.

Sodium salt Pho.Co.Na. Prepared by treating dry sodium phenylate in the cold for a month with CO, until it has taken up the calculated quantity of the gas. The product is a very

hygroscopic powder.

Reactions.-1. With water it gives off half its CO2 in gaseous form: 2PhO.CO2Na+H2O = $PhONa + PhOH + NaHOO_8 + CO_2 - 2$. At 120° it splits up into CO, and PhONa.—3. Heated rapidly to 180°-200° it gives off smaller quantities of CO₂ together with phenol, the residue being sodium salicylate.—4. In a sealed tube at 120°-130° for several hours it changes completely to sodium salicylate: C6H5O.CO2Na = $C_aH_4(OH)CO_2Na$ (R. Schmilt, J. pr. [2] 31,

Di-phenyl carbonate $C_{13}H_{10}O_3$ i.e. $CO(OPh)_2$. [78°] (Kempf, J. pr. [2] 1,404; Hentschel, J. pr. [2] 27, 41; 36, 315); [88°] (Richter, J. pr. [2] 27, 41). (303°). Formed by passing COCl, into an aqueous solution of NaOPh, or into phenol containing AlCl₂. Formed also from ClCO₂CCl₃, phenol, and AlCl₂. Needles (from alcohol). Alcoholic potash forms phenol and K₂CO₃. NaSEt gives NaSPh and (EtS)₂CO (Seiffert, J. pr. [2] 31, 464).

References .- DI-BROMO- and NITRO- PHENYL

CARBONATE

DIPHENYL O-CARBOXYLIC ACID C18H10O2 i.e. C.H. C.H. CO.H [1:2]. Phenyl-benzoic acid. Mol. w. 198. [111°]. Formed by potash-fusion from diphenylene ketone (Fittig a. Ostermaier, B. 5, 933; A. 166, 374; Schmitz, A. 193, 115). Formed also, together with diphenylene ketone oxide, by distilling sodium salicylate with triphenyl phosphate (R. Richter, J. pr. [2] 28, 305). Small needles (from hot alcohol), m. sol. hot

water.—KA'aq.—BaA', aq.—CaA', 2aq.—AgA'.

Ethyl ether EtA'. (300°-305°).

Tetrahydride C_eH₃.C_eH₁₀.CO₂H. Phenylhexamethylene carboxylic acid. [105°]. Formed
by heating the corresponding dicarboxylic acid, and by the hydrolysis of phenylhexamethylenyl methyl ketone carboxylic ether (Kipping a. W. H. Perkin, jun., C. J. 57, 319). Rosettes of

crystals (from ligroïn).—AgA': amorphous pp.
Diphenyl m-carboxylic acid C₆H₅.C₆H₄.CO₂H. [161°]. Formed by oxidation by chromic acid and HOAc from [1:3] C,H,Ph2, C.H.Ph.CH,Br from C_eH₄PhMe, and from (Schmidt a. Schultz, A. 203, 132; Adam, A. Ch. [C] 15, 243). It is also a product of the fusion of benzoic acid with potash (Barth a. Schreder, M. 3, 808). Leaflets (from alcohol), sl. sol. water. Yields isophthalic acid on oxidation .- $NaA'_2 2aq.$ $-CaA'_2 3aq.$ $-BaA'_2 3\frac{1}{2}aq:$ needles. -BaA', 4 aq.

Ethyl ether Eth. (above 360°). Oil. Diphenyl p-carboxylic acid C.H.C.H.CO.H. [219°]. Formed by saponification of its nitrile, which is got by heating potassium diphenyl sulphonate with KCy in a current of dry CO2 (Doebner, A. 172, 109). Formed also by oxidation of [1:4] C.H.Ph., or [1:4] C.H.PhMe (Schultz, A. 174, 213; Carnelley, C. J. 37, 713). It is also a product of the fusion of benzoic acid with potash. Tufts of needles (from alcohol), v. sl. sol. hot water. May be sublimed. Yields

terephthalic acid on oxidation—BaA'₂.—CaA'₂. Ethyl ether EtA'. [46°]. Prisms. Nitrile C₁₂H₂.CN. [85°]. Crystals.

Anilide C₁₂H₂.CO.NHPh. [224°]. Got by heating C₁₂H₂.C(NOH).C₅H₅ with HOAc, Ac₂O, and dry HCl at 100° (Koller, M. 12, 503).

Diphenyl di-o-carboxylic acid C, H, O, i.e. [2:1]CO,H.C,H,.C,H,.CO,H[1:2]. Diphenic acid. Mol. w. 242. [228°]. Formed by the oxidation of phenanthraquinone or phenanthrene (Fittig a. Östermayer, A. 166, 361; Schmitz, A. 193, 116). It is also produced when phenanthraquinone is boiled with conc. alcoholic potash Anschütz a. Schultz, A. 196, 50; 203, 97). Plates or prisms, m. sol. hot water. May be sublimed. By exhaustive chlorination with SbCl, it yields perchloro-diphenyl, together with a small quantity of perchloro-benzene (Merz a. Weith, B. 16, 2872). Distillation over with red-hot CaO yields diphenylene ketone; distilla-

tion with Ca(OH)₂ gives diphenyl.

Salts.—BaA"4aq: v.sol. water.—CaA" 2½aq.

—MgA"4aq.—Ag₂A": bulky white pp.

Methyl ethers Me₂A". [73:5°].—MeHA". [110°]. Plates, m. sol. water. Got by boiling the anhydride with MeOH.

Ethyl ethers Et₂A". [42°]. Crystals, insol. Aq (Hummel, A. 193, 128).—EtHA". [88°].

Anhydride C.H.CO >0. [218°].

Formed by boiling the acid with AcCl or Ac₂O (Anschütz, B. 10, 1884; 13, 1302: Graebe, B. 20, 848; A. 247, 261). Needles, insol. water, sol. alcohol. Decomposed by heat into CO₂ and diphenylene ketone. H₂SO₄ at 100° forms diphenylene ketone carboxylic acid [217°]. Phenyl-hydrazine forms the crystalline compound CO₂H.C₆H₄.C₆H₄.CO.N₂H₂Ph [174° cor.] which at 250° yields C_6 H₄.CO>N.NHPh [150°]. Benzene and AlCl, form an acid [148°].

Chloride C₁₂H_s(COCl)₂. [98°]. Formed from the anhydride and PCl₃. Yields di-oxy-Formed

phenanthrene on reduction.

[212°]. Formed Amide C12H8(CO.NH2)2. by warming the imide with conc. NH, Aq.

Amic acid C₁₂H_s(CO₂H)(CONH₂). [193°]. Got by boiling the anhydride with NH₂Aq. Plates (from alcohol). Converted by heat into the imide.

Imide C.H. CO NH. [220°]. Formed as above, and also by the action of conc. HClAq on the mono-oxim of phenanthraquinone (Wegerhoff, A. 252, 18). Needles (from alcohol), sl. sol. hot water. Yields an acetyl derivative [92°], and forms $C_{14}H_8O_2(NNa)$, $C_{14}H_8O_2(NAg)$, and C14H,O2(NMe).

Diphenyl om-dicarboxylic acid [2:1] CO.H.C.H..C.H..CO.H[1:3]. Isodiphenic acid. [216°]. Formed by fusing diphenylene ketone carboxylic acid with potash (Fittig a. Isodiphenic Lieppmann, B. 12, 163; A. 193, 155; 200, 9). Needles (from hot water). Yields diphenylene ketone on heating with lime. Gives isophthalic acid on oxidation with CrO₃.—BaA"6aq. Crystalline.—CaA"2aq.—Ag₂A": pp., sl. sol. hot water.

Methyl ether Me₂A". [69.5°].

Ethyl ether Et₂A". Thick pil.

Diphenyl op-di-carboxylic acid [2:1] CO₂H.C₆H₄.C₆H₄.CO₂H[1:4]. [252°]. Formed by saponification of its nitrile [158°] which is got by Sandmeyer's reaction from opdi-amido diphenyl (Reuland, B. 22, 3018).

Diphenyl di-m-carboxylic acid. [above 340°]. Formed by the action of nitrous acid on di-pamido-diphenyl dicarboxylic acid (Griess, B. Small needles, almost insol. cold water.—BaA" 3 aq: crystalline aggregates.

Diphenyl di-p-carboxylic acid [4:1] CO₂H.C₆H₄.C₆H₄.CO₂H [1:4]. Formed by saponification of its nitrile, and also by oxidation of p-ditolyl (Doebner, B. 9, 129, 272; A. 172, 109). Amorphous powder, nearly insol. most solvents.—CaA": insoluble pp.—BaA".— Ag_2A'' : white pp.

Ethyl ether Et.A". [112°]. Prisms. Nitrile C₁₂H_s(CN)₂. [234°]. Formed by heating potassium diphenyl disulphonate with KCy. Needles (from alcohol).

Diphenyl dicarboxylic acid

 $C_eH_s.\hat{O}_eH_s(CO_2H)_2$ [1:3:5]. [c. 310°]. Formed by heating benzoic aldehyde with $CH_s.CO.CO_2H$ and baryta-water (Doebner, B. 23, 2381; 1750). Crystals, v. sl. sol. cold HOAc. Yields diphenyl on distillation with lime.—BaA"4 laq: needles.—CaA".—CuA"

Diphenyl tricarboxylic acid

 $\mathbf{C}_{15}\mathbf{H}_{10}\mathbf{O}_{6}$ i.e. $\mathbf{CO}_{2}\mathbf{H}.\mathbf{C}_{6}\mathbf{H}_{4}.\mathbf{C}_{6}\mathbf{H}_{3}(\mathbf{CO}_{2}\mathbf{H})_{2}$. Formed by fusing diphenylene ketone dicarboxylic acid with KOH (Bamberger a. Hooker, B. 18, 1035; A. 229, 160). White crystalline powder, v. sol. alcohol and ether, sl. sol. hot water.—Pb,A"'2: insoluble crystalline pp.—Ag₃A", white pp.

References.—Amido-, Bromo-, Iodo-, Nitro-,

and Oxy- DIPHENIC and DIPHENYL CARBOXYLIC

ACIDS

PHENYL-CETYL-AMINE v. CETYL-ANILINE. PHENYL-CHELIDAMIC ACID v. CHELI-DONIC ACID.

PHENYL-CHLORO-ACETIC ACID v. CHLORO-PHENYL-ACETIC ACID

PHENYL CHLORO-BENZYL KETONE

 C_6H_5 .CO.CHCl.C_pH₅. [65°]. (Curtius, J. pr. [2]) 44, 547). Formed by reducing phenyl dichlorobenzyl ketone with iron and acetic acid (Lachovitch, B. 17, 1163). V. sol. alcohol.

Phenyl di-chloro-benzyl ketone C₆H₅.CO.CCl₂.C₆H₅. [71°]. (Zinin, A. 119, 177); [61°] (L.). Formed from benzil and PCl. Prisms (from ether). Reduced by zinc and HCl

to phenyl benzyl ketone.

PHENYL CHLORO-BENZYL SULPHONE C.H., CO.CHCl.C.H.. Formed by heating benzylidene chloride with C.H., SO.Na and alcohol at 150° (Otto, *J. pr.* [2] 40, 517). Needles (from HOAc), sl. sol. hot alcohol.

PHENYL-CHLORO-ETHANE CHLORO-

ETHYL-BENZENE.

CHLORO-ETHYL PHENYL OXIDE

Chloro-cthyl derivative of PHENOL.

PHENYL CHLORO-ETHYL SULPHONE C.H., SO₂CH₂CH₂Cl. [56°]. Formed by the action of PCl, or HCl on phenyl oxy-ethyl sulphone (Otto, J. pr. [2] 30, 197). Six-sided tablets (from benzene), sl. sol. water.

Reactions.-1. Boiling with alcohol and Ag2O converts it into C₆H₅SO₂C₂H₄OH. -2. By heating with C₆H₅SO₂Na in alcoholic solution it is converted into (C.H., SO₂)₂C₂H₄.—3. Sodium-analgam reduces it, in alcoholic solution, to benzene sulphinic acid.-4. Alcoholic NaOEt forms C.H.SO.C.H.OEt. - 5. Converted by heating in sealed tubes with dry Ag.O and benzene into (C.H. SO.C.H.), O.-6. Ammonia

forms $(C_0H_0.SO_2.C_2H_4)_2NH.$ —7. Alcoholic KHS produces $(C_0H_0.SO_2.C_2H_4)_2S.$ [124°].

Phenyl chloro-ethyl sulphone

C.H.SO.CHCI.CH. [52°]. Formed by heating aqueous CH.CCl.CO.Na (1 mol.) with sodium benzene sulphinate (2 mols.) at 100° (Otto, J. pr. [2] 40, 532). Crystals (from alcohol). PHENYL-CHLORO-IMESATIN v. ISATIN.

PHENYL CHLORO-METHENYL DI-ETHYL TRI-SULPHONE CCl(SO₂Ph)(SO₂Et)₂. [130°]. Got by chlorinating CH(SO₂Ph)(SO₂Et)₂ (Laves, B. 25, 863). Plates, sl. sol. hot water.

TRI-PHENYL CHLORO-METHENYL SULPHONE CCI(SO,Ph),. [260°]. Got by chlorinating CH(SO,Ph), (Laves, B. 25, 350). PHENYL CHLORO-METHYL KETONE v.

ω-Chloro-acetophenone.

PHENYL CHLORO-METHYL SULPHONE

C_eH_a·SO₂·CH_aCl. [53°]. Formed by boiling an aqueous solution of C_eH_aSO₂Na (2 mols.) with CHCl₂·CO₂Na (1 mol.), the yield being nearly the theoretical quantity (Otto, J. pr. [2] 40, 527; B. 21, 656). Formed also from C.H. SO, Na and CH₂Cl₂.

Phenyl di-chloro-methyl surphone

 $C_aH_b.SO_c.CHCl_2$. [59°]. Formed by treatment of $C_aH_b.SO_c.CH_c.CO_c.H$ with Cl in diffused daylight (Otto, J. pr. [2] 40, 541). Monoclinic prismatic tables, v. sol. hot alcohol.

PHENYL-CHLORO-NAPHTHYL-AMINE.

Benzoyl derivative C_eH₂NBz.C₁₆H₆Cl. [152°]. Formed from NPhBz.C₁₆H, and PCl, (Claus a. Richter, B. 17, 1590). Needles, sol. alcohol.

PHENYL-CHLORO-NITRO-PHENYL-HY-**DRAZINE** $C_6H_3.N_2H_2.C_6H_3Cl(NO_2)$ [1:4:6]. [140°]. Formed from phenyl-hydrazine and C,H,Cl(NO,), in the cold (Willgerodt, J. pr. [2] 37, 355). Red prisms (from alcohol and ether).

PHENYL TRI - CHLORO - OXY - PROPYL KETONE C, H, COCH, CH(OH). CH, [66°] and [77°]. Got from acetophenone, chloral, and HOAc (Koenigs, B. 25, 795). V. sl. sol. water. Converted by H2SO4 into C6H2.CO.CH:CH.CCl2 $[102^{\circ}$

PHENYL CHLORO-PHENYL HYDRAZINE C_aH_a , N_zH_z , C_gH_aCl [1:4]. [90°]. Formed by reducing C,H,N2.C,H,Cl with alcoholic ammonium sulphide (Heumann a. Mentha, B. 19, 1688)

PHENYL CHLORO-PHENYL KETONE v.

CHLORO-BENZOPHENONE.

PHENYL-CHLORO-PROPIONIC ACID CHLORO-PHENYL-PROPIONIC ACID.

PHENYL ω-CHLORO-p-TOLYL KETONE C,H,CO.C,H,CH,Cl. Benzoylbenzyl chloride. [98°]. Formed by chlorination of phenyl p-tolyl ketone at 110° (Thörner, A. 189, 89). Silky needles (from dilute alcohol). May be sublimed. On further chlorination at 135° it yields phenyl dichlorotolyl ketone or benzoyl benzylidene chloride C₈H₅.CO.C₈H₄.CHCl₂ [95°] and, at 155°, phenyl tri-chloro-tolyl ketone or benzoyl-benzotrichloride C₆H₄.CO.C₆H₄CCl₂ [111°], which is converted by PCl₂ into C₆H₄.CCl₂.C₆H₄.CCl₂ [80°] crystallising in thin plates.
PHENYL-CHRYSYL-THIO-UREA

C_oH_oNH.CS.NH.C₁₀H₁₁. [186°]. Formed from phenyl-thiocarbimide and chrysylamine and from chrysyl-thiocarbimide and aniline in benzene solution (Abegg, B. 24, 957). Crystals.

PHENYL-CINCHONIC ACID v. PHENYL-

QUINOLINE CARBOXYLIC ACID.

PHENYL-CINCHONIDINE C₁₉H₂₁PhN₂O. { Formed in two modifications (an oil, sol. ether, and an amorphous powder, insol. ether) by heating cinchonidine with aniline (Claus a. Bätcke,

B. 13, 2194). Both give B"H,PtCl_e 2aq.
PHENYL-CINNAMIC ACID C₁₈H₁₂O₂ i.e. C₃H₃.CH:CPh.CO₂H. [170°]. Formed by saponi-ying the nitrile or heating sodium phenylacetate with benzoic aldehyde and Ac2O at 150° (Oglialoro, G. 9, 429). Needles, sol. alcohol and ether, v. sl. sol. cold water. Reduced by sodiumamalgam to a B-di-phenyl-propionic acid.-BaA'24aq: micaceous plates.

Methyl ether MeA'. [78°] (Cabella, G.

14, 114). Needles (from dilute alcohol).

Nitrile C, H, CH: CPh.CN. Phenyl-styryl cyanide. [87°]. (360° cor.). Formed by the action of benzyl cyanide on benzoic aldehyde or benzylidene chloride in presence of NaOEt or solid NaOH (Rossolymo, B. 22, 1235; Meyer, 250, 124; Janssen, A. 250, 129; Neure, A.
 250, 155; Frost, A. 250, 157). White plates (from alcohol). Yields CHPhBr.CPhBr.CN [130°] and CHPhCl.CPhCl.CN. [168°].

o-Carboxylic acid. Imide

C₆H₄<CO—NH $\stackrel{\sim}{-}\stackrel{\sim}{\mathrm{NH}}$. [174°]. Formed by heating the imide of carboxy-phenyl-acetic acid (vol. i. p. 706) with benzoic aldehyde (Gabriel, B. 20, 1205). Needles.

References .- NITRO- and OXY- PHENYL-CIN-

NAMIC ACID.

PHENYL-CINNAMYL-HYDRAZINE

CHPh:CH.CH₂.NPh.NH₂. [54°]. Formed from sodium phenyl-hydrazine and cinnamyl bromide (Michaelis, B. 22, 2233). Crystals. PHENYL-CINNAMYL KETONE v. PHENYL

STYRYL KETONE.

PHENYL-p-COUMARIC ACID [1:4]

C₈H₄(OH).CH: CH.CO₂H. [219°]. Formed by heating p-oxy-benzoic aldehyde with Ac.O and sodium phenyl-acetate and boiling the resulting acetyl derivative with baryta-water (Oglialoro, G. 13, 173).—AgA': nearly insol. water.

Acety'l derivative. [170° Methyl ether MeA'. [170 [170°]. Ethyl ether EtA'. [152°].

Methyl derivative. [189°]. Vol.iii.p.738. PHENYL-COUMARIN [140°]. C15H10O2. Formed, together with C.H.(OAc).CH:CPh.CO2H, heating salicylic aldehyde with sodium pnenyl-acetate and Ac2O for 8 hours at 150° (Oglialoro, G. 9, 428). Prisms, sol. ether.

Mono-sulphonic acid C15HO2(SO2H) 2 aq. White needles (Curatolo, G. 14, 257). -BaA'2. - PbA'2 4aq: needles, m. sol. hot water.

Di-sulphonic acid C13H8O2(SO3H)2 6aq. [89°]. Deliquescent crystals.—BaA'₂ 4aq: white prisms.—PbA'₂ 5aq: needles, v. sol. water.
PHENYL-CROTONIC ACID C₁₆H₁₀O₂ i.e.

C.H. CH: CMe.CO.H. Phenyl-methacrylic acid. (a)-Methyl-cinnamic acid. [82°]. Formed by heating benzoic aldehyde with propionic anhydride and sodium propionate (Perkin, C. J. 31, 391; 32, 661). Formed also by heating benzoic aldehyde with Ac,O and sodium methyl-malonate; and by heating benzoic aldehyde with sodium propionate and HOAc or Ac.O (Stuart, C. J. 43, 404; Slocum, A. 227,57). Obtained also by oxidation of the corresponding aldehyde (Miller a. Kinkelin, B. 19, 526). It is also a product of the action of Na on benzyl propionate (Conrad a

Hodgkinson, A. 193, 314).

Transparent plates (from alcohol), sl. sol. hot water. When prepared by Perkin's method it appears to be accompanied by an isomeride [74°] crystallising in needles, which is converted into the acid [82°] by frequent recrystallisation (Raikoff, B. 20, 3396). Yields methronol (q.v.) when heated with dilute H₂SO₄. Bromine forms CHPhBr.CMeBr.CO₂H[137°] (Körner, B. 21, 276).

Salts.—BaA'2 aq: very minute plates. BaA'2 4aq: small white crystals, got by slow

cooling.—AgA'. S. 374. Needles.

Methylether MeA'. [39°]. (254°). Amide [128°] (Edeleano, B. 20, 619).

Phenyl-isocrotonic acid

C₈H₃.CH:CH.CH₂.CO₂H. [86°] (J.); [88°] (B. a. D.). (302°). Formed by heating benzoic aldehyde with succinic anhydride and sodium succinate (Perkin), the yield being 2 p.c. (Jayne, A. 216, 100). Formed also by heating phenyltrimethylene tricarboxylic acid C,H2Ph(CO2H) in a current of CO2 at 190° (Buchner a. Dessauer, B. 25, 1155). Prepared by heating benzoic aldehyde with sodium succinate and Ac₂O at 125°, and separated from phenyl-paraconic acid by solution in CS2. Long thin needles (from water) or prisms (from CS₂). When heated with H₂SO₄ (4 pts.) and water (4 pts.) it changes to the isomeric phenyl-butyrolactone and an acid [179°] (Erdmann, A. 227, 257). Yields γ-phenyl-n-butyric acid [49°] on reduction. Boiling Boiling converts it into NaOHAq the isomeric C₆H₅.CH₂.CH:CH.CO₂H [65°]. Split up by long boiling into water and (a)-naphthol. Alkaline KMnO, at 0° forms di-oxy-phenyl-butyric lactone (Fittig a. Obermüller, A. 268, 44).—BaA', 3aq.— CaA' 2 3aq. - AgA': bulky pp

a-Phenyl-crotonic acid CH, CH:CPh.CO, H. Methylatropic acid. [135°]. Formed by heating sodium phenyl-acetate with paraldehyde and Ac₂O (Oglialoro, G. 15, 514). Minute prisms (from water), sol. alcohol and ether .-AgA': m. sol. hot water.

Phenyl-crotonic acid

 C_0H_3 .CH:C(CH₂Ph).CO₂H. [158°]. Formed from sodium phenyl-propionate, benzoic aldehyde, and Ac₂O at 160° (Oglialoro, G. 20, 162). Needles (from alcohol).

Reference .- OXY-PHENYL-CROTONIC ACID.

PHENYL-CROTONIC ALDEHYDE v. PHENYL-METHACRYLIC ALDEHYDE.

PHENYL-CUMAZONIC ACID

i.e. $C_0H_3(CO_2H) < CMe_2.O (?)$. C₁₇H₁₅NO₃ [220°]. Formed by heating (3,4,1)-amido-oxypropyl-benzoic acid with BzCl at 100°-120° (Widmann, B. 16, 2585). White crystals. Sol. alcohol, insol. water.—HA'H2SO, 2aq: thin plates

PHENYL-CUMINYL-AMINE C.H,,N C.H., NH.CH., C.H.Pr. [41°]. Formed by reducing C.H., N:CH.C.H.Pr (Uebel, A. 245, 289). Plates or prisms. Yields a nitrosamine C₁₀H₁₀(NO)N [94°].—B'HCl: white crystals. PHENYL-CUMINYL-THIO-UREA

[100°-106°]. C.H.Pr.CH.NH.CS.NHPh. Formed from cuminylamine and phenyl-thiocarbimide (Goldschmidt a. Gessner, B. 20, 2416).

PHENYL-CUMINYL-UREA C₁, H₂₀N₂O i.e. C₄H₂NH₂CO.NHC₁₀H₁₂. [144°]. Formed from cuminyl cyanate and aniline (Basb, B. 8, 1151).

and from phenyl cyanate and cuminylamine (Goldschmidt, B. 20, 2415). Small needles.

PHENYL-DI-CUMYL-GUANIDINE

 $NPh:C(NH.C_6H_4(C_8H_7)[1:4])_2$. Formed from dicumyl-thio-urea and aniline (Francksen, B. 17, 1226). Resin.

PHENYL 4-CUMYL KETONE C16H16O i.e. C₃H₁,CO,C₆H₂Me₃ [1:2:4:5]. (329°). From B2Cl, ψ-cumene, CS₂, and AlCl₃ (Elbs, J. pr. [2] 35, 491; cf. Claus, B. 19, 2881); the yield is

65 p.c. of the calculated.

Reactions.-1. Boiling produces small quantities of needles [180°], probably (B. 2,3)-dimethyl-anthraquinone.—2. Conc. or fuming H₂SO₄ at 100° forms benzoic acid and ψ-cumene sulphonic acid $C_8H_2Me_8(SO_3H)$ [1:2:4:5].—3. Fuming HNO₃ mixed with H₂S₂O₇ gives two isomeric tri-nitro-derivatives; one [185°] is v. sl. sol. alcohol, the other [155°] is m. sol. alcohol .- 4. Dilute nitric acid forms benzoyltrimellitic acid, C.H. CO.C.H. (CO.H) C₈H₂.CO.C₈H₂Me(CO₂H)₂. Phenyl cumyl ketone

C₆H₅.CO.C₆H₄Pr. (343°) at 738 mm. Formed from cuminyl chloride, benzene, and AlCl₃ (Smith). Yields two oxims [106°] and [132°], the latter of which

forms an acetyl derivative [90°].

Phenyl n-cumyl ketone C₆H₄.CO.C₆H₄Pr. (345° at 716 mm.). Formed from n-cumene, BzCl, and AlCl₂ (Smith, B. 24, 4033). Prisms. Yields two oxims [130°] and [104°], which yield acetyl derivatives, melting at 116° and 66°

PHENYL-y-CUMYL-METHANE CARBOXY-LIC ACID [1:2:4] C₆H₂Me₃.OH₂.C₆H₄.CO₂H [1:2] [186°]. Got by reducing C₆H₂Me.CO.C₆H₄.CO₂H

with Zn and NH₂Aq (Gresly, A. 234, 238). DI-PHENYL-\(\psi\)-CUMYL-METAPYRAZOLE

TETRAHYDRIDE C21H26N2 i.e.

 $C_9H_{11}.CH < NPh > C_2H_4$. [125°]. Formed from p-cuminic aldehyde and di-phenyl-ethylene-diamine at 100° (Moos, B. 20, 733). Silky needles.

PHENYL-4-CUMYL-UREA C16H18N2O i.e. NHPh.CO.NHC₆H₂Me₈ [211°]. Formed from ψ-cumidine and phenyl cyanate (Goldschmidt a. Bardach, B. 25, 1861). White needles.

PHENYL-CYANAMIDE v. vol. ii. p. 316.
Di-phenyl-cyanamide C₁₈H₁₀N₂ i.e. C(NPh)₂. Carbodiphenylimide. Mol. w. 194. (331° cor.). Formed by warming a solution of s-di-phenylthio-urea in benzene with HgO (Weith, B. 7, 10, 1303; 9, 810). Syrup, drying up to a vitreous mass. Changes spontaneously into an opaque polymeride [170°] resembling porcelain.

Reactions.-1. Boiling dilute alcohol converts it into s-di-phenyl-urea. —2. Aniline yields (a)-tri-phenyl-guanidine.—3. H₂S passed into its solution in benzene forms $CS(NHPh)_z$ -4. s-Di-phenyl-thio-urea at 150° forms phenyl-thiocarbimide and (a)-tri-phenyl-guanidine. — 5. Diphenyl-urea at 150° acts in like manner, forming phenyl cyanate and tri-phenyl-guanidine.—6. Phenyl-hydrazine at 120° forms the compound C₄H₃N₂:C(NHPh)₂ [204°] which gives the salts B'HC!, B'₂H,PtCl₈, and B'H₂SO₄, and is converted, by heating with phenyl-hydrazine at 185°, into $C_{32}H_{28}N_6$ [200°], which gives the salts B'_3H_1 , C_{4} and B'_42H_2 PtCl₅. The compound PhN₂:C(NHPh)₂ is converted by heating with phenyl-thiocarbimide at 190° into $O_{28}H_{23}N_3$ S [175°], and by heating with di-p-tolyl-cyanamide

into $C_{24}H_{22}N_s$ [128°], which gives the salts $B'_{2}H_{4}Cl_{4}$ and $B'_{2}2H_{2}PtCl_{5}$ (Wessel, B. 21, 2272). 7. HCy passed into its solution in benzene forms C14H11Na crystallising from alcohol in prisms [137°]. A little of its solution in H2SO, mixed with water is coloured deep blue by NaOH (Laubenheimer, B. 13, 2155).—8. Tolylene-o-diamine at 135° forms C₂₀H₂₀N₄ [161°] which yields B'2H3Cl, [174°] (Dahm a. Gasiorowski, B. 19, 3057).

Salt.—B'HCl. Needles (from benzene).

Tri-phenyl-di-cyan-di-amide $C_{20}H_{18}N$ Formed from NHPh.CS.NH.C(NPh).NHPh, alcohol, and HgO, and also by heating di-phenylguanidine at 175° (Rathke a. Oppenheim, B. 23, 1673). Yellow needles (containing EtOH) (from alcohol).-B'HCl. Needles, insol. water. $-B'_{2}H_{2}PtCl_{6}2aq.-B'_{2}H_{2}SO_{4}2aq.$

Reference .- OXY-PHENYL-CYANAMIDE. PHENYL CYANATE v. vol. ii. p. 315. PHENYL-CYANIC ACID v. CYANIC ACID. PHENYL CYANIDE v. BENZONITRILE and PHENYL-CARBAMINE.

Di-phenyl dicyanide C11H10N2 i.e.

PhC $\langle N \rangle$ CPh? [152°]. Formed from benzamidine and benzoic aldehyde (Pinner, B. 22, 1610; 23, 2925). Prisms. Changes at 260°-270° into cyaphenine.

PHENYL-CYANURIC ACID v. Cyanuric

acid in the article on CYANIC ACID PHENYL-p-CYMYL-CARBINOL

 $C_6H_3.CH(OH).C_6H_3(CH_3)(C_3H_7)$ [1:2:5]. (327° uncor.). Formed by reduction of phenyl p-cymyl ketone (Claus a. Elbs, B. 18, 1798; f. pr. [2] 35, 497). Thick colourless oil.

PHENYL p-CYMYL KETONE C₁₇H₁₈O i.e.

 C_6H_8 .CO. C_6H_3 MePr [1:2:5]. (224° at 40 mm.);

(340°) at 760 mm.

Formation.—1. From cymene by heating with benzoic acid and P2O3 (Kollarits a. Merz, B. 6, 546).—2. From cymene, BzCl, and zine (Grucarevic a. Merz, B. 6, 546, 1246).—3. From cymene, BzCl, CS2, and AlCl2 (Elbs, J. pr. [2] 35, 494].

Properties.—Yellowish oil, v. sol. alcohol and ether, m. sol. HOAc. Very slightly volatile

with steam.

Reactions .- 1. Bromine and iron powder at 140° give benzoic acid and bromo-cymene, C.H.BrMePr [1:2:5] [232°]. - 2. Conc. H.SO. and KNO, give an amorphous di-nitro-deri tive.—4. Dilute HNO, gives benzoyl-terephthalic acid.—5. Concentrated H₂SO₄ at 100° gives benzoic acid and cymene sulphonic acid, $C_6H_3MePr(SO_3H)$ [1:4:2].

PHENYL-CYSTEINE C.H., NSO, i.e. Formed by boiling CH_2 . $C(SPh)(NH_2)$. CO_2H . phenyl-mercapturic acid with dilute H2SO4 (Baumann a. Preusse, H. 5, 337). Plates (from NH.Aq) or six-sided tables (from water), v. sol. acids and alkalis, sl. sol. cold water. phenyl mercaptan on boiling with NaOHAq.

Reference.—Bromo-Phenyl-Cysteine. PHENYL-DECOIC ACID. Nitrile Formed from $C_sH_s.CH(C_sH_1,).CN.$ (328°). benzyl cyanide, octyl iodide, and NaOH (Rossolymo, B. 22, 1237). Yellow oil.

PHENYL-DURYL-CARBINOL [6:4:3:2:1] $C_0HMe_1.CH(OH).C_0H_0$. (above 360°). Formed by reduction of phenyl iso-duryl ketone (Essner a. Gossin, Bl. [2] 42, 172). Liquid. Yields an acetyl derivative boiling above 360° and a benzoyl derivative [75°].

DI - PHENYL DURYLENE DIKETONE C₂₁H₂₂O₂ i.e. (C₆H₃.CO)₂C₄Me₄. [270°]. (above 380°). Formed from durene, BzCl, and AlCl₃ (Friedel a. Crafts, A. Ch. [6] 1, 512). Prisms.

PHENYL DURYL KETONE C17H18O i.e. C.H. CO.C. HMe. [119°]. (343°) at 725 mm. Got from durene, BzCl, and AlCl, (Friedel a. Crafts, A. Ch. [6] 1,511). Acicular prisms, v. sol. warm alcohol. Reduced by HI and P to C₁₇H₂₀ [60.5°] (310° at 716 mm.). Br forms C₁₇H₁₂Br₂O

[225°] and other products.

Phenyl isoduryl ketone. [63°]. Formed by the action of BzCl and AlCl₃ on the isodurene derived from the isodurene derived from toluene (Essner a. Gos-in, Bl. [2] 42, 170). Successive treatment with HCy and alcoholic potash gives the acid C₆HMe₄.CPh(OH).CO₂H, sol. water and alcohol.

PHENYLENE. The radicle C.H.. PHENYLENE-ACETAMIDINE

 $C_eH_4 < NH > C.CH_e$. [170°] (H.); [175°] (L.). Formed by boiling phenylene-o-diamine with HOAc, or by reducing C₆H₄(NO₂)(NHAc) with tin and HCl (Ladenburg, B. 8, 677; Hübner, A. 209, 353). Needles.—B'HCl.—B'₂H₂PtCl₆ aq.— B'HNO3.-B'H2SO4.

Phenylene - di - acetamidine $C_{10}H_{14}N_{4}$ [1:4] C₆H₄(CH₂.C(NH).NH₂)₂. [182°]. Formed by the action of alcoholic NH₃, followed by NaOHAq, upon the hydrochloride of phenylenedi-acetimido-ether (Glock, B. 21, 2660). Plates.

o-PHENYLENE-DI-ACETIC ACID C,0H,0O, i.e. [1:2] C₆H₄(CH₂.CO₂H)₂. Mol. w. 194. [150°]. Formed by saponification of its nitrile by boiling with dilute H₂SO₄ (Baeyer a. Pape, B. 17, 447). Slender needles, v. sol. alcohol, m.

sol. cold water.—Ag₂A": insoluble pp.

Nitrile C₈H₄(CH₂.CN)₂. [60°]. Formed by the action of KCy on di-ω-bromo-o-xylene.

m-Phenylene-di-acetic acid C₆H₄(CH₂.CO₂H)₂. [170°]. Formed by boiling its nitrile with alcoholic potash (Kipping, C. J. 53, 42; B. 21, 42).

Needles, v. sol. water.—Ag₂A": white pp.

Nitrile C₆H₁(CH₂CN)₂. [29°]. (308° a
300 mm.). From [1:3] C₆H₁(CH₂Br)₂ and KCy. (308° at p-Phenylene-di-acetic acid C.H. (CH2.CO2H)2 [241°]. Got from its nitrile (Biedermann, B. 5, 762, Klippert, B. 9, 1766; Kipping, C. J. 53, 705, Rhppert, B. 5, 1705, Rhping, C. 3. 5.
44). Needles, m. sol. water.—CaA"2aq.—CaA"3aq.—BaA"2½aq: needles, v. sol. water.—ZnA".—CuA".—Ag₂A": white amorphous pp.

Ethers.—MoA". [57°].—Et₂A". [58°].

Chloride. Öil (Klippert).

Amide. [above 290°]. Plates and needles. Nitrile C₆H₄(CH₂.CN)₂. [96°]. Formed Formed by the action of KCy on C₈H₄(CH₂Br)₂ or C₈H₄(CH₂Cl)₂. Three-sided prisms (from ether). Alcoholic ammonium sulphide at 100° converts it into C₈H₄(CH₂·CS.NH₂)₂ [206°] which crystallises from HOAc.

Di-phenylene-acetic acid C.H. CO.H. [222°]. Formed by heating C₁₂H₈:C(OH).CO₂H with HI and P at 140° (Friedländer, B. 10, 536). Small crystals (from alcohol). Decomposed by heating with sods-lime into CO, and fluorene. AgA': unstable pp.

Ethyl ether EtA'. [165°].

Reference .- OXY-DIPHENYLENE-ACETIC ACID. p-PHENYLENE DI-ACETIMIDO-ETHER

C₆H₄(CH₂.C(NH).OEt)₂. The hydrochloride B"2HCl [above 240°] formed by the action of alcohol and HCl on C₅H₄(CH₂.CN)₂, crystallises in needles (Glock, B. 21, 2660). It is insol. ether. o-PHENYLENE-DI-ACRYLIC ACID

C_sH_s(CH:CH.CO₂H)₂. [above 300°]. Formed by boiling C_sH_s(CH₂.CCl(CO₂Et)₂)₂ with alcoholic by both (Parkin CH₂.CCl (CO₂Et)₂)₂ and (25) potash (Perkin, C. J. 53, 14; B. 19, 435). Sl. sol. water, v. sl. sol. alcohol. - Ag, A": pp.

p-Phenylene di-acrylic acid C.H.(CH:CH.CO2H)2. [310°]. Obtained saponifying the mono-ethyl ether HEtA" [200°] which is got by heating p-aldehydo-cinnamic ether with NaOAc and Ac.Ō (Löw, A. 231, 377). Got also by boiling C₆H₄(CH₂.CBr(CO₂Et)₂), with alcoholic potash (Kipping, C. J. 53, 41). Yields a tetrabromide crystallising in needles. - Ag2A"

PHENYLENE - DI - ALLYL - DI - THIO - DI -UREA C₆H₄(NH.CS.NHC₂H₅)₂. The o- [159°], m- [105°], and p- [200°] compounds are formed by heating the corresponding phenylene-diamines with alcohol and allyl-thiocarbimide (oil of mustard) at 100° (Lellmann a. Würthner, A. 228, 201). The o- and p- compounds decom-

pose when heated. They are all sol. HOAc.
o-PHENYLENE-DI-AMIDO-DI-ACETIC ACID [1:2] $C_0H_4(NH.CH_2.CO_2H)_2$. The ether $\mathrm{Et_2A''}$ [135°] is got from o-phenylene diamine and ClCO₂Et (Zimmermann a. Knyrim, B. 16, 515).

m-Phenylene-di-amido-di-acetic acid m-Franylene-ul-aminu-ul-action acide [1:3] C_sH₄(NH.CH₂.CO₂H)₂. The ethyl ether Et₄A" [73°] is got by heating m-phenylene-diamine with CH₂Cl.CO₂Et (Zimmermann B. 15, 518; 16, 514). It is converted by conc. HClAq into crystalline H₂A"H₂Cl₂, v. sol. water.

p-Phenylene-di-amido-di-acetic acid. ethyl ether Et₂A" [83°] and the hydrochloride H₂A''H₂Cl₂ are formed in like manner from p-phenylene-diamine.

o-PHENYLENE-DIAMINE [1:2] C.H.(NH.)2. Mol. w. 108. [104°]. (252°). Prepared by reducing o-nitro-aniline with tin and HCl (Zincke a. Sintenis, B. 6, 123; Vignon, Bl. [8] 2, 675; Sandmeyer, B. 19, 2654). Formed also by heating the corresponding di-amido-benzoic acids with baryta (Griess, J. pr. [2] 3, 143; Salkowski, A. 173, 58) and by the action of tin and HCl on bromo-o-nitro-aniline (Wurster, A. 171, 63; Hübner, A. 209, 360) and on o-di-nitro-benzene (Zincke, B. 7, 1374). Plates (from water) or tables (from chloroform), v. e. sol. hot water, alcohol, and ether.

Reactions.—1. FeCl, added to a solution of the hydrochloride forms di-amido-phenazine hydrochloride, which crystallises in ruby-red needles C12H10N4HCl (Fischer, B. 22, 355) .- 2. Pyrocatechin at 200° forms phenazine C.H.: N.: C.H. 3. (β)-Naphthoquinone forms naphthophenazine C₁₀H₄:N₂:C₈H₄.—4. Cyanogen gas forms C₈H₈N₄ a yellow crystalline powder [c. 280°], al. sol. water, forming B'₂H₂PtCl₈3aq and B'₂H₂PtCl₈aq. By heating with HCl it is converted into C.F.N.O [280°] and di-oxy-quinoxaline (Black, Bl. [2] 42, 104; B. 18, 672).-5. The sulphocyanide is converted by heat into a mixture of phenylenethio-urea and thio-urea (Lellmann, A. 221, 8) .-6. KNO2 added to a very dilute solution of the sulphate forms 'amidoazophenylene' C.H.N.

which crystallises from benzene in pearly white needles [98.5°] (Ladenburg, B. 9, 219).—7. ICy forms C₁,H₁,N₄, which yields the salts B"H₂Cl₂, $B''HCl 2\frac{1}{2}aq$, $B''2HNO_2 2\frac{1}{2}aq$, and $B''H_2SO_4 2\frac{1}{2}aq$, and the benzoyl derivative C₁₃H₁₁BzN₄, and is converted by nitrous acid into C₁₅H₈N₆O₂ (Hübner, B. 9, 778; 10, 1716).—8. The hydrochloride is converted by benzoic aldehyde into dibenzylidene-phenylene diamine (C₆H₄(N:CHPh)₂ [134°], yielding B'_H.PtCl, B'HNO, B'H,SO, B'MeI, and B'EtI [213°] (Ladenburg, B. 11, 1653).

Salts.—B"2HCl. Needles, v. sol. water.—
B"H2PtCl_s.—B"2H2SO₄.—B"H2SO₄1laq: plates.
Di-acetyl derivative C_sH₄(NHAc)₂.

[186°]. Long silky needles, v. sol. hot water

and alcohol (Bistrzycki, B. 23, 1878).

Benzoyl derivative C.H.(NH.)(NHBz). [140°]. Got by reducing C.H.(NO.)(NHBz) with ammonium sulphide (Mixter, Am. 6, 26). Minute crystals, sol. water. When the reduction is effected by tin and HCl the product is benzenyl-phenylene-diamine (q. v.).
Dibenzoyl derivative C.H. (NHBz).

[c. 300°]. Thin prisms (from HOAc) (Bistrzycki).

 $Di-p-toluyl\ derivative$ $C_sH_4(NH.CO.C_sH_4Me)_2$. [228°]. Got, together with tolenyl-benzamidine $C_sH_4:N_2H:C.C_sH_4Me$ [268°], by the action of p-toluyl chloride on ophenylene-diamine dissolved in benzene (Hübner, A. 210, 328). Colourless needles, sl. sol. alcohol.

m-Phenylene - diamine C₈H₄(NH₂)₂. (277° i.V.). Prepared by reduction of m-dinitro-benzene or m-nitro-aniline (Hofmann, Pr. 11, 518; 12, 639; Zincke, B. 5, 792). Formed also by heating s-di-amido-benzoic acid with baryta (Wurster, B. 7, 214; Hübner, A. 222, 86), and by the action of tin and HCl on (4,2,1)-dinitro-benzoic acid (Wurster, B. 7, 149) and on (1,2,4)-bromo-di-nitro-benzene (Z.). Got also by heating resorcin (1 pt.) with ammoniacal CaCl₂ (4 pts. containing 35 p.c. NH₃) for eight hours at 275° (Seyewitz, C. R. 109, 814). Crystalline, m. sol. water, v. sol. alcohol and ether. Its solutions are alkaline in reaction. KNO₂ added to a solution of a salt of m-phenylene-diamine forms a brown pp. containing $C_sH_3(NH_2)_2\cdot N_2\cdot C_sH_1\cdot NH_2$. A solution in dilute H₂SO₄ is coloured yellow by a trace of nitrous acid. Aldehyde added to an alcoholic solution of m-phenylene-diamine hydrochloride forms an orange-red colour, with a green fluorescence. From this liquid NH, Aq ppts. di-methyl-phenanthroline tetrahydride C₁₄H₁₄N₂ (Girard a. Roques, C. R. 107, 1158; Schiff, A. 253, 328). Benzoic aldehyde in alcohol forms with the hydrochloride the base $CHPh(C_0H_4(NH_2)_2)_2$. The hydroide the base CHPh(CeH4(NH2)2)2. chloride of m-phenylene-diamine gives a crimson colour when boiled with oxygenated water (Deniges); in presence of ammonia the liquid becomes blue, but is turned red by NaOHAq. Hydrogen peroxide gives a bluish-green tint (Cazeneuve, Bl. [3] 5, 855) PbO₂ and dilute HOAc give a brown colour (Lauth, C. R. 111, 975). Alkaline NaOBr gives a maroon-red pp. (Deniges, C. R. 107, 662). CS₂ in alcohol at 35° forms yellow,monoclinic prisms of $C_0H_4 < NH > CS$ and several other bodies (Gucci, G. 17, 523, 2657).

ous than the p- isomeride (Dubois a. Vignon, C. R. 107, 533)

Salts.—B"HCl. Needles, v. sol. water, sl. sol. conc. HClAq. — B"2HCl. — B"HPtCl.; needles. — B"H₂Sn₂Cl. (Gerdemann, Z. [2] 1, 51).—B"H₂SnCl.,—B"H₂SO₄.

Hydriodide: Mills, C. J. 17, 153.

By driodide: Mills, C. J. (H. 19) (19)

Di-formyl derivative C.H.(NH.CHO)r [155°]. Small crystals (Tobias, B. 15, 2447).

Mono-acetyl derivative $C_eH_4(NH_2)(NHAc)$. Crystalline solid, v. sol. water (Wallach a. Schulze, B. 15, 3020).—B'HCl. [280°]. [280°].

Di-acetyl C.H.(NHAc)2. derivative [189°] (Kelbe, B. 16, 1200; cf. Barbaglia, B. 7,

Mono-benzoyl derivative C_cH₄(NH₂)(NHBz). [125°] (Bell, B. 7, 498); [260°] (Hübner, A. 208, 298). Got by reducing C_cH₄(NO₂).NHBz.—B"HCl.—B"H₂SO₄. Needles.

Di-benzoyl derivative C,H,(NHBz)2. [240°] (Ruhemann, B. 14, 2651; Hinsberg, A.

254, 255). White plates.

m-Amido-benzoyl derivative C_cH₄(NH₂)(NH.CO.C_cH₁NH₂). [129°]. (from dilute alcohol) (Hugh, B. 7, 1268). Needles

o-Oxy-benzoyl derivative $C_6H_4(NH_2).NH.CO.C_6H_4.OH.$ [143°]. (Bell, C. J. 28, 1201).

p-Phenylene-diamine $C_6H_4(NH_2)_2$. [140°].

Formation.-1. By reduction of p-nitroaniline and of p-di-nitro-benzene (Hofmann, Pr. 11, 518; 12, 639; Zincke, B. 7, 871).—2. By reduction of C₆H₅, N₂.C₆H₄NH₂ (Martius a. Griess, Z. 1866, 136).—3. By distilling (5,2,1)-di-amidobenzoic acid (Griess, B. 5, 201).—4. In the pre-paration of induline by heating C₈H₈,N₂,C₈H₄,NH₂ with aniline hydrochloride (Witt; Fischer a. Hepp, B. 23, 839).

Preparation.—From acetanilide by nitration

and reduction with tin and HCl.

Properties. - Plates (by sublimation) or monoclinic crystals (from water) changing in the solution to trimetric plates. M. sol. water, v. sol. alcohol and ether. Forms a hydrate C₆H₈N₂2aq [80°]. With PbO₂ and aqueous HOAc it gives a bright bluish-green colour (Lauth, C. R. 111, 975). p-Phenylene-diamine is very poisonous (Vignon, C. R. 107, 533). A mere trace of a mixture of m- and p- phenylene-diamine gives on warming with HOAc and K.C. a blue colour, changing on boiling to red (Witt, C. J. 35, 356; Meldola, C. J. 51, 105).

Reactions. - 1. Oxidised to quinone by H,SO, and MnO. Quinone is also formed together with NH, by oxidation of the sulphate with K₂Cr₂O₇ (Meldola a. Evans, C. J. Proc. 5, 115).—2. When exposed to air, O, H₂O₂, or K₂FeCy₆ p-phenylene-diamine and its salts are oxidised to C₆H₆N₂ a feeble red base with green lustre [230°] yielding C.H.AcN, [294°] (Bandrowski, M. 10, 126).—3. Chlorine passed into a solution in HOAc forms tetra-chloro-quinone. -4. FeCl, added to a solution of the hydrochloride saturated with H2S ppts. Lauth's violet, which is v. sol. pure water, but ppd. by salts (Lauth, C. R. 82, 1441).-5. Potassium nitrite The sulphocyanide is converted by heat into $C_cH_4(NH.CS.NH_2)_2$ (Lellmann, A. 221, 8). m-Phenylene-diamine is somewhat less poisonadded to a solution of the hydrochloride forms C_eH₄(NH₂Cl)N₂Cl, and in presence of a large excess of HCl it forms C_eH₄(N₂Cl)₂.—6. Mixed

with aniline (2 mols.) and oxidised by K2Cr2O, it yields pheno-safranine. - 7. The sulphocyanide is converted by heat into $C_0H_1(NH.CS.NH_2)_2$ (Lellmann, A. 221, 8).—8. Bleaching-powder solution added to a solution in HClAq ppts. yellowish-white quinone di-chloro-di-imide which detonates at 124° (Krause, B. 12, 47).—9. Aceto-acetic ether at 170° forms C₁₄H₁₆N₂O₄ [176°] which is probably C.H.(NH.CO.CH2.CO.CH2)2 (Knorr,

B. 17, 545; 19, 3303).
Salts.—B"H_{**}Cl_{**}. Triclinic tables, v. so water, almost insol. HClAq.—B"H_{**}PtCl_{**}. Triclinic tables, v. sol. B'H₂Sn₂Cl₅ (Hübner, A. 209, 366).—B"H₂Br₃. $B''H_2S_2O_3$ (Bernthsen, A. 251, 62).— $B''H_2SO_3$ Micaceous plates, sl. sol. water (Vignon, Bl. [2] 50, 152).—B"H₂C₂O₄. S. 15 at 15°.

Di-formyl derivative C,H,(NH.CHO)2. [204°]. Amorphous mass (Wundt, B. 11, 828).

Mono-acetyl derivative

 $C_cH_4(NH_2)(NHAc)$. [161°]. Formed by reduction of $C_0H_4(NO_2).NHAc$ with iron and acetic acid (Nietzki, B. 17, 343). Needles, sl. sol. cold water. -B'2H2PtCl

C6H4(NHAc)2. Di-acetylderivative [above 295°]. Formed by boiling the base with HOAc (Biedermann a. Ledoux, B. 7, 1531).

Small octahedra, sol. HOAc.

Benzoyl derivative C.H.(NH2) (NHBz). [128°]. Formed by reducing benzoyl-p-nitro-aniline (Hübner, A. 208, 295). Plates, sl. sol. water.—B'HCl.—B'2H2SO4. Needles.

Di-benzoyl derivative C.H.(NHBz). [above 300°]. Formed from the base, BzCl, and NaOHAq (Hinsberg, A. 254, 254). Plates. Yields a nitro- product [251°].

o-Oxy-benzoyl derivative C₄H₄(NH₂)(NH.CO.C₅H₄.OH). [158°]. Glittering

needles (Bell)

Hexahydride CeH10(NH2)2. Di-amidohexamethylene. Formed by reducing the dioxim of quinone tetrahydride in alcohol with Na (Baeyer a. Noyes, B. 22, 2172). Liquid, with faint ammoniacal odour, forming crystalline salts and an acetyl derivative melting above 310°.

References.—Bromo-, Bromo-nitro-, Chloro-, NITRO-, and OXY- PHENYLENE-DIAMINE.

PHENYLENE-DIAMINE SULPHONIC ACID v. DI-AMIDO-BENZENE SULPHONIC ACID.

PHENYLENE-DI-AMYL-DIAMINE

[1:4] $C_0H_4(NH_*C_5H_{11})_2$. [49°]. Colourless crystail (Baeyer a. Noyes, B. 22, 2173)

DIPHENYLENE AZONE $\stackrel{C.H.N}{C_{3}H.N}$. [156°]. Formed, together with $C_{12}H_{3}N_{2}O_{2}$ [240°] and $C_{12}H_6N_2O$ [152°] by boiling di-o-nitro-diphenyl with alcohol and zinc-dust (Täuber, B. 24, 3081, 3883). Greenish-yellow needles by sublimation, not volatile with steam. Almost insol. water, v. sol. alcohol, m. sol. ether, v. e. sol. HOAc. Reduced by tin and HCl to C₁₂H₁₀N₂HCl, which crystallises in white needles, rapidly reoxidised by air to diphenylene-azone.

Salts. - B'HCl. Flat bronzed prisms .-B'HAuCl₄. — B'₂H₂PtCl₆: yellow needles. — B'H₂CrO₄.—B'C₆H₃N₈O₇. [194°]. Brown needles.

PHENYLENE-BENZALDEHYDINE v. vol. i. p. 109.

PHENYLENE-BENZAMIDINE v. BENZENYL-PURNYLENE-DIAMINE.

Vol. IV.

PHENYLENE-DI-BENZIDENE-DIAMINE v. DI-BENZYLIDENE-PHENYLENE-DIAMINE

PHENYLENE-BENZYL-m-DIAMINE

C₆H₄(NH₂).NHC,H₇. Formed by reducing benzyl. m-nitro-aniline (Meldola a. Coste, C. J. 55, 597). Oil. -B"2HCl: hexagonal tablets. FeCl. colours its solution red and gives a white pp. on heating. Di-bensoyl derivative

 $C_6H_4(NHBz).NBz.C_7H_7.$ [178°]. Sol. alcohol.

Phenylene-benzyl-p-diamine. [30°]. Got by reducing benzyl-p-nitro-aniline (Meldola a. Coste, C. J. 55, 590) or p-nitroso-benzyl-aniline (Boeddinghaus, A. 263, 302). Wax-like mass of crystals, turning red in air. When mixed with the m-isomeride it yields di-benzyl-amidoindamine on oxidation.—B'2HCl. White plates, v. sol. water. FeCl, colours its aqueous solution green, changing to red. After saturation with H2S it gives a blue pp. with FeCl.

Di-acetyl derivative. [117°]. Prisms. Di-benzoyl derivative. [124°]. Needles. Phenylene-di-benzyl-diamine v. vol. i. p.

Phenylene-tetra-benzyl-m-diamine

 $C_6H_4(N(C_7H_7)_2)_2$. [81°]. Formed by heating phenylene-m-diamine with NaOHAq and benzyl chloride. Amorphous powder, sol. HOAc.

Phenylene-tetra-benzyl-p-diamine. White crystalline powder. HOAc and CrO,

form a blue-violet colouring matter.

PHENYLENE - BENZYLIDENE - DI - METHYL-DIAMINE C₁₅H₁₆N₂ i.e. C₆H₅.CH:N.C₆H₄.NMe₂. [93°] (Calm, B. 17, 2940); [101°] (Köhler, A. 241, 361). Formed from phenylene-di-methyl-p-diamine and benzoic aldehyde. -B"2HCl.

PHENYLENE - BENZYL - DI - METHYL -DIAMINE C.H.CH.NH.C.H.NMer. [48]. Formed by reducing the preceding body in alcohol with sodium-amalgam (Köhler, A. 241, 861). Plates. Yields a nitrosamine [128°].

PHENYLENE-BLUE or Lauth's violet v. MIDO-IMIDO-DI-PHENYL-SULPHIDE.

PHENYLENE - BROMO - ACETYLENE -KETONE v. Bromo-oxy-indonaphthene.

C.H.CH.CH. DI-PHENYLENE-BUTANE Diphensuccindene. [100°]. Formed by heating (β)-di-phenyl-succinic anhydride with HI and P and 180° (Roser, A. 247, 156). White needles.

PHENYLENE-ISOBUTYL-p-DIAMINE
C,H,NH.C,H,NH2. [39°]. Formed by reduction of p-nitroso-isobutyl-aniline (Wacker, A. 243, 299). Plates (from benzene-ligroin).
PHENYLENE-BUTYLIDENE-o-DIAMINE

C,H,<NH>CHPr. [233°]. Formed from phenylene-o-diamine and isobutyric aldehyde (Lassar-Cohn, B. 22, 2724). Needles (by sublimation).—B'HCl. [184°].—B'zH_PtCl.

The isomeric compound from phenylene-m-diamine melts at 216°, while that from phenylene-p-diamine is oily, and yields B'2H,PtCl

DI - PHENYLENE - ISOBUTYL - QUINOX -ALINE C.H.C:N C.H.CH.Pr. [147°]. Fo med from phenanthraquinone in HOAc and isobutylphenylene-diamine in alcohol (Gelzer, B. 20, 3256). Yellowish needles.—B'HCl: needles. o-PHENYLENE-DICARBAMIC ETHER

J.H.(NH.CO.Et). [88°]. Formed by heating

2-phenylene-diamine with ClCO, Et (Snape, C. J.) 49, 259). Needles (from alcohol).

[145°]. m-Phenylene-dicarbamic ether. p-Phenylene-dicarbamic ether. [193°]. Got by heating p-phenylene dicyanate with alcohol (Gattermann a. Wrampelmeyer, B. 18, 2605).

Di-phenylene-dicarbamic acid. ether CO2Et.NH.C6H4.C6H4.NH.CO2Et. [230° Formed from benzidine and ClCO, Et or by boiling diphenylene dicyanate with alcohol (Snape, C. J. 49, 256; Schiff a. Vanni, A. 258, 369).

Di-phenyl ether. [c. 240°]. Made by heating diphenylene dicyanate with phenol at 140° (Snape). Tables, sl. sol. alcohol.

DI-PHENYLENE-CARBINOL C13H10O i.e. С.Н. >CH.OH. [153°]. Fluorene alcohol. Formed by reducing diphenylene ketone with sodium-amalgam (Barbier, A. Ch. [5] 7, 504). Hexagonal plates (from benzene). Oxidised by CrO₃ to diphenylene ketone. Ac₂O at 100° forms an acetyl derivative [75°]. When heated above its melting-point, diphenylene carbinol yields C₂₄H₁₈O [290°], a nearly colourless resin. o-PHENYLENE CARBONATE C₆H₄CO₈.

[118°]. (225°-230°). Formed from sodium pyrocatechin and ClCO, Et (Wallach, A. 226, 84). Four-sided prisms (from alcohol-benzene).

m-Phenylene carbonate $C_6H_4 < 0 > CO$

Formed by heating resorcin with cyanuric acid and ZnCl2, and also by the action of COCl2 on resorcin (Birnbaum, B. 14, 1753). Amorphous solid, sol. HOAc

m-PHENYLENE DICARBONIC ETHER (298°-302°). C_eH₄(O.CO₂Et)₂. (298°-302°). Formed from sodium resorcin and ClCO₂Et in ether (Wallach, A. 226, 84). The p-isomeride, $[100^{\circ}]$, (310°) , is got in like manner from hydroquinone (Bender, B. 13, 697)

PHENYLENE-CHLORO-OXY-ACETYLENE KETONE v. CHLOBO-DI-OXY-INDONAPHTHENE.

PHENYLENE - CINNAMYLIDENE - DI-METHYL DI-AMINE C17H18N2 i.e.

C.H.OH.CH:N.C.H.NMe2. •[141°]. Formed from cinnamic aldehyde and phenylene-di-methyl-p-diamine (Nuth, B. 18, 575). Needles. PHENYLENE - CUMINYLIDENE - DI -

METHYL-DIAMINE C18H22N; i.e.

[100°]. C₂H₄,C₆H₄,CH:N.C₆H₄,NMe₂. Formed from cuminol and phenylene-di-methyl-diamine (Nuth, B. 18, 575; Uebel, A. 245, 299). Needles. m-PHENYLENE DICYANATE C. H. (N:CO)2.

[51°]. Got by distilling m-phenylene-dicarbamic ether (Bender, Bn. 3, 897). Crystals.

p-PHENYLENE DI-CYANATE O.H. (N:C:0)2 [91⁵]. (231° at 745 mm. i.V.). V.D. 5·79 (for 5·54). Prepared by passing a stream of carbonyl chloride COCl₂ through p-phenylene-diamine hydrochloride heated to 200°-250° (Gattermann a. Wrampelmeyer, B. 18, 2604). Pungent vapour. less needles. Sublimable. Reacts like phenyl-cyanate, forming di-ureas with amines, and di-carbamic ethers with alcohols.

PHENYLENE DI-CYANIDE v. Nitrile of

PHTHALIC ACLD.

PHENYLENE ETHYL-ACETAMIDINE

 $C_{\bullet}H_{\bullet} < NEt > C.OH_{\bullet}$. [180°]. Formed by the action of Ac2O on phenylene-ethyl-o-diamine (Hempel, J. pr. [2] 89, 200; 41, 166). Plates.

PHENYLENE-ETHYL-o-DIAMINE

 $C_6H_4(NH_2)(NHEt)$. (249°). Formed by reducing o-nitro-ethylaniline with tin and HCl (Hempel, J. pr. [2] 39, 199; 41, 164). turning red in air and light.

Acetyl derivative C.H.N.EtAc. [104°].

Phenylene-ethyl-m-diamine

C₆H₄(NH₂)(NHEt). (276° uncor.). Oil, formed in like manner (Nölting a. Stricker, B. 19, 546). B"H2Cl2: white powder.

 $(270^{\circ}).$ sol. benzene (Nölting a. Collin, B. 17, 267; Schweizer, B. 19, 149; Fischer a. Hepp, B. 19, 2994; Bernthsen a. Goske, B. 20, 930).—
B'H.SO.: needles or prigns. Phenylene-ethyl-p-diamine. B"H2SO4: needles or prisms.—B"H2Cl2. Plates, v. sol. water.—B"H2PtCls.

Phenylene-di-ethyl-m-diamine

 $C_6H_4(NH_2)(NEt_2)$. (277°). Formed by reducing m-nitro-di-ethyl-aniline (N. a. S.; Groll, B. 19,

200). Oil.

Phenylene-di-ethyl-p-diamine. (261° cor.). Formed by reducing nitroso-diethyl aniline or C₆H₄(NEt₂).N₂.C₆H₄.NEt₂ (Lippmann a. Fleissner, B. 16, 1424). Oil.—B"₂H₄Cl₄PtCl₄. Red crystals.—B"ZnCl₂ 2aq. Converted by treatment in aqueous solution with aluminium sulphate, Na₂S₂O₃, and K₂Cr₂O₇ into $NEt_2.C_6H_3(NH_2).S.SO_3H$ [229°], whence by diazotisation $C_6H_3(NEt_2) < \frac{N}{8}^2$ [1076] may be got (Bernthsen, A. 251, 1).

Phenylene-s-di-ethyl-p-diamine.

Nitrosamine C₆H₄(NEt.NO)₂. [90°]. Formed from p-phenylene-diamine by successive treatment with EtBr and HNO₂ (Nietzki, B. 16, 465). Yellowish plates (from benzene-ligroin).

Phenylene-tetra-ethyl-p-diamine

 $C_6H_4(NEt_2)_2$. [52°]. (280° uncor.). Got by heating phenylene-di-ethyl-diamine or N2(C6H4.NEt2), with EtI (Lippmann a. Fleissner, B. 16, 1427; M. 4, 301). Monoclinic plates (from dilute alcohol), a:b:c = .99:1:1.833; $\beta = .90°30'.--B''H_2PtCl_{\bullet}$. — B"H,Hg,Cl. Monoclinic prisms. — B",I.: black prisms. — B"2HI.—B"HI,.

PHENYLENE-ETHYL-BENZAMIDINE

C.H. CPh. [81°]. Got by ethylation of phenylene-benzamidine (Howe, Am. 5, 421) .-B'HČl 3aq.—B',H,PtCl,.—B'HNO, aq [155°],-B'EtOH [186°].— B'EtI aq. — B'EtCl 2aq. — B'sEt2PtCl .- B'EtHSO, aq

PHENYLENE-ETHYLENE-o-DIAMINE

C.H. NH.CH. Quinoxaline tetrahydride.
[97°]. (290°). V.D. 4.44 (calc. 4.64). Formed by heating pyrocatechin with ethylene-diamine hydrate at 200°-210° (Merz a. Ris, B. 20, 1191). Formed also by reducing quinoxaline with alcohol and Na. Plates (from water). FeCl, colours its dilute aqueous solutions blue or violet. Alkaline K₃FeCy₆ oxidises it to quinoxaline.—B"₂3HCl. Scales. $B''_2H_2C_2O_4$. [184°]. — [above 120°]. Yellow crystal-[above 150°]. B",2C,H,N,O,. line pp., sl. sol. ether.

C.H.: N.Ac.: C.H. Acetyl derivative [144°]. [850° uncor.). Granules, sl. sol. cold water.

Di-phenylene-ethylene-tetramine v. vol. ii. p. 491

PHENYLENE-ETHYLENE-DISULPHONE C₂H₄(SO₂)₂C₂H₄. Formed by heating ethylene bromide with potassium benzene m-disulphinate (Otto, J. pr. [2] 86, 450). White plates, m. sol. conc. HNO..

PHENYLENE DI-ETHYL DIKETONE

C₆H₄(CO.C₂H₅)₂. [220°]. Formed from terephthalic chloride and ZnEt₂ in ether (Münchmeyer, B. 19, 1850). Needles (from ether).

PHENYLENE DI-ETHYL DISULPHONE $C_8H_4(SO_2Et)_2$. [142°]. Obtained by heating $C_8H_4(SO_2K)_2$ [1:3] with EtBr at 100° (Otto, J. pr. [2] 36, 449). Colourless plates.

PHENYLENE-FURFURALDEHYDINE vol. i. p. 109.

PHENYLENE-DIGLYCOLLIC ACID v. Oxy-DIPHENYLENE-ACETIC ACID.

DIPHENYLENE-DIHYDRAZINE C, H, N, i.e. N₂H₃, C₆H₄, C₆H₄, N₂H₃. [167°]. Formed from benzidine hydrochloride by diazotisation and reduction by Na, SO, or SnCl₂ (Arheidt, A. 239, 206). Small white plates, sol. hot water. Aqueous potassium cyanate forms the semicarbazide $C_{11}H_{16}N_0O_2$ [307°]. Nitrous acid forms a dinitrosamine $C_{12}H_{12}N_4O_2$ [112°]. Pyruvicacid forms $C_{18}H_{14}N_4O_4$ [197°]. Acetone yields $C_{18}H_{22}N_4$ [198°], whence fasion with ZnCl2 yields diphenyl-di-methyl-indole C₁₈H₁₆N₂ [270°].

Reference. - DI-OXY-DI-PHENYLENE DIHYDRAZ-INE.

DIPHENYLENE-IMIDE v. CARBAZOLE. C,H,>co. KETONE DIPHENYLENE

[83°]. (above 300°). Formation.—1. By distilling phenanthraquinone or diphenyl di-o-carboxylic acid with CaO (Fittig a. Ostermayer, A. 166, 373; Anschütz a. Schultz, B. 9, 1400).—2. By heating anthraquinone with lime (Anschütz, B. 11, 1213).-3. By dry distillation of the Ag salt of its dicarboxylic acid (Bamberger a. Hooker, B. 18, 1034; A. 229, 156).-4. By passing a mixture of phenol and CS₂ over hot copper (Carnelley a. Dunn, C. J. Proc. 4, 53; B. 21, 2005, who regard the product thus obtained as isomeride).

Properties .- Yellow needles or plates (from alcohol). Yields phenyl-benzoic acid by potashfusion. Distillation over zinc-dust gives diphenylene-methane [113°] (Fittig, B. 6, 187). Forms a nitro-derivative [220°] and a di-nitroderivative [290°] (Schultz, A. 203, 104). Alkaline KMnO4 oxidises it to phthalic acid (Anschütz a. Tapp, B. 11, 213). H₂SO, forms at 260° a disulphonic acid, whence CaA" (Schmidt a. Schultz, A. 207, 345).

Oxim. [192°] (Spiegler, M.5, 195).

References .- BROMO-, DI-CHLORO-, NITRO-, and OXY- DIPHENYLENE KETONE.

DIPHENYLENE KETONE CARBOXYLIC ACID [1:2] $C_0H_4 < \frac{CO}{2} > C_0H_3$, $CO_2H \begin{bmatrix} 1 & 6 \end{bmatrix}$, [192°]. Formed by oxidation of fluoranthene by chromic acid mixture (Fittig, A. 193, 149; 200, 6).

Orange-red needles (from dilute alcohol).

Reactions.—1. Potash-fusion gives diphenyl
om-dicarboxylic acid.—2. Yields diphenylene
ketone on distillation.—3. Distillation with zincdust gives diphenylene-methane (fluorene).

Salts.—BaA', 4aq.—CaA', 2aq.—AgA'. Diphenylene ketone carboxylic acid. [above 275°]. Formed by heating the dicarboxylic acid *Bamberger a. Hooker, B. 18, 1034; A. 229, 158). Pale yellow needles, sl. sol. alcohol. May be sublimed.—BaA'2.—AgA': yellow pp. Diphenylene ketone carboxylic acid

[1:2] $C_0H_4 < {}^{CO} > C_0H_3 \cdot CO_2H \left[{1 \over 2} 8 \right]$. [227° cor.].

Formation (Graebe, B. 13, 1303; B. 20 845; A. 247, 275).-1. From diphenyl dive carboxylic acid by heating with H2SO, at 120° or with POCl₃ or ZnCl₂.—2. By heating diphenylene dicarboxylic acid with H,SO, for 10 minutes at 150°.

Properties. -Yellow needles, v. sol. hot alcohol. Conc. H2SO4 forms a red solution.

Reactions.—1. Potash-fusion yields diphenyl di-o-carboxylic acid. - 2. PCl, (1 mol.) forms C_8H_4 COCl) CO [128°], which may be crystallised from ligroin. — 3. PCl, (2 pts.) at 160° yields C.H. (COCI) CCl₂ [95°], which yields fluorene carboxylic acid on reduction with zincammonia reduce it to C_sH₄(CO₂H) > CH(OH).—

Dhonol and dust and dilute acetic acid .- 4. Zinc-dust and 5. HI and P give fluorene. 6. Phenol and SnCl₄ at 120° give C_0H_4 (CO₂H) $C(C_0H_4.OH)_2$ [165°], which yields AgA' and $C_{2b}H_{1c}Ac_2O_1$ [c. 130°].-7. Resorcin gives rise, on heating, to C.H. $C_{\mathfrak{g}}^{\mathfrak{g}^{-1}}_{\mathfrak{g}}(CO_{\mathfrak{g}}H)$ $C(C_{\mathfrak{g}}H_{\mathfrak{g}}(OH)_{\mathfrak{g}})_{\mathfrak{g}}$, a yellowish-brown powder which exhibits green fluorescence in alkaline solutions.

Salts. - NH, A' aq. - NaA' 6aq: yellow needles, v. sol. hot water.—AgA': yellow pp.

Methyl ether Med'. [132°]. Needles.

Ethyl ether Eth'. [103°]. Needles.

Chloride C₁₄H,ClO₂. [128°].

Antide C₁H,(NH₂)O₂. [226°]. Silky yellow
needles (containing ½HOEt). Formed from the chloride, and also by heating phenanthraquinone mono-oxim with H₂SO₄ at 100° (Wegerhoff, B. 21, 2357).

 $Oxim\ C_{14}H_7(CO_2H)(C:NOH).\ [263°].$ Phenyl-hydrazide $C_{14}H_{7}(CO_{2}H)(C:N_{2}HPh)$. [205°]. Brownish-

yellow prisms, m. sol. alcohol. Diphenylene ketone dicarboxylic acid

 $C_eH_4 < CO > C_eH_2(CO_2H)_2 [\frac{1}{2}3:6].$ Formed by oxidation of retene-quinone and of oxyisopropyldiphenylene-ketone carboxylic acid (Bamberger a. Hooker, B. 18, 1033; A. 229, 151). Yellow needles (from HOAc). Not melted at 270°, but loses CO₂ at a higher temperature. Yields diphenyl on heating with CaO. Potash-fusion forms diphenyl tricarboxylic acid.—BaA".—Ag₂A": yellow pp.

Methyl ether Me.A". [184°]. Yellow needles.

Ethyl ether EtA". [115°]. Needles. Oxim C₁₂H₃(CO₂H)₂:C(NOH). Yellow pp., not melted at 280°

DIPHENYLENE KETONE OXIDE

 $CO < \stackrel{C_0H_4}{C_0H_4} > 0$. Xanthone. [174°]. (350°) at 780 mm. (Graebe, A. 254, 280). S. (alcohol) · 7 in the cold; 8.5 at 78°

Formation .- 1. Obtained by oxidation of $CH_2 < C_0H_4 > O$ with CrO_0 and HOAc (Me:z a. Weith, B. 14, 192).—2. By the dry distillation of salicylide; the yield being 35 p.c. of the theoretical (Perkin, B. 16, 339).—3. By distilling phenyl phosphate with sodium m- or p-oxybenzoate (Richter).-4. By the action of POCl, on potassium-salicylate of ethyl.-5. By distilling neutral or basic sodium salicylate with P₂O₅.—6. In small quantities, when salicylic acid is distilled .-- 7. By strongly heating sodium o-chlorobenzoate.-8. From sodium o-phenylbenzoate and POCl_s.—9. To the amount of 12 p.c. of the theoretical quantity, by the action of ochlorobenzoyl chloride on basic sodium salicylate.—10. By the action of conc. H₂SO₄ on [1:2] C_oH₄(OPh).CO₂H (Graebe, B. 21, 503).— 11. By distilling phenyl salicylate (Seiffert, J. pr. [2] 31, 479).—12. By heating salicylic acid (1 mol.) with PCl₅ (1 mol.), then adding phenol and distilling (Klepl, J. pr. [2] 28, 217. 13. By distilling calcium p-oxy-benzoate (Goldschmidt, M. 4, 128). -14. A product of the electrolysis of a solution of phenol in KOHAq at 2005 (Bamberger a. Berlé, B. 24, 3212).

Preparation.—1. Phenyl phosphate (20 g.) and sodic salicylate (20 g.) are distilled together; a violent action ensues; when it is over the residue is distilled over at 400°, best in a current of air. The crude product is shaken with solution of NaOH and distilled with steam. The residue is extracted with alcohol, from which, on cooling, the ketone crystallises (R. Richter, J. pr. [2] 28, 276).—2. By distilling salicylic acid with Ac₂O, the salicylide which is first formed splitting off CO₂ when the temperature rises (Perkin, C. J. 43, 35). In this process there is formed a by-product C₁₄H_aO₃ [192°] which is converted by boiling alcoholic potash into an acid C₁₄H_aO₄ [275°] (A. G. Perkin, C. J. 43, 187).

Properties.—Long white needles, grouped concentrically. M. sol. ether, alcohol and benzoline, more sol. benzene and chloroform, insol. water. The solution in conc. H., SO, shows blue fluorescence (Graebe, B. 15, 1679). Does not react with hydroxylamine or phenyl-hydrazine (Spiegler, B. 17, 808).

Reactions.—1. Reduced by zinc-dust or by HI to CH₂ $< \stackrel{C_0}{C_0} \stackrel{H}{H} > 0$ [100.5°] but no further.—

2. Potash-fusion converts it into salicylic acid and phenol.—3. If the fusion be stopped when the mass is pasty (at about 200°), dissolved in water and ppd. by HCl, di-oxy-benzophenone is obtained, CO(C₈H₁OH)₂.—4. Sodium-amalgam reduces the di-phenylene-ketone oxide (in dilute alcoholic solution) to C₂H₁₈O₃ [200°]. This body crystallises from CHCl₃ in prisms. It is split up by chromic acid into di-phenylene-ketone oxide and diphenylene methane oxide, so that it is perhaps a molecular compound of these two bodies.—5. Bromine at 200° forms a dibromo-derivative [212°].—6. Fuming H₂SO₄ gives a disulphonic acid which forms BaA^maq, crystallising from hot water in needles (A. G. Perkin, C. J. 43, 192).

Isomer.de C₁₃H₄O₂. [91°] (R.); [96°] (G.). A product of the action of POCl, on sodium salicylate (Richter, J. pr. [2] 28, 294). Formed also by heating the sulphate of o-diazobenzoic acid with phenol (Griess, B. 21, 981). Needles, v. e. sol. ether. Yields diphenyl on distillation

with zinc-dust. Br gives C₁₂H₂BrO₂ [198°]. H₂SO₄ yields a disulphonic acid, which gives BaA"aq crystallising in needles.

Dixanthone C₆H₄ C_{CO} C₅H₂ C_{CO} C₅H₄.

[256°]. Occurs in the product of the action of salicylic acid on resorcin (Kostanecki a. Seidmann, B. 25, 1654). It crystallises from HOAc in groups of needles, and its solution in H₂SC₄ shows greenish fluorescence.

References. — DI-AMIDO-, NITRO-, and Oxy-

DI-PHENYLENE KETONE OXIDE.

PHENYLENE MERCAPTAN v. DI-THIO-RESORCIN and DI-THIO-HYDROQUINONE.

Diphenylene mercaptan C_sH₄(SH).C_sH₄(SH). [176°]. Formed by diazotising benzidine, and treating the product with potassium xanthate and alcoholic potash successively (Leuckart, J. pr. [2] 41, 212; cf. Gabriel, B. 13, 590). White plates (from alcohol).

Di-methyl ether Me₂A". [184°]. Plates. Di-ethyl ether Et₂A". [135°]. Plates.

oo-DIPHENYLENE-METHANE v. FLUOR-

(γ) - Diphenylene - methane CH₂ C_sH₄.

Methylene-diphenyl. [118°]. (295° uncor.). V.D.
84 (calc. 83). Occurs, together with the (δ)isomeride, among the products of the passage of
a mixture of benzene and toluene through a
red-hot tube (Carnelley, C. J. 37, 708). Pearly
plates (from alcohol), sl. sol. cold alcohol, v. sol.
ether. Its alcoholic solution shows faint blue
fluorescence. CrO_s and HOAc oxidise it to
C₁₃H₅O₂ [281° cor.]. Bromine in ether gives
C₁₃H₅Br₂ [162°]. Picric acid forms a compound
C₁₃H₁₀C₆H₃N₃O₇ crystallising in blood-red needles

(δ)-Diphenylene-methane $C_{13}H_{10}$. [205°]. (320°). V.D. 86°2 (calc. 83). Formed as above. Pearly plates (from alcohol). Oxidised by CrO_s in HOAc to $C_{13}H_sO_2$ [288° cor.] which sublimes in white needles.

DIPHENYLENE-METHANE OXIDE

CH₂ C_8H_4 O. [100°]. (315° cor.). Formed, to the amount of 2 p.c., in the preparation of Ph₂O by heating phenol with AlCl₃ (Merz a. Weith, B. 14, 191). Formed also from diphenylene ketone oxide by heating with fuming HIAq at 160° or by distilling with zinc-dust (R. Richter, J. pr. [2] 28, 280; Graebe, \triangle . 254, 282); and by distilling euxanthone over heated zinc-dust (Wichelhaus a. Salzmann, B. 10, 1399; Graebe a. Ebrard, B₁ 15, 1678). White plates (from alcohol).

Reactions.—1. Oxidised by CrO₂ or dilute HNO₃ to diphenylene ketone oxide.—2. PCl₃ forms HCl and a product whence water at 0° produces C₁₂H₂O.PO(OH)₂ [255°-260°] an acid forming Ag₂A" and (NH₂)₂A".—3. Br forms C₁₂H₄Br₆O and C₁₃H₄Br₇O [136°].

DIPHENYLENE - METHANE SULPHIDE CH₂ C₆H₄ S. [128°]. (342°). Formed by

reducing $CO < C_0H_4 > S$ with HI and P at 170°; and also by passing phenyltolyl sulphide through a red-hot tube (Graebe a. Schultess, A. 263, 12). Needles, sl. sol. cold alcohol, v. e. sol. CHCl₄.

DIPHENYLENE - METHANE SULPHONE $\mathbf{CH_2} < \mathbf{C_6H_4} < \mathbf{SO_2}$. [170°]. Formed by heating $CO < \stackrel{C_0 H}{C_0 H} > SO_2$ with HI and P at 170° (Graebe a. Schultess, A. 263, 15). Needles, v. sol. hot alcohol.

PHENYLENE-METHENYL-AMIDINE

 $C_{\bullet}H_{\bullet} < N^{H} > CH$. [167°]. (above 360°). Formed by boiling phenylene-o-diamine with formic acid (Wundt, B. 11, 826; Fischer, B. 22, 645). Trimetric crystals (from alcohol).—B'HCl aq.—B'HAuCl. Yields on methylation $C_8H_8N_2$ [33°] (278° at 730 mm.).

 $CO_2H.C_6H_3 < NH > CH.$ Carboxylic acid Formed by reducing the formyl derivative of (2,1,4)-nitro-amido-benzoic acid (Zehra, B. 23, 3634). White needles, blackening without fusion when heated.—HA'HCl: white needles.

m - PHENYLENE - METHYL - DIAMINE NH₂).NHMe. Methyl - phenylene -Amido-methyl-aniline (265°-270° Formed by reduction of m-nitro-[1:3] $C_6H_4(NH_2).NHMe$. diamine. uncor.). methyl-aniline (Nolting a. Stricker, B. 19, 548). Oil. Gives a methyl-chrysoidine with diazobenzene chloride.

 $p extsf{-} extbf{Phenylene-methyl-diamine}$ [1:4] $C_0H_4(NH_2)(NHMe)$. (258° uncor.). Formed by reduction of p-nitroso-methyl-aniline (Fischer, B. 19, 2992) or of C₆H₄(NHMe).N₂.C₆H₄.SO₃H (Bernthsen a. Goske, B. 20, 929). Oil, v. sol. water. FeCl, gives a red colour, which on successive treatment with HCl and H2S becomes blue (dimethylthionine). — B'2H2SO4: white needles (from water).

m-Phenylene-di-methyl-diamine

[1:3] $C_6H_4(NH_2)(NMe_2)$. (269°) at 740 mm. S.G. 25 995 (Groll, B. 19, 200); (258°) (Staedel a. Bauer, B. 19, 1945). Formed by reduction of m-nitro-di-methyl-aniline. Oil. With nitrous acid it yields a colouring-matter closely resembling Bismarck brown. With diazobenzene chloride it yields a di-methyl-chrysoidine which dies a somewhat redder shade than chrysoidine. PbO₂ and aqueous HOAc gives a yellowish-brown colour (Lauth, C. R. 111, 975).—B'HCl.

Acetyl derivative CaH4(NMe2)(NHAc).

 $[87^{\circ}].$

p-Phenylene-di-methyl-diamine

[1: $^{\uparrow}$] $C_6H_4(NH_2)(NMe_2)$. [41°]. (257° i.V.). Prepared by reduction of nitroso-di-methyl aniline and of (C₈H₄NMe₂).N₂O by tin and HCl (Schraube, B. 8, 616; Wurster, B. 12, 522; Weber, B. 10, 760). Prepared also by reduction of C₆H₄(NMe₂).N₂.C₆H₄.SO₃H (E. Fischer, B. 16. 2235). Formed also together with di-chloro-pphenylene-diamine and di-chloro-p-phenylenedi-methyl diamine by boiling nitroso-di-methylaniline with HCl (S.G. 1.2) (Möhlau, B. 19, 2010).

Properties.—Long white needles or short prisms, v. sol. water, alcohol, and ether. PbO₂ and aqueous HOAc give a magenta-red solution changing to violet-black (Lauth, C. R. 111, 975). Colours wood, and paper made from wood,

crimson (Wurster, B. 20, 808).

Reactions.—1. Yields quinone on oxidation with MnO₂ and H₂SO₄.—2. Bromine in HOAc forms C₂H₁₁N₂Br, which crystallises from its red

alcoholic solution in green scales [146°] (Wurster a. Sendtner, B. 12, 1803).—3. H₂S and an oxidising agent (e.g. FeCl,) gives methylene-blue (v. vol. iii. p. 265). Methylenered $S_2 < \stackrel{C_6H_2}{N} > NMe_2Cl$ by-product is (Bernthsen, A. 230, 137; 251, 1).-4. Na₂S₂O₃ and CrO, give C,H,(NMe,)(NH,)S.SO,H, which yields methylene-blue on further oxidation.-5. Aldehydes and ketones form products of condensation, e.g. C₆H₄(NMe₂).N:CHPh [98°] (Oalm, B. 17, 2938), C₆H₄(NMe₂).N:CH.C₆H₄OH [1:2] [134°], and C₆H₄(NMe₂).N:CH.C₆H₄.OMe [1:4] [148°] (Nuth, B. 18, 574; Steinhart, A. 241, 343). Benzil and alcoholic KOH form CPhBz:N.C₆H₄NMe₂ [139°]; benzoyl-acetone forms in like manner CH2Bz.CMe:N.C6H4NMe2 [136°] (Vogtherr, B. 25, 635). Benzoin gives CHPh(OH).CPh:N.C. H., NMe. [127°] on heating, while deoxybenzoin forms C22H22N2 [139°].— 6. Oxalic ether forms, on heating, the ether C₆H₄(NMe₂).NH.CO.CO₂Et [117°] which yields a corresponding acid [192°] and amide [259°] (Sendtner, B. 12, 530). Salts.—B"2HCl. Very deliquescent plates.

-B"H₂SnCl₄. Cubes.—B"H₂PtCl₆.

Acetyl derivative C₀H₄(NMe₂)(NHAc). [130°]. (355°). Small leaflets or needles.

p-Phenylene-tri-methyl-diamine [1:4] C₆H₄(NMe₂)(NMeH). (265°). Obtained by reduction of its nitrosamine, which is formed by the action of nitrous acid on p-phenylene-tetramethyl-diamine (Wurster a. Schobig, B. 12, 1809). Oil.

Acetyl derivative. [95°]. Prisms. Nitrosamine C₆H₄(NMe₂)(NMe.NO). [99°]. Leaflets or tables, sl. sol. cold water.

m-Phenylene-tetra-methyl-diamine [1:3] C₃H₁(NMe₂)₂. [-2°]. (266° i.V.). S.G. ¹³ 992. Prepared by heating phenylene-*m*-diamine with methyl alcohol and HCl at 190° (Wurster a. Morley, B. 12, 1814). Purified by distilling with Ac_2O (Romburgh, R. T. C. 7, 2). Oil, sl. HOAc and nitric acid give sol. water. Sol. water. Hole and intro acid give C₆H(NO₂)₁N₂Me₁NO [132°]. s-Tri-nitro-benzene forms B"C₆H₄(NO₂)₂ [21°]. m-Di-nitro-benzene gives B"C₆H₄(NO₂)₂ [58°] crystallising in needles. Bromine forms oily C₁₆H₁,Br₂N₂. NaNO₂ and dilute HClAq give oily C₁H₁(NO)(NMe₂)₂, which forms a hydrochloride B'HCl, crystallising in dark-red needles (Witt, B. 18, 877).

Salts.—B"H2Cl2 2aq: hygroscopic crystals. -B"H, FeCy, aq. Roser, B. 12, 1826). Pearly plates (Wurster a.

Methyliodide B"Mel. [192°]. Crystals, v. sol. water.—B"MeHI,. Di-methylo-di-iodide B"Me, I. Plates,

v. sol. water (Hofmann, Pr. 12, 639).

p-Phenylene-tetra-methyl-diamine [1:4] $C_0H_4(NMe_2)_2$. [51°]. (260° i.V.). Prepared by heating p-phenylene-di-methyl-diamine with MeOH and HClAq at 200° (Wurster, B. 12, 526, 1803, 1827). White leaflets, sl. sol. cold water, v. sol. alcohol. Its aqueous solution becomes violet-blue on exposure to air or additior of oxidising agents. The blue substance appears only to exist as a salt, e.g. C.H. NMe_Cl_>CH. (Wurster, B. 19, 3195; 20, 256; 21, 921). It becomes colourless on further oxidation.

Reactions.-1. Bromine in HOAc forms &

blue colouring matter C10H14N2, which forms blue solutions in water and alcohol.—2. KaFeCy added to a solution of the sulphate forms C₁₀H₁₄N₂2H₄FeCy₆ crystallising in lustrous blue needles. - 3. Sodium nitrite and hydrochloric acid give C₆H₄(NMe₂)(NMeNO) [98°] C₆H₄(NO₂)(NMe₂).NMe.NO [87°]. Salts.—B"2HCl; crystals.—B"H₂PtCl₆.

B"2H2SO4-B"H4FeCy6: white needles.

Methylo-iodides B"MeI. [above 270°]. Leaflets (W.).—B"Me₂I₂. Plates (Hofmann).

Reference. - NITRO-PHENYLENE-DI-METHYL-DI-

AMINE.

PHENYLENE - DI - METHYL - DI - AMINE **SULPHONIC** ACID C₆H₃(NH₂)(NMe₂).SO₃H. Got by reduction of p-nitro-di-methyl-amidobenzene sulphonic acid (Michler a. Walder, B. 14, 2176). Large rhombohedra, v. sol. water.-BaA'2.—CaA'2: plates.

p - PHENYLENE - DI - METHYL-DI - ETHYL-**DIAMINE** $C_6H_4(NMe_2)(NEt_2)$. (264°). Formed from [1:4] C.H. (NEt.) (NH2), MeOH, and HClAq at 200° (Lippmann a. Fleissner, M. 4, 791). Liquid. Coloured blue by oxidising agents. Methylo-iodide B"2MeI. [218°]. Yields

B"Me2AuCla, B"Me2CdI4, B"Me2PtCl6,

 $B''Me_2(C_3H_2N_3O_7)_2$ [235°].

o-PHENYLENE-METHYL-ETHYLENEdiamine $C_6H_4{<}^{\rm NMe}_{\rm NH}{>}C_2H_4$. v-Methyl-quin-

oxaline tetrahydride. (274°). Formed by heating phenylene-ethylene-diamine with McI at 110° (Ris, B. 21, 381). Liquid Its square (Ris, B. 21, 381). Liquid. Its aqueous solution is coloured blue by FeCl3.

Methylo-iodide B'MeI. [above 200°]. Crystals.—B'2Me2PtCl6. Yellow needles.

DIPHENYLENE - METHYL - FURFURANE C₆H₄.C.O.OMe [124°]. Got by reducing C₆H₄.C(OH).CH₂Ac (Japp a. Klingemann, B. 21, C,H,.CO 2932).

PHENYLENE - DI - METHYL - DI - MALONIC ACIDS C₆H₄(CH₂·CH(CO₂H)₂)₂. The ethers of these acids are formed by the action of zincdust on the acids C₆H₄(CH₂·CCl(CO₂Et)₂)₂ which are formed from sodium chloro-malonic ether and w-di-bromo-xylenes (Perkin a. Kipping, C. J. 53, 16; B. 19, 437; 21, 36). The o- and methers are oily, the p- ether is crystalline [51°]. The ethers form deliquescent sodium derivatives $Na_2C_{22}H_{28}O_8$, and the \tilde{o} -compound of this formula is converted by iodine into the tetrahydride of naphthalene tetracarboxylic ether. The macid is a liquid v. sol. water, the p-acid is crystalline [195°]. The p-acid forms Ag₂A", crystalline [195°]. The p-acid forms Ag_2A'' , and when heated splits up into CO_2 and C₆H₄(CH₂.CH₂.CO₂H)₂ [223°].

PHENYLENE - METHYL - METHENYL -

AMIDINE C₆H₄< $\stackrel{\rm NMe}{\sim}$ CH. [33°]. (278°) at

730 mm. Formed from phenylene-methenylamidine, MeI, and MeOH at 100° (O. Fischer, Formed from phenylene-methenyl-B. 22, 644). Prisms.—B'HAuCl. Needles.

DIPHENYLENE-METHYL-PYRAZINE O_0H_4 .C.N.CH . [128°]. Formed from phenanthraquinone and propylene-diamine (Strache, B. 21, 2362). White needles.—B',H,PtCl,.

v-p-DIPHENYLENE - TETRA - METHYL - DI. PYRROLE C24H24N2 i.e.

CH:CMe N.C_eH₄.C_eH₄.N < CMe:CH CMe:CH[130°]. Got by heating an alcoholic solution of acetonylacetone (2 mols.) and benzidine (1 mol.) (Paal a. Schneider, B. 19, 3158). Colourless tables. Sol. alcohol, ether, benzene, and ligroin.

DIPHENYLENE-METHYL-QUINOXALINE $C_{21}H_{14}N_2 \ \textit{i.e.} \ C_6H_3Mo < N:C,C_6H_4. \ [213^{\circ}]. \ Formed$ from (1,3,4)-tolylene-diamine and phenanthraquinone in HOAc (Hinsberg, B. 17, 323). Yellow hair-like crystals, v. sol. benzene.

DIPHENYLENE - NAPHTHOQUINOXAL -

INE $C_{10}H_6 < N:C.C_6H_4$ [273°]. Formed by boiling an alcoholic solution of (1,2)-naphthylene-diamine with phenanthraquinone in HOAc (Lawson, B. 18, 2426). Small yellow crystals.

The sulphonic acid, got by mixing a hot aqueous solution of naphthylene-o-diamine sulphonic acid with phenanthraquinone dissolved in NaHSO3Aq with addition of NaOAc, forms a yellow crystalline Na salt, sol. hot water, but ppd. by alkalis (Witt, B. 19, 1719, 2791). When fused with KOH it yields the insoluble eurhodol

 $C_{12}H_8 < \stackrel{N}{N} > C_{10}H_5(OH) \begin{bmatrix} 1 \\ 2 \end{bmatrix}$, which forms a scarlet hydrochloride.

m - PHENYLENE - DI - (β) - NAPHTHYL -DIAMINE C_eH₄(NHC₁₀H₇)₂. [126³]. Formed by heating *m*-phenylene-diamine with (8)naphthol (Ruhemann, B. 14, 2654). Felted needles, sol. alcohol and ether.

p-Phenylene-di- (β) -naphthyl-diamine $C_{b}H_{4}(NHC_{10}H_{7})_{2}$. [235°]. (over 400° in vacuo). Formed by heating p-phenylene-diamine with (β) -naphthol at 200° (Rueff, B. 22, 1080). White plates, almost insol. boiling alcohol, v. sol. nitro-benzene. KNO₂ added to its solution in conc. H₂SO₄ gives a greenish-violet colour, changing to blue.—B'2C₆H₃N₃O₇. [c. 217°].

Acetyl derivative C₆H₄(NAc.C₁₆H₇)₂.

[210°]. Plates (from benzene).

Bensoyl derivative C20H18Bz2N2. [220].

Reference. - NITRO-PHENYLENE-NAPHTHYL-DI-AMINE PHENYLENE - NAPHTHYLENE - AMINE

C₁₆H₁₁N i.e. [1:2] C₆ H₄ NH. Phenyl-naphthyl-[330°]. (454°) (Schweizer, A. 264, carbazole. 195). S. (alcohol) 25 at 78°; S. (toluene) 5 at 111° (Bechi, B. 12, 1978). Occurs in crude anthracene, from which it may be got from the residues after sublimation of the anthracene (Graebe a. Knecht, B. 12, 341, 2242; A. 202, 1). Formed by passing phenyl- (β) -naphthyl-amine through a red-hot tube. Colourless plates, v. sl. sol. HOAc. Its solutions show blue fluorescence. Oxidation by K2Cr2O, and H2SO, forms the quinone $\stackrel{C_0H_4}{C_{10}H_4O_2}$ NH [307°] which crystallises from HOAc in reddish-yellow needles, and is accompanied by $\overset{C_0H_4}{C_{10}H_4O_2}\!\!>\!\!0$ which crystallises from benzene in red prisms.

Acetyl derivative C_{1e}H₁₂NO. [121°]. Nitrosamine C_{1e}H_{1e}N₂O. [240°]. Tetrahydride C_{1e}H_{1e}N. Formed

reduction with HIAq and P at 210°. Syrup.— B'HI. Long needles.

Phenylene-naphthylene-amine

[1:2] C₆ H₄ NH. [225°]. Formed by heating [2:1] C₁₀ H₆ NH with reduced copper at 240° (Kym, B. 23, 2465). Greenish-yellow plates, sl. sol. cold benzene, v. sol. hot alcohol.

PHENYLENE - NAPHTHYLENE - KETONE **OXIDE** CO< $C_{10}H_{6}$ >0. [155°]. Formed by boiling (a)-naphthyl salicylate (Graebe a. Feer, B. 19, 2612). Formed also by heating (a)-naphthol with salicylic acid and HOAc; and by distilling (1,2)-(a)-naphthol carboxylic acid with salicylic acid (Kostanecki, B. 25, 1643). Crystals, sl. sol. fit rescence. Converted by potash-fusion into o-(a)-di-oxy-phenyl-naphthyl-ketone (Phomina, A. 257, 92).

Isomeride. [140°]. Formed in like manner from (β) -naphthyl salicylate; and by heating (2,1)-(\(\beta\))-naphthol carboxylic acid with salicylic acid. Needles. Reduced by zinc-dust to Needles. Reduced by zinc-dust to $CH_2 < C_0 H_0 H_0 > 0.$ [80°].

p-PHENYLENE-DI-(β)-NAPHTHYL-DI-**METHYL-DIAMINE** $C_6H_4(NMe.C_{10}H_7)_{2*}$ [180°]. Formed from $C_6H_4(NHC_{10}H_7)_2$, MeI, and MeOH at 140° (Rueff, B. 22, 1080). Plates.

PHENYLENE - NAPHTHYLENE $\frac{C_0}{C_{10}}$ $\frac{H_4}{G_0}$ O. [178°]. Formed by distilling a mixture of phenol and (a)-naphthol with PbO (Arx, B. 13, 1726; A. 209, 141). Yellow needles, sl. sol. alcohol. H₂SO₄ forms a tetrasulphonic acid which gives Ba₂C₁₆H₆S₁O₁₃4aq. CrO₃ and HOAc form C₁₆H₈O₃ [140°].—C₁₆H₁₀O2C₆H₃N₃O_. [165°]. Isomeride C₁₆H₁₀O. [296°] (A.); [300°] (G.

K.). Formed in like manner from (β)naphthol (A.). Got also by distilling the compound C18H8O3 (from phenylene-naphthyleneamine) with zinc-dust (Graebe a. Knecht, A. 202, 15). Plates, v. sol. toluene.

References .- DI-BROMO-, DI-CHLORO-, and DI-NITRO- PHENYLENE-NAPHTHYLENE OXIDE.

PHENYLENE OXIDE C4H4O? [103°]. A product of the distillation of o-oxy-benzoyl-ooxy-benzoic acid (Märker, A. 124, 249). Silky needles (from alcohol). HNO, yields the nitrocompound C₆H₃(NO₂)Ó [150°].

Diphenylene oxide $C_{12}H_8O$ i.e. [87°]. (283°) (Galewsky, A. 264, 188).

Formation.—1. By distilling Ph.PO, over lime, MgO, or PbO (Lesimple, A. 188, 375; Hoffmeister, Z. [2] 7, 24; A. 159, 211; Kreysler, B. 18, 1720).—2. By distilling phenol (2 pts.) with PbO (3 pts.) (Behr a. Dorp, B. 7, 398; Charles (1998). Graebe, A. 174, 190).—3. By distilling Ca(OPh). (Niederhäusern, B. 15, 1120).-4. By the action of red-hot lime on di-phenylene-ketone oxide.-5. One of the products of the action of HCl on mucic acid at 150°.

Properties .- White plates (from alcohol). Converted by AcCl into CH, CO.C. H, O [81°], which yields an oxim [146°] and a phenylhydrazide [133°]. H₂SO₄ forms a deliquescent disalphonic acid, which yields BaC₁₂H_aS₂O₇ aq.

Picric acid compound C12H2OC2H3N2O2 [24°]. Yellow crystals (Goldschmiedt a. Schmidt, M. 2, 14).

References .- AMIDO-, BROMO-, TETRA-CHLORO-, DI-10DO, DINITRO- PHENYLENE OXIDES.

o-PHENYLENE - OXY - BENZAMIDINE $C_{e}H_{\bullet} < NH > C.C_{e}H_{\bullet}.OH.$ [223°]. Formed by reducing salicyl-o-nitro-aniline with tin and HCl (Mensching, A. 210, 345). Needles.—B'HCl aq.—B'₂H₂SO, 4aq. Needles, sl. sol. Aq.

o-PHENYLENE-PHENYL-DIAMINE. The sulphonic acid which gives the barium salt ${\rm BaA'_2\,2aq}$ may be got from its anilide [1:2:4] ${\rm C_8H_3(NHPh)(NH_2).SO_2NHPh}$ [157°] which is got by reduction of C₈H₈(NHPh)(NO₂).SO₂NHPh, and yields B'HCl [182°] (Fischer, B. 24, 3794).

p-Phenylene-phenyl-diamine C₆H₄(NHPh).NH₂. Amido-diphenylamine. [67°]

and [75°]. (354°). Formation. - 1. By reduction of nitrodiphenylamine (Nietzki a. Witt, B. 12, 1401). 2. From p-nitroso-diphenylamine by boiling with alcoholic potash, by reduction with tin and HCl (Ikuta, A. 243, 279), or by treatment with phenyl-hydrazine in ether (O. Fischer, B. 21, 2615).—3. Together with azophenine, by heating p-nitroso-diphenylamine with aniline and aniline hydrochloride (O. Fischer a. Hepp, B. 20, 2480). 4. By reducing C₆H₄(NHPh)N₂C₆H₄SO₃Na (Hess a. Bernthsen, B. 18, 692).

Properties.—Plates or needles, sl. sol. water, v. sol. alcohol. Melts at 75° after crystallisation from ligroin. FeCl, added to a solution of its salts gives a red colour, turned green by excess.

Yields quinone on oxidation.

Nalt.— $B'_2H_2SO_4$. Plates. Nitrosamine $C_{18}H_{14}N_4O_2$. [c. 130°]. Acetyl derivative $C_{14}H_{14}N_4O$. [158°]. Bensoyl derivative (Lellmann, B. 15,

Sulphonic acid. [1:4:2] C₆H₃(NHPh)(NH₂).SO₃H. Formed by reducing C₂H₂(NHPh)(NO₂).SO₂H (Fischer, B. 24, 3800). Yields BaA'₂ aq and an anilide [171°], which gives B'HCl [215°].

m-Phenylene-di-phenyl-diamine C₆H₄(NHPh)₂. [95°]. Formed by heating resorcin (2 mols.) with aniline (8 mols.), CaCl₂ (8 mols.) and ZnCl₂ (1 mol.) for 35 hours at 210° (Calm, B. 16, 2792). Flat needles, v. sol. ether. Oxidising agents give a green colour passing to bluish violet. Amyl nitrite and alcohol HCl form the p-nitroso- derivative C18H15N.O, crystallising in brown-red prisms and yielding B"2HCl (Fischer a. Hepp, A. 255, 145). B"2HCl:

needles decomposed by water. Acetyl derivative C.H. (NPhAc)2. [163°]. C.H.(NPhBz)2. Benzoylderivative [184°].

Nitrosamine C,H,(NPh.NO)2. Yellow needles, forming a violet solution in H,SO,

p-Phenylene-di-phenyl-diamine C.H.(NHPh), [132°] (Bandrowski, M. 8, 475); [141°] (Limpricht, B. 22, 2910); [145°] (O. Fischer, B. 21, 2615); [152°] (Calm, B. 16, 2803). Formed by heating hydroquinone (5 753.) with aniline (17 pts.), CaCl, (20 g.), and ZnCl, (5 g.) for 18 hours at 210°. Formed also by the action of phenyl-hydrazine on nitroso-diphenylamine in alcohol, and from the hydrochloride of amido-salicylic acid and aniline. Plates, v. sol. hot benzene. HNO, added to its solution in H.SO, gives a blue liquid, changing to red.-B"2HCl: needles, decomposed by water.

Acetyl derivative C.H. (NPhAc)2. [192°]. C.H.(NPhBz)2 Benzoylderivative [219°].

Nitrosamine C₁₆H₁₄N₄O₂. [c. 120°].

Phenylene-di-phenyl-diamine

 $C_6H_1(NH_2).NPh_2$ Formed by reducing nitrotri-phenyl-amine (Herz, B. 23, 2537). Crystalline.-B'HCl: silvery needles.

C₆H₄(NHAc).NPh₂. Acetyl derivative

[197°]

o-PHENYLENE-PHENYL-GUANIDINE

 $C_{\bullet}H_{\bullet} < NH > C:NPh.$ [190°]. (440°-450°).

Formed by heating phenylene o-diamine with di-phenyl-cyanamide at 215° (Keller, B. 24, 2499). White needles (from benzene) or prisms (from alcohol). Phenyl-cyanate forms the compound NPh CO N2(CNPh)C.H. [266°]. -

B'HCl.-B'₂H₂PtCl₆.-B'H₂SO₄: long needles. Mono-acetyl derivative. [160°]. Di-benzoyl derivative. [1710]. Needles. PHENYLENE - DI - PHENYL - DÍHYDRAZ-Hexahydride $C_6H_{10}(NH.NHPh)_2$ [148°]. Formed by reducing the di-phenyl-dihydrazide of quinone with sodium and alcohol (Baeyer a. Noyes, B. 22, 2175). Crystalline, sl.

sol. alcohol.—B'H,C,O, aq.

m-PHENYLENE DI-PHENYL DIKETONE $C_6H_4(CO.C_6H_3)_2$. Isophthalophenone. [95°] (M.); [100°] (A.). (above 360°). Formed from isophthalyl chloride, benzene, and AlCl₃ (Ador, Bl. [2] 33, 56; B. 13, 320; Münchmeyer, B. 19. 1848). Plates (from alcohol). Yields a monooxim [201°] (Nölting a. Kohn, B. 19, 146), a dioxim [70°-75°], and two di-nitro-derivatives [260°] and [c. 100°].

p-Phenylene diphenyl diketone

[160]. Formed from tere- $C_6\hat{H}_4(CO.C_6H_5)_2$. phthalic chloride, benzene, and AlCl₃ (M.). Yields a dioxim [235°].

Diphenylene di-phenyl diketone $C_{12}H_{3}(CO.C_{6}H_{5})_{2}$. [218°]. Got from diphenyl, BzCl, and AlCl₃ (Wolf, B. 14, 2031). Crystals. DIPHENYLENE-PHENYL-METHANE

 $C_{19}H_{14}$ i.e. $C_{cH}^{eH_4}$ CHPh. [145.5°]. Formed by the action of P.O. or AlCl. on a mixture of benzene and diphenylene-carbinol; and also by heating CPh_sCl at 200° (Hemilian, B. 11, 202, 837; Bl. [2] 34, 325). Formed also in small quantity in the distillation of calcium benzoate or phthalate (Kekulé, B. 5, 910; Miller, B. 12, 1489). Needles, sl. sol. ether. Yields o-benzoylbenzoic acid on oxidation.

Reference.—DI-BROMO-DI-PHENYLENE-PHENYL-METHANE.

ν - m - PHENYLENE - DI - PHENYL - DI -METHYL-DI-PYRROLE DI-CARBOXYLIC-ETHER C₃₄H₉₂O₄N₂ i.e.

CO.Et.C:CMe N.C.H.N CMe:C.CO.Et

[185°]. Obtained by mixing together in acetic acid solution acetophenone-aceto-acetic ether (2 mols.) and m-phenylene diamine (1 mol.), and allowing the mixture to stand a long time

(Paal a. Schneider, B. 19, 3161). Slender white needles. V. sol. ordinary solvents, except Aq.

v-p-Diphenylene-di-phenyl-di-methyl-dipyrrole di-carboxylic-ether C40H36O4N2 i.e. CO₂Et.C:CMe HC:CPh N.C₆H₄.O₆H₄.N CPh:CH

[179]. Obtained by mixing together in acetic acid solution acetophenone-aceto-acetic ether (2 mols.) and benzidine (1 mol.), and allowing the mixture to stand for a long time (P. a. S.). Hair-like needles. Sol. alcohol and acetic acid, v. sol. ether, chloroform, and petroleum-spirit.

p - PHENYLENE - TETRA - PHENYL - DI -PYŘROLE DICARBOXYLIC ACID

 $CO_2H.C:CPh$ HC:CPh $N.C_6H_4.N$ $CPh:C.CO_2H$ CPh:CH 300°]. Formed by saponification of its ether [250°] which is got from phenacyl-benzeylacetic ether and phenylene p-diamine (Paal a. Braikoff, B. 22, 3095). Small grains.

m-PHENYLENE DI-PHENYL SULPHONE C₆H₄(SO₂Ph)₂. [191°]. Formed by heating benzene with P₂O₅ and benzene m-disulphonic [191°]. acid or di-phenyl sulphone m-sulphonic acid (Otto, B. 19, 2421; 20, 185). Minute needles (from HOAc). Alcoholic potash at 170° forms $O(C_6H_4.SO_2Ph)_2$ [70°] and $C_8H_5.SO_2K$.

Di-phenylene di-phenyl trisulphone $SO_2(C_6H_4.SO_2C_6H_b)_2$. [193°]. Formed from SO₂(C₆H₄.SO₇H)₂, benzene, and P₂O₈ at 200° (Otto a. Rössing, B. 19, 3127).

PHENYLENE-DI-PHENYL-DI-THIO-DI-UREA C₀H₄(NH.CS.NHPh)₂. Formed from phenylene-diamine and phenyl thiocarbimide in alcohol (Lellman a. Wurthner, A. 221, 28; 228, 200). The o- and p- compounds are decomposed by heat into di-phenyl-thio-urea and phenylene-thio-urea; the m- compound melts at 161°, and the o- isomeride at about 290°. They are insol. ether.

m-PHENYLENE-DI-PHENYL-UREA C_bH₄(NH.CO.NHPh)₂. Formed from m-phenyl-

ene-diamine and phenyl cyanate (Kuhn, B. 18, 1478). Amorphous.

PHENYLENE-PROPENYL-DIAMINE

 $C_{e}H_{4} < N^{H} > CEt$. [169°]. (above 360°). by boiling o-phenylene-diamine with propionic acid (Wundt, B. 11, 829), and by reducing propionyl-o-nitro-aniline with tin and HOAc (Smith, Am. 6, 172). Plates (from water).—B'HC'.—B'HgCl₂.—B'₂H₂PtCl₆ 2aq.—B'₂H₂Cr₂O₇.

PHENYLENE-DI-PROPIONIC ACIDS $C_6H_1(CH_2.CH_2.CO_2H)_2$. Formed by heating $C_{2}H_{2}(CH_{2}\cdot CH(CO_{2}H)_{2})_{2}$ (Perkin a. Kipping, C. J. 53, 8, 32; B. 21, 27, 40). The o-acid [162°] forms Ag₂A". The m-acid [147°] forms Me₂A" [51°] and Et₂A" (250° at 60 mm.). T [224°] forms Ag_2A " and Me_2A " [115°]. The p- acid

 $p extsf{-}\mathsf{PHENYLENE-PROPYL-DIAMINE}$ $C_6\hat{H}_4(NH_2)(NHPr)$. (281°). Got by reducing p-nitroso-propyl-aniline with SnCl. (Wacker, A. 243, 294). Plates.—B"2HCl. Plates, v. e. sol. Aq.

o-PHENYLENE-PROPYLENE-DIAMINE $C_{e}H_{4} < NH > C_{s}H_{e}$. [72°]. (283°). Formed by heating pyrocatechin with propylene-diamine at 200° (Ris, B. 21, 382). Plates (from ligroin).

-B',3HCl.-B',2C,H,N,O,. Thin needles.

DIPHENYLENE-PYRAZINE. Dihydride | alcohol at 100° (Gabriel, B. 10, 184). Needles, $\textbf{C}_{16}\textbf{H}_{12}\textbf{N}_2 ~~\textit{i.e.} ~~ \begin{matrix} \textbf{C}_6\textbf{H}_1, \textbf{C.N.CH}_2 \\ \textbf{C}_8\textbf{H}_4, \textbf{C.N.CH}_2 \end{matrix}.$ Formed from phenanthraquinone and ethylene-diamine (Mason, B. 19, 112; 20, 267). _B',H,PCl

(a)-PHENYLENE-PYRIDYL-KETONE

C₆H₄CO.C.CH:CH C-N:CH. [141°]. Formed by distilling the dicarboxylic acid

 $C_7H_4O < \stackrel{C.C(CO,H):CH}{C.N} : \stackrel{CCO_2H)}{CCO_2H} >$ [284°], which is formed by oxidising (a)-styryl-(β)-naphthoquin-oline carboxylic acid (Doebner a. Peters, B. 23, 1241). Yields a picric acid compound The dicarboxylic acid yields Ag₂A".

(β)-Phenylene-pyridyl-ketone

C₆H₄<0.C.N:CH [129°]. Formed in like C.CH:CH* manner from the corresponding dicarboxylic acid [264°] which is got by oxidising (a)-styryl-(a)-naphthoquinoline carboxylic acid with KMnO. Needles. Yields B',H.,PtCl, 2a., The dicarboxylic acid forms Ag,A" as a pp.
PHENYLENE - QUINOLYLENE KETONE

OXIDE $C_{1e}H_{\varphi}N\mathring{O}_{2}$ i.e. $O < \overset{C_{e}H_{\varphi}}{C_{u}H_{v}N} > CO$. Phenoquinoxanthone. [188°]. Formed by distilling p-oxy-quinoline with salicylic acid. (from alcohol).-B'HCl: needles.

DIPHENÝLENE-QUINOXALINE C20H12N2 i.e. $\frac{C_0H_1.Q.N}{C_0H_4.Q.N} > C_0H_4.$ [217°]. Got from phenyleneo-diamine and phenanthraquinone (Hinsberg, A. 237, 340). V. sl. sol. alcohol.

DIPHENYLENE-STYRYL-OXAZOLE $C_{cH_{+},C,N} > C.CH:CHPh.$ [172°]. Formed from phenanthraquinone, cinnamic aldehyde, and alcoholic NH, at 100° (Wadsworth, C. J. 57, 11). Yellow needles.

p-PHENYLENE DISULPHIDE $\mathbf{C}_{a}\mathbf{H}_{a}\mathbf{S}_{c}$ Formed by oxidising dithiohydroquinone with an alkaline ferricyanide solution (Leuckart, J. pr. [2] 41, 206). Chars without melting at 300°, insol. ordinary solvents. Is reduced by fusing with potash to dithiohydroquinone.

Diphenylene sulphide C.H. S. [97°]. (333° i.V.). Formed by passing Ph2S through a redhot tube (Stenhouse, A. 156, 332; Graebe, A. 174, 185). Needles, m. sol. alcohol. Yields on dxidation the sulphone C₁₂H₈SO₂ [230°].

 $C_6H_4 < S > C_6H_1$ Diphenylene - disulphide [154°]. (366°). A product of the distillation of sodium benzene sulphonate (Stenhouse, Pr. 17, 62; A. 149, 252). Formed also by heating phenol with P2S, (Graebe, A. 174, 185; 179, 178), by heating benzene with S and AlCl, at 80° (Friedel a. Crafts, A. Ch. [6] 1, 530; 14, 439), and by heating $C_6H_4 < N > N$ (Jacobson a. Elley, B. 22, 910). Prisms (from CS2). Conc. H.SO₄ forms a purple solution. CrO₃ in HOAc gives the sulphone C₁₂H₈S₂O₄ [325°] and the sulphoxide C₁₂H₈S₂O₂ [241°], sl. sol. cold benzene. Br forms C₁₂H₄S₂Br₄ crystallising from CS in small black rejents CS2 in small black prisms.

v. e. sol. hot alcohol. Yields a nitro- compound $C_{\rm e}H_{2}({
m NO}_{2})({
m SCN})_{2}$ [150°].

DIPHENYLENE SULPHONE v. DIPHENYL-ENE SULPHIDE.

DI-PHENYLENE SULPHONE KETONE

 $C_eH_4 < SO_2 > C_eH_4$. [185°]. Formed by heating benzophenone with fuming H.SO. (Beckmann, B. 6, 1112; 8, 992), and by oxidising $C_sH_4 < S > C_sH_4$ (Graebe a. Schultess, A. 263, Yellowish needles, v. sol. alcohol.

PHENYLENE DITHIOCARBAMIC ETHERS C, H, (NH.CO.SEt),. Formed by boiling from phenylene-thiocarbimide with alcohol (Billeter a. Steiner, B. 20, 230). The m-compound [116°] and the p- compound [197°] are crystalline and yield Ag_2A'' as white pps.

PHENYLENE-DI-THIO-CARBIMIDE

C₆H₄(NCS)₂. Formed from phenylene-diamine and CSCl, (Billeter a. Steiner, B. 20, 231). The o- compound [59°], m- compound [53°] (250°), and p- compound [130°] all crystallise in needles.

 \hat{m} -PHENYLENE - DI - THIO - DIGLYCOLLIC ACID C₆H₄(SCH₂.CO₂H)₂. [127°]. Formed from di-thio-resorcin and ClCH₂CO₂H (Gabriel, B. 12, 1639). Microcrystalline powder.

Diphenylene-di-thio-di-glycollic acid $C_{12}H_s(S.CH_2.CO.H)$. [252°]. Formed from $C_{12}H_s(SH)_2$, NaOHAq, and chloro-acetic acid (Gabriel, B. 13, 390).

o-PHENYLENE-THIO-UREA

 $C_6H_4 < NH > CS$. [298°]. Formed by heating o-phenylene-diamine hydrochloride with am monium sulphocyanide and water at 130 (Lellmann, B. 15, 2146; A. 221, 9). Formed also from o-phenylene-diamine and CSCl, (Billeter a. Steiner, B. 20, 231). Plates (from dilute alcohol, sl. sol. water). The crystalline m-isomeride is got by heating m-phenylene-diamine with CS, and some alcohol at 150° (Gucci, G. 17, 524). The p-isomeride melts at 271° .

Phenylene-di-thio-di-ureas C,H,(NH.CS.NH2)2. $m \cdot [215^{\circ}].$ Formed by heating phenylene-diamine hydrochloride with ammonium sulphocyanide (Lellmann, A. 221, 11; B. 15, 2840). The p- compound is also formed from p-phenylene dithiocarbimide and alcoholic NH_3 (Billeter a. Steiner, B. 20, 230).

 $C_0H_1.NH > CS.$ Diphenylene - thio - urea [238°]. Formed by heating di-op-amido-diphenyl with alcohol and CS2 (Reuland, C. J. 58, 167; B. 22, 3014)

o-PHENYLENE-TOLENYL-AMIDINE

 $C_6H_4Me.C \leqslant {NH \atop N} > C_6H_4$. [268°]. Formed from o-phenylene-diamine and p-toluic chloride, and also by reduction of p-toluyl-o-nitro-aniline (Hübner, A. 210, 328; cf. Brückner, A. 205, 115). Prisms (from alcohol).—B'HCl.—B'2H2PtClg.— B'HNO₃, -B'₂H₂SO₄: slender needles.
PHENYLENE-DI-p-TOLYL-m- PIAMINE

 $\mathbf{C}_{6}\mathbf{H}_{4}(\mathbf{NH}.\mathbf{C}_{6}\mathbf{H}_{4}\mathbf{Me})_{2}$ C₂₀H₂₀N₂ i.e. Formed by heating resorcin with p-toluidine, m-PHENYLENE SULPHOCYANIDE

Complex Sulphocyanide

ZnCl₂, and CaCl₂ (Hatschek a. Zega, J. pr. [2]

Complex Sulphocyanide

ZnCl₂, and CaCl₂ (Hatschek a. Zega, J. pr. [2]

33, 218). Needles. Yields a di-nitrosamine lead salt of di-thio-resorcin with ICy and $C_{20}H_{18}N_4U_2$ [c. 150°], a di-acetyl derivative $\substack{C_{.0}H_{18}Ac_2N_2\\C_{20}H_{18}Bz_2N_2}}$ [176°], and a di-benzoyl derivative crystalline

powder, decomposed by water.

Phenylene-di-o-tolyl-p-diamine [135°]. (420°). Formed by heating hydroquinone with o-toluidine and CaCl₂ at 290° (Philip, J. pp. [2] 34,65). Plates (from HOAc). Yields a dinitrosamine [140°]. Yields C₂₀H₁₈Bz₂N₂ [235°] and C₂₀H₁₈Az₂N₂ [189°].—B"2HCl: needles.

Phenylene-di-p-tolyl-p-diamine. [182°]. Formed by heating hydroquinone or phenyl-p-amido-phenol with p-toluidine and ZnCl₂ at 220° (H. a. Z.); Calm, B. 16, 2810). Plates. Fields a di-nitrosamine [152°], a di-acetyl derivative [173°], and a di-benzoyl derivative [222°].—B"2HCl.

PHENYLENE-TOLYLENE-KETONE OXIDE

 $C_eH_i < {{\rm CO} \atop {\rm O}} > C_eH_3$ Me. Methyl-xanthone. [105°]. Formed by boiling p-tolyl salicylate (Graebe a. Feer, B. 19, 2612; cf. Seifert, J. pr.[2] 31, 478).

Isomeride $C_bH_4 < {{O}\atop{CO}} > C_bH_4Mc \left[{{3}\atop{4}} 1\right]$. [176°]. Made by heating $C_bH_3Me(OH)CO_2Ph$ [49°] (Weber, B. 25, 1745). Crystals (from alcohol).

o-PHENYLENE-p-TOLYL-GUANIDINE

C_bH_i $\stackrel{\text{NH}}{\sim}$ C:N.C_bH_iMe. [209°]. Formed by heating o-phenylene-diamine with di-p-tolyl-cyanamide at 210° (Keller, B. 24, 2509). Tablos. Yields a mono-acetyl derivative [152°], a dibenzoyl derivative [191°], and a nitrosamine C₁₄H₁₂(NO)N₃ [150°-160°]. Phenyl cyanate forms C₆H₃N $\stackrel{\text{CO}}{\sim}$ N₂(C₆H₄):C:NC₆H₄Me [254°] crystallising from benzene in small needles. Di-p-tolyl-cyanamide at 210° forms the compound (NHC,H₇)₂C:N₂(C₆H₄):C:N.C₆H₄Me [188°]. Salts.—B'HCl.—B'₂H₂PtCl₆,—B'₂H₂SO₄.

i.e. C_0H_1 CH.C. H.Me. [128°]. Formed by the action of P_2O_5 on a mixture of diphenylene-carbinol and toluene, or of AlCl. on diphenylene-carbinyl chloride dissolved in toluene (Hemilian, B. 11, 203; Bl. [2] 34, 325). Silky needles.

PHENYLENE-DI-p-TOLYL-DI-METHYL-m-DIAMINE $C_0H_+(NMe.C_0H_+(Me)_2$. (c. 400°). Got by heating phenylene-di-p-tolyl-m-diamine with MeI and KOH at 150° (Hatschek a. Zega, J. pr. [2] 33, 223). Liquid smelling like geraniums.

The isomeric phenyler e-di-o-tolyl-di-methyl-p-diamine (385°-390°) and phenylene-di-p-tolyl-di-methyl-diamine [153°] may be prepared in like manner (Philip, J. pr. [2] 34, 57; H. a. Z.).

o-Phenylene-urea C_uH , $<_{NH}^{NH}>CO$.

[308°]. Formed by heating o-amido-phenyl-carbamic ether at 90° (Rudolph, B. 12, 1296), by heating o-amido-di-phenyl-urea (Lellmann a. Würthner, A. 228, 220), by heating its ethyl derivative with HCl (Sandmeyer, B. 19, 2654), and by the action of COCl, in toluene on a solution of o-phenylene-diamine hydrochloride at 160° (Hartmann, B. 23, 1046). Leaflets, sl. sol. water.—B'HCl. Decomposed by water.

Ethyl derivative $C_eH_e < \stackrel{NH}{N} > C.OEt$. [160°]. Formed from o-phenylene-diamine and NH:C(OEt). Plates.

m-Phenylene-urea. [above 300°]. Formed by similar methods (Michler a. Zimmermann, B. 14, 2177; L. a. W.). Insoluble powder.

p-Phenylene-urea. [above 320°]. Got by heating p-amido-di-phenyl-urea (L. a. W.).

Isomeride, [130°]. Got by boiling an alcoholic solution of oxy-phenyl-thio-urea with HgO (Bendix, B. 11, 2264). Tables, sol. water.

Phenylene - di - ureas C₆H₄(NH.CO.NH₄)₂. o- [290°]. m- [above 300°]. Formed from phenylene-diamine hydrochloride and potassum cyanate in cold aqueous solutions (Warder, B. 8, 1180; Lellmann, A. 221, 13; B. 16, 592).

The p-compound can be heated to carbonisa

tion without melting.

References. —Amido- and Nitro- phenylene-

PHENYL-ENNOIC ACID.

Nitrile C₆H₅.CH(C₇H₁₅).CN. (327°). Form 2d from CH₂Ph.CN, heptyl iodide, and NaOH (Rossolymo, B. 22, 1237). Yellow oil.

DI-PHENYL-ENNYL TRICYANIDE

C₀H₁₉C₃N₄Ph₂. [c. 38°]. (c. 293° at 15 mm.). Formed from decoic chloride, benzonitrile, and AlCl₄ (Krafft a. Koenig, B. 23, 2384).

PHENYL-ENNYL-THIO-UREA C₁₆H₂₆N₂S i.e. NHPh.CS.NHC₆H₁₈. [60°]. Formed from ennylthiocarbimide and aniline (Freund a. Schönfeld, B. 24, 3359). Tables, v. sol. alcohol and ligroïn. PHENYL-ENNYL-UREA C₁₆H₂₆N₂O i.c.

NHPh.CO.NHC₀H₁₀. [63°]. Formed from phenyl cyanate and aniline in alcohol (Freund a. Schönfeld, B. 24, 3358). Long prisms.

s-DI-PHENYL-ETHANE $C_{11}\Pi_{11}$ i.e. CH.Ph.CH.Ph. Dibenzyl. Mol. w. 182. [53°]. (277°). S.V.S. 174·2 (Schiff, A. 223, 261). H.F. -31,200. H.C.v. 1,828,300. H.C.p. 1,830,200 (Berthelot a. Vieille, Bl. [2] 47,866).

Formation.—1. By the action of Na on benzyl chloride (Cannizzaro a. Rossi, A. 121, 250; Fittig, A. 137, 257).—2. From ethylene chloride, benzene, and AlCl₃ (Silva, C. R. 89, 606; A. Ph. S. 18, 345).—3. From CHBr:CHBr, benzene, and AlCl₃ (Anschütz, A. 235, 155).—4. From acetylene, benzene, and AlCl₃ (Varet a. Vienne, Bl. [2] 47, 919).—5. By heating benzyl chloride with copper powder (Onufrovitch, B. 17, 833).—6. A product of the action of Na on o-bromo-benzyl-bromide (Jackson a. White, Am. 2, 390).—7. By adding sodium (70 g.) to an alcoholic solution of phenyl-cinnamonitrile (Freund a. Remse, B. 23, 2859).

Properties.—Long colourless needles, sol. alcohol, ether, and CS₂.

Reactions.—1. Yields toluene and di-phenylethylene when passed through a red-hot tube (Otto, Z. [2] 6, 22; A. 154, 176; Barbier, C. R. 78, 1769).—2. Chlorine passed into fused s-di-phenyl-ethane forms first CHPh:CHPh and then C₆H₄Cl.CH:CH.C₆H₄Cl (Kade, J. pr. [2] 19, 466). In presence of I, chlorine forms p-di-chloro-di-phenyl-ethane in the cold. Exhaustive chlorination yields C₂Cl₆ and C₆Cl₆ (Merz a. Weith, B. 16, 2877).—3. H₂SO₄ forms a disulphonic acid C₁₄H₁₂(SO₃H)₂ 5aq, which yields K.A'' 2aq, BaA'' ½aq, and PbA'' aq (Kade, B. 6, 953). A tetra-sulphonic acid, C₁₄H₁₀(SO₃H)₄ 3aq, is also formed.

 $u\text{-Di-phenyl-ethane CH}_{\rm s}.\text{CHPh}_{\rm 2}.~(270^\circ)$ (O.); $(286^\circ)~(\text{H.}).$

Formation.—1. From CCl, CHPh, alcohol,

and Na (Goldschmiedt, B. 6, 1501).--2. From CH2Ph.CH2Br, benzene, and zinc-dust (Radziszewski, B. 7, 140).—3. From paraldehyde, benzene, and H₂SO₄ (Baeyer, B. 7, 1190).—4. Together with ethyl-benzene and di-methyl-anthracene dihydride by heating ethylidene chloride with benzene and AlCl, (Silva, Bl. [2] 41, 448; Anschütz, B. 17, 165).—5. By distilling its dicarboxylic acid (Haiss, B. 15, 1481).

Properties. - Oil, with blue fluorescence. Solidifies in a freezing mixture. Yields benzophenone on oxidation. Fuming HNO, added to its solution in HOAc forms benzophenone, CPh₂(OH).CH₂.O.NO₂ [107°], di-phenyl-vinyl nitrite [87°], and a body [148°] which yields diphenyl-acetonitrile on reduction (Anschutz a. Romig, A. 233, 329).

Tri-phenyl-ethane CH.Ph.CHPh. (above 360°). Formed from CH.Cl.CHCl.OEt, benz-Tri-phenyl-ethane CH2Ph.CHPh2. ene, and AlCl₃ (Waas, B. 15, 1128). with violet fluorescence. Insol. cold alcohol.

s-Tetra-phenyl-ethane C H2. i.e. CHPh₂. CHPh₂. Mol. w. 334. [210°]. S. (benzene) 14 at 80°. S. (95 p.c. alcohol) 8 on boil-

Formation.-1. By distilling benzoyl- and succinyl- di-phenyl-carbinol and by distilling diphenyl-carbinol with succinic acid (Linnemann, A. 133, 24).—2. By distilling benzophenone with zinc-dust (Staedel, B. 6, 1401).-3. By reducing benzpinacone CPh2(OH).CPh2(OH) with HI and P (Graebe, B. 8, 1055).—4. From di-phenylcarbinol, glacial HOAc, conc. HClAq, and zinc (Zagumenny, A. 184, 176; Bl. [2] 34, 329).--5. By reducing (3)-benzpinacolin CPh, CO.C. H. with HI and P (Zincke a. Thörner, B. 11, 67).— 6. By boiling (CHPh₂)₂S₂ with alcohol and copper powder (Engler, B. 11, 926).—7. By reducing CPh₂:CPh₂ (Friedel a. Balsohn, Bl. [2] 33, 338).—8. From CHPh2Cl and Na (Engler). 9. By the action of benzene and AlCl. on CBr₃.CH₂Br, on CHBr₂.CHBr₂, on CPhHBr.CHBr₂, on CPhBr2.CPhBr2, and on CHPhBr.CHPhBr, the yield in the last case being excellent (Anschütz, A. 235, 196).—10. By distilling CPh₂HCl (Anschütz, A. 235, 220).

Properties.—Needles (by sublimation), sl. sol. alcohol. Crystallises from benzene with C.H. Yields a crystalline tetra-nitro- derivative, a crystalline tetra-sulphonic acid, which gives Ba_2A^{1v} and $C_{2s}H_{1s}(OH)_4$ [248°].

u-Tetra-phenyl-ethane CPh₃.CH₂Ph. [140°]. Formed from CPh₃K and benzyl chloride (Hanriot, C. R. 108, 1119).

References .- AMIDO-, BROMO-, BROMO-AMIDO-, DI-BROMO-DI-NITRO-, CHLORO-, PENTA-CHLORO-DI-NITEO-, NITEO-, and OXY-, PHENYL-ETHANES.

PHENYL-ETHANE DICARBOXYLIC ACID v. CARBOXY-PHENYL-PROPIONIC ACID and PHENYL-SUCCINIC ACID.

Phenyl-ethane tricarboxylic acid

CHPh(CO₂H).CH(CO₂H)₂. [171°]. Got by sa ponifying its ether, which is made by the action of a-chloro- or a-bromo- phenyl-acetic ether on sodium malonic ether (Spiegel, A. 219, 31; Alexander, A. 258, 71). Small tablets, v. sol. hot water. Decomposed on fusion into CO₂ and phenyl-succinic acid. Salts.—Ca,A''', 10aq. -Ca,A''', 5aq: ppd. from aqueous solution by alco-hol.—Ag,A''': crystalline pp.

Ethyl ether Et.A". [46°]. 10 mm.). Needles (from dilute alcohol).

Di-phenyl-ethane o-carboxylic acid C15H14O2 i.e. C.H., CH., CH., C.H., CO.H. [131°]. Formed by the action of HI and P at 200° on isobenzylidene-phthalide, and on deoxybenzom carboxylic acid (Gabriel, B. 11, 1019; 18, 2446). Tablets (from dilute alcohol). - AgA': pp.

Isomerides v. DI-PHENYL-PROPIONIC ACID.

Di-phenyl-ethane di-o-carboxylic acid C₁₀H₁, O₄ i.e. CO.H.C₀H₄,CH₂,CH₂,C₀H₄,CO₂H. [186°] (H.); [229°] (D.).

Formation.—1. By heating diphthalyl with

HI and P (Graebe, B. 8, 1055).-2. By the action of HI and phosphorus on the acid $CO < C_0^H > CH.CH^-C'H^4.CO^H$ (Wislicenus, B. 17, 2181; Hasselbach, A. 243, 254), and on

diphthalylic acid (Dobreff, A. 239, 66). Properties. - Small needles, v. sol. alcohol and dilute HOAc. KMnO, in alkaline solution forms diphthalylic acid [263°]. Distillation

over soda-lime forms s-di-phenyl-ethylene. Salts.—(NH₁).A" (dried at 100°).—CaA' (at 100°). — BaA''. — $Cu_2A''O$. — $Pb_2A''O$. — Zu,A"O.—AgHA".

Methyl ether Me₂A". [101^c].

Ethyl ether Et₂A". [71°]. Conby alcoholic NH₃ into the amic C₂H₄(C₆H₄,CO₂Et)(C₆H₄,CONH₂) [c. 67°]. Converted

Di-phenyl-ethane dicarboxylic acid $C_{\theta}H_{3}.CH_{2}.CH(CO_{2}H).C_{\theta}H_{1}.CO_{2}H$ [1:2]. (above 300°). Formed by heating the nitrile with conc. HCl at 220° (Eichelbaum, B. 21, 2682). Small prisms, sol. alcohol, insol. ether.

Nitrile C₆H₄,CH₂,CH(CN),C₆H₄,CN. [110°]. ove 300°). Formed from benzyl chloride (above 300°). Formed from and [1:2] C₈H₁(CN).CH₂.CN. Plates, insol. water, alkalis, and acids.

Amide. [224°]. Formed by the action of conc. H₂SO, on the nitrile. Converted by conc. HCl (S.G. 1·19) at 100° into the imide

Di-phenyl-ethane di-carboxylic acid CH₃.CH(C₈H₄.CO₂H)₂. [275°]. Got by heating the tri-carboxylic acid (Hass, B. 15, 1481). Long needles. May be sublimed.—CaA".

Isomeride v. DI-PHENYL-SUCCINIC ACID.

Di-phenyl-ethane tri-carboxylic acid $CO_2H.CMe(C_eH_4.CO_2H)_2$. [255°]. Formed by oxidation of di-a-tolyl-propionic acid KMnO₄ (Haiss, B. 15, 1479). Sol. alcohol and ether.—Ag₂HA'''.—Ag₃A'''.

Di-phenyl-ethane tri-carboxylic acid C₁₇H₁₄O₆ i.e. CHPh(CO₂H).CPh(CO₂H)₂.

Amide CHPh(CO2H).CPh(CO2H).CONH2. [190°]. Got from CHPh(CO,Et).CPh(CN).CO,Et [105°] which is got by heating a-chloro-phenylacetic ether with alcoholic KCy on a water-bath (Poppe, B. 23, 114).

Mono-nitrile CO₂H CHPh.CPh(CN).CO₂H. The ethers Me₂A" [101°] and Et₂A" (v. supra) are got from a-chloro-phenyl-acetic acid and KC in MeC.H or EtOH. Conc. H₂SO₄ converts Et₂A" into CO Et.CHPh.CPh(CO₂Et).CONH₂ [157°] orystallising in aggregates of needles.

References .- NITRO- and OXY- DI-PHENYL-ETHANE CARBOXYLIC ACID.

PHENYL-ETHENYL-AMIDINE v. PHENYL-

PHENYL - ETHENYL - AMIDO - PHENYL -MERCAPTAN C,4H,1NS i.e.

 $C_{\bullet}H_{\bullet} <_{S}^{N} > C.CH_{2}Ph.$ Formed by heating phenyl-acetic chloride with o-amido-phenyl mercaptan (Hofmann, B. 13, 1234). Oil, sol. alcohol and ether. On fusion with potash it yields phenyl-acetic acid and amido-phenyl mercaptan.—B'HCl.—B'2H2PtCl6 5aq: needles.

PHENYL-ETHENYL-AMIDÖXIM C8H10N2O i.e. C₆H₅.CH₂.C(NOH).NH₂. Phenyl-acetamidoxim. [67°]. Got by heating phenyl-aceto-nitrile with a solution of hydroxylamine in dilute alcohol (Knudson, B. 18, 1068, 2482). Thin prisms, v. sol. water. Converted by phenyl cyanate into C₆H₂.CH₂.C(NH.CO.NHPh):NOH [123°].—B'HCl. [155°]. White prisms.

Acetyl derivative. [124°]. Plates.

Benzoyl derivative C,H,C(NOBz).NH2.

[144°]. Prisms, v. sol. alcohol.

Ethyl ether C,H,C(NOEt).NH2. [58°]. Converted by potassium cyans CH,Ph.C(NOEt).NH.CO.NHPh [148°]. into cyanate

Benzyl ether. [55°]. Prisms.

Phenyl-ethenyl-amidoxim

CH₈.C(NOH).NHPh. [121°]. Formed by heatthioacetic anilide with hydroxylamine solution at 100° (Müller, B. 22, 2408; cf. Nordmann, B. 17, 2746). Satiny plates, v. sol. alcohol. FeCl₃ gives a violet colour, changing to olive green and, on heating, to red.—B'HCl: needles.

-B'₂H₂PtCl₈: yellow needles.

Benzoyl derivative C₂H₄(NOBz).NHPh. [110°]. White needles (from dilute alcohol).

PHENYL-ETHENYL-AZOXIM v. AZOXIMS. PHENYL-ETHENYL-DI-ETHYL-TRI-SUL-PHONE CH₃.C(SO₂Et)₂(SO₂Ph). [109°]. Got by oxidation of CH_3 . $C(SPh)(SO_2Et)_2$ (Laves, B.

25, 364). Needles, v. sol. alcohol.
TRI-PHENYL ETHENYL TRISULPHONE CH₃.C(SO₂Ph)₃. [182°]. Formed by the action of alcoholic NaOH, and MeI on CH(SO2Ph), Got also by oxidation of CH, C(SPh), by KMnO₄ (Laves, B. 25, 352). Needles, v. sol. CHCl₃.

DI-PHENYL-ETHENYL-DI-UREA

NHPh.CO.N:CMe.NH.CO.NHPh. [169°]. Formed by adding aqueous NaOH (2 mols.) to an aqueous solution of acetamidine hydrochloride (2 mols.) shaken with phenyl cyanate (1 mol.) (Pinner, B. 23, 2923). Needles, m. sol. alcohol. Converted by boiling dilute (50 p.c.) acetic acid into acetyl-phenyl-urea [183°].

PHENYL ETHER v. DI-PHENYL OXIDE.

a-PHENYL-ETHYL ALCOHOL C,H,O i.e. CH₂.CHPh.OH. Mol. w. 122. (203°). 1.013. Formed from CH₃.CHBr.C₅H₅ by by successive treatment with AgOAc and NaOH (Radziszewski, B. 7, 141; Berthelot, Z. 1868, 589). Got also by reducing acetophenone with sodiumamalgam (Emmerling a. Engler, B. 6, 1005). Yields an acetyl derivative C, H, OAc (217°-220° which yields styrene on treatment with alcoholic

Ethylether C₈H₉OEt. (186°). S.G. 22 .931. Formed from CH₃.CHBr.C₆H₅ and alcoholic

NH, at 100° (Thorpe, Z. 1871, 131). β-Phenyl - ethyl alcohol CH CH,Ph.CH,OH. Benzyl-carbinol. (212°). S.G. 21 1.034. Formed amalgam (Radziszewski, B. 9, 373). Oil. Ac₂O at 150° converts it into an acetyl derivative C₈H₉OAc, (224°), S.G. 1.029.

PHENYL-ETHYL-ALLOPHANIC ETHER C₁₂H₁₈N₂O₃ i.e. C₈H₉.NH.CO.NH.CO₂Et. [106°]. Formed from phenyl-ethyl-urea and ClCO₂Et (Neubert, B. 19, 1825). Needles (from water). PHENYL-ETHYL-ALLYL-GUANIDINE

C₁₂H₁₇N₃ i.e. CN₃H₂PhEt(C₃H₅). Formed by boiling ethyl-allyl-thio-urea with NHPh.HgCl and alcohol (Forster, A. 175, 41).—B'HgCl₂ aq.

PHENYL-ETHYL-ALLYL-THIO-UREA NH(C₂H₃).CS.NPhEt. [c. 26°]. Formed from allyl-thiocarbimide and ethyl-aniline (Gebbardt, B. 17, 3037). Colourless very soluble crystals.

PHENYL-ETHYL-w-AMIDO-ACETOPHEN-**ONE** C₁₆H₁₇NO *i.e.* C₆H₅.CO.CH₂.NPhEt. [95°]. Formed from ω -bromo-acetophenone and diethylaniline (Weller, B. 16, 26). Needles.

PHENYL ETHYLAMIDO - ETHYL PHONE C₆H₅.SO₂.C₂H₄.NHEt. Formed from C₂H₄(SO₂C₆H₅), and ethylamine at 85° (Otto,

J. pr. [2] 30, 337).—B'HCl. [130°]. Needles. PHENYL-ETHYL-AMIDO-(a)-NAPHTHO-QUINONE $C_{10}H_5(NPhEt)O_2$. [155°]. Formed by heating (a)-naphthoquinone (2 pts.) with ethyl-aniline (3 pts.) and HOAc (5 pts.) (Elsbach, B. 15, 1810). Violet needles (from alcohol).—B'HCl. [c. 230°]. Yellow needles, decomposed by water.

Phenyl - ethyl - amido - (β) - naphthoquinone. [165°]. Formed by heating (\$\beta\$)-naphthoquinone with ethyl-aniline and alcohol (Elsbach, B. 15. 691). Dark-red needles (from ether). Decomposed by boiling HClAq into ethyl-ariline and oxy-naphthoquinone.

PHENYL-ETHYL-AMIDO-PHENOL. Ethyl ether C, H, NO i.e. NPhEt. C, H, OEt. (319). Formed by heating phenyl-p-amido-phenol with alcoholic potash and EtI (Philip a. Calm, B. 17, 2434). Oil.

a-PHENYL-ETHYL-AMINE $C_sH_{11}N$ CH₃.CHPh.NH₂. (187.5° i.V.). S. 4.166 at 20°. Formed by reducing the phenyl-hydrazide or the oxim of acetophenone in alcoholic solution with HOAc and sodium-amalgam (Tafel, B. 19, 1929; 22, 1856; Kraft, B. 23, 2783). Formed also from acetophenone and sodium formate (Leuckart a. Janssen, B. 22, 1413). Liquid, miscible with alcohol and ether.—B'HCl. [158°]. $-B'_2H_2PtCl_a$. $-B'_2H_2SO_4$. [238°]. Prisms, almost [170°].—B'2H2C2O4. insol. alcohol.-B'H2CO4. Plates, m. sol. boiling alcohol.

Isomerides: - AMIDO-PHENYL-ETHANE ETHYL-ANILINE.

Di-phenyl-ethyl-amine C14H15N i.e. NPh.Et. Ethyl diphenylamine. (296°) (Girard, Bl. [2] 23, 3); (286°) (Lippmann a. Fleissner, M. 4, Got by heating diphenylamine with 797). alcohol and HClAq. Oil.

Di-phenyl-ethyl-amine CHPh,.CH,NH2. Formed by reducing di-phenyl-acetonitrile in alcohol with Na (Freund a. Immerwahr, B. 23,

2845). Oil.—B'HCl. [255°]. V. sol. water. Di-phenyl-ethyl-amine CH.Ph.CHPh.NH. (310°) at 737 mm. Formed by heating deoxybenzoïn (1 pt.) with ammonium formate (2 pts.) at 225° (Leuckhart a. Janssen, B. 22, 1409). Liquid, sl. sol. water. Potassium cyanate forms C₂H₃Ph₂.NH.CO.NH₂[99°]. Phenylthiocarbimide by reducing phenyl-acetic aldehyde with sodium- forms the corresponding C.H.Ph.NH.CS.NHPh

CPh, CH2.NH2.

[170°]; while phenyl cyanate gives the compound C2H2Ph2.NH.CO.NHPh [129].-B'HCl.-B'2H2PtCla.—B'HNO3: needles.—B'H2SO4.

Acetyl derivative. [148°]. Needles. Benzoyl derivative. [178°]. Needles. Di-phenyl-di-ethyl-amine NH(CH2CH2Ph) (336° at 603 mm.). Formed, together with NH2.CH2.CH2Ph and N(CH2.CH2Ph), by the action of zinc and HClAq on phenyl-acetonitrile (Spica, G. 9, 567). Formed also by distilling

CH₂Ph.CH₂·NH₃Cl (Fileti a. Piccini, G. 9, 294). Liquid, sl. sol. water.—B'HCl. [270°]. Pearly scales. -B'2H2PtCl, m. sol. water. Tri-phenyl-ethyl-amine

[116°]. Formed by reducing CPh. CN with zinc

and HClAq (Elbs, B. 17, 700).—B'HCl. [247°]. Tri-phenyl-tri-ethyl-amine N(CH2.CH2Ph)s. A product of the action of zinc and HCl on phenyl-acetonitrile (Spica, G. 9, 567). Oil.-B'HCl. [138°]. Needles, sl. sol. water.

PHENYL-ETHYL-ARSINE v. vol. i. p. 321. PHENYL-ETHYL-TRIAZOLE CARBOXYLIC

 $\begin{array}{ccc} \textbf{ACID} & \stackrel{N.NPh}{\sim} \textbf{C.CO}_2\textbf{H.} \end{array}$ [145°]. Got saponifying its hitrile (Bladin, B. 18, 1548; 25, 175). Melts at 123° when crystallised from water or alcohol, but at 145° when crystallised from benzene. - B'HCl: plates. - CuA'2 3 aq. -AgA': crystalline pp.

Methyl ether MeA'. [41°]. Ethyl ether EtA'. Oil.

Nitrile C₁₁H₁₀N₄. [38°]. Formed by the action of propionic anhydride on phenyl-hydrazine dicyanide. Converted by alcoholic NH, and H₂S into C₂N₃PhEt.CS.NH₂ [150°] which crystallises in yellow prisms

AmideC2N, PhEt.CO.NH2. [152·5°]. Formed from the nitrile, alcoholic potash, and H₂O₂. Small prisms, sl. sol. water.

DI-PHENYL-DI-ETHYL-DITRIAZYL

C₂₀H₂₀N_e i.e. CEt.N N.NPh C.C N-CEt [187°]. Formed by boiling phenyl-hydrazine dicyanide with propionic anhydride (Bladin, B. 22, 3115). Groups of prisms.—B"2HCl: minute prisms, decomposed by water.

PHENYL-ETHYL-CARBAMIC CHLORIDE NPhEt.COCl. [c. 52°]. Formed from ethylaniline and CO_2 (Michler, B. 9, 396). Needles.

PHENYL-ETHYL-SEMICARBAZIDE C9H13N3O i.e. NHPh.CO.NH.NHEt. [112°]. Formed from ethyl-hydrazine and phenyl cyanate (E. Fischer, A. 199, 295). Thin plates, m. sol. hot water. Yields a crystalline nitrosamine.

Phenyl-ethyl-semicarbazide NHEt.CO.NH.NHPh. [151°]. Formed from phenyl-hydrazine and ethyl cyanate (Fischer, A. 190, 109). Monoclinic tables (from dilute alcohol) a:b:c = .827:1:1.146; $\beta = 61^{\circ}$. a bluish-black pp. with cold Fehling's solution, and Cu,O on warming. Its nitrosamine crystal-

lises from acetone in yellow needles [86.5°].

PHENYL ETHYL CARBONATE C. H. O. 2 i.e.

C. H. O.CO.OEt. (c. 234°) (P.). S.G. 2 1.1134 (P.). Formed from KOPh and ClCO, Et (Fatianoff, 1864, 77). Formed also by the action of AlCl, on a mixture of phenol and ClCO, Et (Pawlewski, B. 17, 1205). Liquid. By long heating at 300° it is split up into Ph.CO, and Et.CO. (Bonder, B. 19, 2268).

Reference .- NITRO-PHENYL ETHYL CARBONATE. PHENYL-ETHYL-CYANAMIDE. by boiling phenyl-ethyl-thio-urea in benzene with PbO (Weith, B. 8, 1530). Vitreous mass.

DI-PHENYL-ETHYL-TRICYANIDE C₃N₃Ph₂Et. V.D. 129. [67°]. (234° at 15 mm.). Formed by the action of AlCl, on a mixture of benzonitrile and propionyl chloride at 70° (Krafft a. von Hausen, B. 22, 806). - B"H2PtCla.

PHENYL-ETHYLENE v. STYRENE.

s-Di-phenyl-ethylene C11H12, i.e. CHPh: CHPh. Stilbene. Mol. w. 180. [124°]. (307° i.V.). H.C.v. 1,775,600. H.C.p. 1,777,300 (from diamond) (Berthelot a. Vieille, A. Ch. [10] 4, 50);

1,773,331 (Ossipoff, Z. P. C. 2, 646).

Formation.—1. By the dry distillation of benzyl, or benzylidene, sulphide (Laurent, B. J. 25, 616; Maercker, A. 136, 91; Anschütz, A., 235, 206).—2. By distilling benzoic aldehyde with Na (Williams, Z. 1867, 432).-3. By heating benzoic aldehyde with phenyl-acetic acid and NaOAc at 250°; the yield being 55 p.c. (Michael, Am. 1, 313).—4. From s-di-phenylethane and Cl (Kade, J. pr. [2] 19, 465).—5. By heating benzylidene chloride with Na or with alcohol and zinc-dust (Limpricht, A. 139, 318; Lippmann, J. 1877, 405).—6. By passing s-diphenyl-ethane through a red-hot tube (Otto a. Dreher, A. 154, 177).—7. By heating benzoin with zinc-dust (Limpricht, A. 155, 80).—8. By passing toluene over heated PbO (Behr a. Dorp, B. 6, 754).—9. By heating di-phenyl-acetylene with HI and P at 175° (Barbier, J. 1874, 421).-10. By distilling lead phenyl-acetate with sulphur (Radziszewski, B. 6, 390).—11. Together with benzonitrile by the action of zinc-dust and HCl on C₆H₅.CSNH₂ (Bamberger, B. 21, 55).— 12. By heating di-phenyl-fumarate or di-phenylcinnamate (Anschütz, B. 18, 1945).—13. By heating CHPhBr.CHPhBr with alcoholic KSH in a sealed tube at 100° (Auwers, B. 24, 1779). 14. By heating thio-benzoic aldehyde at 190° (Baumann a. Klett, B. 24, 3308).

Properties .- Monoclinic plates, v. sol. ether, sl. sol. cold alcohol. Combines with N2O4 forming C₂H₂Ph₂N₂O₄ [c. 300°] crystallising in needles, sl. sol. hot alcohol (Gabriel, B. 18, 2438). Picryl chloride forms a combination C₁₄H'₁₂C₆H₂(NO₂)₂Cl [71°] (Liebermann, B. 8,

Reactions.-1. Yields phenanthrene and toluene when passed through a red-hot tube (Graebe, B. 6, 126) .- 2. Reduced by HIAq at 150° to s-di-phenyl-ethane (Limpricht Schwanert, A. 145, 333) .- 3. Bromine added to an ethereal solution forms a product containing $C_{14}H_8Br_2O_2$ [121°] which gives rise to $C_{14}H_8Br_4O_2$ [150°] and $C_{14}H_8Br_4O_2$ [206°] and when dissolved in alcohol and reduced by sodium-amalgam yields C14H10O2, crystallising from alcohol in flat plates [172°], whence PCl, forms C_{1,}H₂ClO₂ [58°], C_{1,}H₂ClO₂ [87°], and C_{1,}H₂ClO₂ [190°] (Limpricht a. Schwanert, A. 153, 121).—4. Fuming HNO, added to an ethereal colution forms C. H. NO, 5000°3. solution forms $C_{14}H_{11}N_2O_2$ [220°], which is converted by boiling alcohol into $C_{29}H_{22}N_2O_4$ [57'-78°] (Lorenz, B. 7, 1097; 8, 1050)

u-Di-phenyl-ethylene CH2:CPh2 (162° at 15 mm.). Formed by boiling CHPh, CH, Cl with alcoholic potash (Hepp, B.7, 1409). Formed also by the action of benzene and AlCl. on CH₂:CBr₂ and on CHBr:CBr₂ (Demole, B. 12, 2245; Anschütz, A. 235, 154. Liquid. Oxidised by CrO₃ to benzophenone. Combines with Br, forming CPh, Br. CH, Br. which readily gives off HBr, and forms bromo-di-phenyl-ethylene [40°] (c. 170° at 11 mm.).

Isomeride. [190°]. A product of the action of alcoholic potash on exo-chloro-di-phenylethane (Hepp, B. 7, 1412). Small plates (from ether), v. sl. sol. alcohol.

Tetra-phenyl-ethylene $\mathbf{C_{26}H_{20}}$ i.e. $\mathbf{C_{2}Ph_{4}}$.

Mol. w. 382. [221°]. (193° at 30 mm.).

Formation.—1. By heating CPh.Cl. with finely-divided silver (Behr, B. 3, 751; 5, 277).— 2. By heating benzophenone with zinc-dust (Staedel, B. 6, 178; A. 194, 307). — 3. By strongly heating chloro-di-phenyl-ethane (Engler a. Bethge, A. 174, 194).—4. From CPh₂Br₂ by repeated distillation (Friedel a. Balsohn, Bl. [2] 33, 337).—5. A by-product in the preparation of tri-phenyl-methane from benzene, chloroform, and AlCl₃ (Schwarz, B. 14, 1526).

Preparation.—1. By adding Br to di-phenylmethane and warming the resulting CHPh₂Br; the yield is 80 p.c. (Boissieu, Bl. [2] 49, 681). 2. By heating di-phenyl-methane (20 g.) with sulphur (8 g.) for 9 hours to 250°, and finally for one hour to 290°, exhausting with ether, and recrystallising the residue from benzene; the yield is 75 p.c. (Ziegler, B. 21, 780).

Properties.—White needles, v. sol. hot benz-

ene and CS2, v. sl. sol. ether. Yields benzophenone (2 mols.) on oxidation (Anschütz, A. 235, 221). Yields a tetrasulphonic acid.

Isomeride C₁₃H₁₀? [244°]. Formed by heating (α)-benzpinacolin with soda-lime at 370° (Zincke a. Thörner, B. 11, 1397). Needles (from alcohol). Perhaps identical with the preceding.

References .- DI-AMIDO-, BROMO-, DI-CHLORO-, DI-10DO-, NITRO-, NITRO-AMIDO-, and OXY- PHENYL-ETHYLENE.

PHENYL-ETHYLENE-DIAMINE $C_8H_{12}N_2$ $(262^{\circ}$ C₆H₅NH.CH₂.CH₂.NH₂. uncor.). Formed by boiling phenyl-amido-ethyl-phthalimide (got from bromo-ethyl-phthalimide and aniline) with conc. HClAq (Gabriel, B. 22, 2224). Liquid, miscible with water, forming an Absorbs CO2 from the air, alkaline solution. forming a crystalline carbonate.—B"2HCl: small greenish needles, acid in reaction .-B"HCl. Neutral to methyl-orange. - B"H, Br2. -Picrate [143°]. Flat yellow tables.

Di-acetyl derivative C₈H₁₀Ac₂N₂. [116°]. Crystals, v. e. sol. Aq (Newman, B. 24, 2193). Di-benzoylderivative. [143.5°]. Prisms.

Phenyl-di-ethylene-triamine $C_0H_4N(C_2H_4,NH_2)_2$. (above 300°). Formed by the action of boiling cone. HBr upon its diphthalyl derivative $NPh(C_2H_4,N:C_6H_4O_2)_2$ [211°], which is a product of the action of aniline on bromo-ethyl-phthalimide at 100°-180° (G.). Thick ammoniacal liquid, miscible with water. Absorbs CO₂ from the air.—B"H₂Br₂.—Picrate. [202°]. Needles (from alcohol).

Di-phenyl-ethylene-diamine C14H16N2 i.e. C.H.(NHPh). [63°]. Prepared by heating ethylene bromide (1 mol.) with aniline (4 mols.); the yield being 80 p.c. of the theoretical (Morley, B. 12, 1794; cf. Hofmann, Pr. 10, 104; Gretillat, M. S. [3] 3, 383). Plates, v. sol. alcohol. Yields a di-nitrosamine C₂H₄(NPh.NO)₂ [157°].

Reacts with benzoic aldehyde forming the compound C2H4(NPh)2:CHPh [187°], while cuminic, salicylic, anisic, isobutyric, and heptoic aldebydes yield corresponding compounds [125°], [116°], [164°], and [95°] respectively (Moos, B. 20, 732).—B"2HCl.—B"H₂PtCl_e.

Mono-acetyl derivative. [128°]. Got by heating the base with chloro-acetic acid and NaOAc at 170° (Bischoff a. Nastvogel, B. 22,

1783).

Di-acetyl derivative. [158°]. Crystals.

Di-phenyl-ethylene-diamine $C_{14}H_{16}N_2$ i.e. CHPh(NH₂).CHPh(NH₂). [121°]. Formed, together with benzoic aldehyde, by the action of boiling HClAq on C₂₈H₂₄N₂, which is a product of the action of Na on amarine (Grossmann, B. 22, 2299). Formed also by the action of ammonia on the hydrocyanide of benzoic aldehyde (Limpricht a. Müller, A. 111, 142). Is perhaps identical with lophine. Plates (from hot water). Reacts with benzoic aldehyde forming CHPh:N.CHPh.CHPh.N:CHPh [164°], m-nitrobenzoic aldehyde forming C₂₈H₂₂N₄O₂ [161°], with salicylic aldehyde forming C2, H24N2O2 [205°], and with cuminol forming a compound [168°].—B"2HCl. White needles (from hot water) .- B"H2PtCl : dark-yellow crystals.

Di-acetyl derivative. [above 350°]. Phthalyl derivative $C_{8}H_{4}:C_{2}O_{2}:N_{2}H_{2}:C_{2}H_{2}Ph_{2}$. [213°]. Minute crys-

Di-phenyl-di-ethylene-diamine C16H18N2 i.e. NPh C.H. NPh. Diphenylpiperazine. phenylpyrazine hexahydride. [163°]. (c. 300°). Prepared by heating ethylene bromide (1 pt.) with aniline (1 pt.) and NaOAc, and by the action of ethylene bromide on di-phenyl-ethylene-diamine at 120° (Morley, B. 12, 1795; Bischoff, B. 22, 1777; cf. Hofmann, Pr. 9, 277; 10, 104; Lellmann a. Schleich, B. 22, 1387; Bischoff, B. 22, 1778). Formed also by heating pyrazine hexahydride (1 pt.) with bromo-benzene (11 pts.) at 270° (Schmidt a. Wichmann, B. 24, 3239). Needles, sol. alcohol and ether. Its solutions are neutral to litmus. Yields a crystalline di-nitrosoderivative which may be reduced by tin and HCl

to $C_6H_s(NH_2)N < CH_2, CH_2 > NC_6H_3NH_2$, which, when diazotised and combined with naphthylamine sulphonic acid, yields a colouring matter which dyes cotton.—B"2HCl.—B"H,PtCl.

Methyl-iodide B'MeI. Crystalline. Yields B'2Me2PtCl6.

Ethylo-iodide B'Etl. [100°]. Yields B'2Et2PtCla.

References .- NITRO- and OXY- DI-PHENYL-ETHYLENE-DIAMINE.

DI-PHENYL - ETHYLENE - DI-BENZYL-DI. CHPh(NHCH,Ph).CHPh(NHCH,Ph). [153°]. Formed by reducing the compound CHPh(N:CHPh).CHPh(N:CHPh) [163°], which is itself got by reducing amarine with Na (Grossmann, B. 22, 2301). White needles.

DI - PHENYL - ETHYLENE - DICARBAMIC

ACID C₂H₁(NPh.CO₂H)₂. Ethylether Et₂A" [88°]. Needles. C2H, (NPh.COCI). [183°]. Chloride Formed from di-phenyl-ethylene-diamine and COCl₂ (Hanssen, B. 20, 781). Prisms.

DI-PHENYL-ETHVLENE DI-o-CARBOXY-LIC $C_gH_4(CO_gH).CH:CH.C_gH_4.CO_gH.$ ACID [264°]. Formed by heating for four hours at 215° CO COH CH.CH, C, H4.CO, H (2 pts.) with KCy (5 pts.) (Hasselbach, A. 243, 258). Small needles (from dilute HOAc). Changes on heating into the parent acid [196°]. Reduced by conc. HIAq to C₂H₄(C₆H₄.CO₂H)₂ [185°].—Ag₂A". Flocculent pp.

Ethylether Et,A". [80°]. Needles. Isomeride v. DI-PHENYL-MALEIC ACID.

References .- NITRO- and OXY- DI-PHENYL-

ETHYLENE CARBOXYLIC ACIDS.

DI - PHENYL - ETHYLENE - DI - ETHYL -**DIAMINE** $C_{18}H_{24}N_2$ *i.e.* $C_2H_4(NPhEt)_2$. [70°]. Formed from di-phenyl-ethylene-diamine and EtI (Hofmann, Pr. 10, 104).—B"2HI.— B'H,PtCla: needles.

PHENYL-ETHYLENE-GLYCOL v. DI-OXY-

ETHYL-BENZENE

DI-PHENYL-ETHYLENE-DI-HYDRAZINE $C_{14}H_{18}N_4$ i.t. $C_2H_4(NPh.NH_2)_2$. [90°]. Formed from sodium phenyl-hydrazine and ethylene bromide in bertzene (Burchard a. Michaelis, B. 21, 3202; A. 254, 116). Prisms or plates. Aldehyde forms C₂H₄(NPh.N:CHMe), [82°]. Benzoic aldehyde forms an analogous body [193°]. Acetone and acetophenone form analogous compounds [72°] and [118°]. Phenyl-thiocarbimide forms NH₂,NPh.C₂H₄,NPh.NH.CS.NHPh [164°] and C₂H₄(NPh.NH.CS.NHPh)₂ [194°]. Yields a crystalline nitroso- derivative [160°]. SOCl₂ forms C₂H₄(NPh.N:SO)₂ [123°] (Michaelis a. Ruhl, A. 170, 122).—B"H₂Ol₂. [212°]. Needles, sl. sol. HClAq.—B"H₂SO₄.—B"2HNO₅ [173°].— B"H2C2O4 [183°].

Acetyl derivative C14H16Ac2N2. [222°] Succinyl derivative C14H16N2:C4H4O2.

[c. 126°].

Succinoxyl derivative C₂H₄(NPh.NH.CO.C₂H₄.CO₂H)₂. [203°]. Oxalyl derivative C14H16N2:C2O2.

Di-phenyl-di-ethylene-dihydrazine

 $(C_2H_4)_2(N_1HPh)_2$. [178°]. Got by heating phenyl-hydrazine with $C_2H_4Br_2$ and alcohol (Marckwald, C. C. 1888, 1410).

DI-PHENYL ETHYLENE DIKETONE

C.H., CO.CH, CH, CO.C. H, Diphenacyl. Succinophenone. [134°] (A.); [140°] (C.); [142°-145°] (P.). Formation. -1. Together with the isomeric CH_xCO CH_xCPh_x>0, by the action of succinyl chloride and AlCl₃ on benzene (Auger, A. Ch. [6] 22, 312; Claus, B. 20, 1374).—2. From acetophenone by treatment with fuming HNO, and reduction of the resulting $C_{16}H_{10}N_{.}O_{4}$ by zinc-dust and HOAc (Hollemann, B. 20, 3361).—3. By the action of KOHAq on di-benzoyl-propionic acid suspended in alcohol (Paal, B. 21, 3056).

Properties. Needles, v. sol. ether. Oxim C2H4(CPh:NOH)2. $[204^{\circ}]$

Phenyl-hydrazide C₂H₄(CPh:N₂HPh)₂.

[1809]. Needles, v. sol. ether.

DI-PHENYL ETHYLENE DIKETONE CARBOXYLIC ACID v. PHENACYL-BENZOYL-ACRTIC ETHER.

Di-phenyl-ethylene diketone di-o-carboxylic acid C₁₈H₁₄O₆ i.e. C₂H₄(CO.C₆H₄.CO₂H)₂ [172°]. Obtained by boiling di-phthalyl-ethane with

alkalis (Gabriel a. Michael, B. 10, 1561, 2199; Roser, B. 17, 2622; 18, 803, 3115; Baumann, B. 20, 1486). Prisms (from water), v. sol. alcohol. Reconverted by conc. H₂SO, into di-phthalyl ethane C₂H₂(C₂O₂C₆H₄)₂. A boiling alcoholic solution of phenyl-hydrazine forms C,0H,22N,O, [237]. Hydroxylamine at 100° gives rise to C_2H_4 ($C \leqslant_{N.O}^{C_5H_4} > CO$)₂ [270°] (Baumann, B. 20, 1492).—Ag₂A": small plates, sl. sol. hot water.

(a)-Anhydride C₁₄H₁₂O₅. [230°]. Got by heating the acid alone, or together with diphthalyl-ethane, by heating it for a short time with HCl. Needles (from alcohol).

(β)- $Anhydride C_{18}H_{12}O_{5}$. [202°]. Formed by more prolonged heating of the acid with HCl, and also, together with the (a)-isomeride, by heating the acid by itself. Prisms (from Prisms (from alcohol). Both anhydrides are reconverted into the acid by boiling alkalis, and into di-phthalylethane by elimination of H₂O.

Isomeride v. DI-BENZOYL-SUCCINIC ACID.

PHENYL-ETHYLENE OXIDE $_{\text{CHPh}}^{\text{CH}_2}$ >0.

(260° at 50 mm.). Formed by heating di-oxyethyl-benzene with dilute H2SO4 (Breuer a. Zincke, B. 11, 1402). Oil. Converted by PBr, into CHPhBr.CH2Br.

Tetra-phenyl-ethylene oxide v. Benzpinaco-

PHENYL-ETHYLENE SULPHIDE

CH. cHPh>s. 1·099. S.G. Formed from PhCHBr.CH2Br by successive treatment with alcoholic KSHAq (Spring a. Marsenille, Bl. [3] 7, 13). Oil with strong smell, sol. alcohol-ether. Oxidised by CrO, to benzoic acid.

CHPh S. Di-phenyl-ethylene sulphide [169°]. A product of the distillation of benzyl

sulphide (Barbier, J. 1876, 421). Needles. Di-phenyl-ethylene disulphide C2H4(SPh)2. [65°]. Formed from NaSPh and C,H,Br, (Ewerlöf, B. 4, 716). Needles, insol. water.

DI-PHENYL-ETHYLENE DISULPHONE

 $C_2H_4(SO_2C_4H_5)_2$. [180°]. Formation.—1. By oxidation of $C_2H_4(SPh)_2$ (Ewerlöf, B. 4, 717).—2. By boiling sodium benzène sulphinate (100 pts.) with C₂H₄Br₂ (58 pts.) in alcohol (Otto, B. 13, 1279; J. pr. [2] 30, 174). 3. By adding CH₂.CCl₂.CO₂Na (1 mol.) to C₆H₅.SO₂Na (2 mols.) in weak alcoholic solution, kept neutral by Na,CO₃ (Otto, J. pr. [2] 40,531). 4. By heating CH₃.CCl₃ with C₈H₅.SO₂Na at 160° (Otto, B. 21, 1691).

Properties.—Triclinic needles or plates, sl. sol. water, m. sol. alcohol, v. sol. HOAc.

Reactions.—1. Sodium-amalgam reduces it to alcohol and C.H.SO.Na, which is finally reduced to C.H.SNa.—2. Chlorine in diffused daylight forms C2H4Cl2 and benzene sulphonic chloride. In sunlight the products are C2H1Cl2, SO₂Cl₂, and chlorinated benzenes.—3. Boiling aqueous KOH splits it up into C,H,SO,K and C.H.SO.C.H.OH. Conc. KOHAq forms a compound [88°] crystallising from alcohol.—4. Aqueous NH, forms C.H. SO.ONH, and C.H.SO.ONH (C.H., SO, CH. CH.), NH [78°] which yields B'HCl [193°], B',H,PtCl, a nitrate [190°], and the derivatives (C.H.,SO, CH.,CH.), NMe and (C.H.SO,CH,CH,2),NMeHCl[221°].-5.Aqueous

ethylamine C,H,SO,NH,Et forms C.H. SO. C.H. NHEt, an oil which yields B'HCl [130°].—6. Alcoholic KCy forms C₆H₄SO₂K and C2H4(CN)2.

PHENYL-ETHYLENE-THIO-UREA

CS NPh C₂H₁. [155°]. Formed from phenylethylene-diamine and CS₂ (Newman, B. 24, 2191). White plates, v. sol. alcohol.

Di-phenyl-ethylene-ψ-thio-urea

 $\stackrel{\text{NPh}}{\sim}$ C:NPh. [136°]. (above 300°). Formed by heating di-phenyl-thio-urea with ethylene bromide (Will, B. 14, 1490; 15, 343). Plates (from alcohol). KClO, and HClAq form $C_{15}H_{14}N_2SO_3$ [187°] (Andreasch, M. 4, 134).— B'H2SO4: thick prisms, v. sol. water.

Di-phenyl-ethylene-di-thio-di-urea

 $C_{16}H_{16}N_4S_2$ i.e. $C_2H_4(NH.CS.NHPh)_2$. [193°]. Formed from $C_2H_4(NH_2)_2$ and phenyl-thio-carbimide in alcohol (Lellmann a. Wurthner, A. 228, 234). White scales, insol. alcohol, sl. sol. HOAc. Decomposed by heat, giving di-phenyl-thio-urea and a crystalline body [164°].

PHENYL-ETHYLENE-UREA

 $C_2H_4 < NH > CO$. [161°]. Formed from phenylethylene-diamine hydrochloride and potassium cyanate, NH3 being given off (Newman, B. 24, 2192). Plates, v. sol. alcohol, insol. cold water.

Di - phenyl - ethylene - urea CH. NPh CO.

[209°]. Formed by the action of COCl₂ in benzene on $C_2H_1(NHPh)_2$; an intermediate body being C₃₁H₂₈N₄Cl₂O₃ (Michler a. Keller, B. 14, 2183; Hanssen, B. 20, 784). Plates.

PHENYLETHYL-ETHYL-PYRIDINE

 $C_{13}H_{17}N$ i.e. $N \leqslant \stackrel{C(CH_2\cdot CH_2Ph).}{CH:CEt} \stackrel{CH}{\longrightarrow} CH$ (316° cor.). -CH. S.G. 9 1.016. Formed by reducing styryl-ethylpyridine with HIAq at 165° (Plath, B. 21, 3093; 22, 1057). Oil, v. sl. sol. water, v. sol. alcohol and ether, volatile with steam. Yields C15H1,Br2N [128°] whence AgOAc forms C₁₅H₁₅(OAc)₂N (315°-320°). — B'₂H₂PtCl₆. [168°]. Yellow - B'HHgCl. [136°]. Ñeedles. needles. -B'HAuCl, aq: crystalline mass.

(314°). S.G. 2 Hexahydride C, H23N. 9663. Got by reducing the preceding body in alcohol with Na. Oil, sl. sol. water, miscible

with alcohol and ether.

PHENYL-ETHYL-FURFURANE

C,H,CH,CH,CH,C,H,O. A product of (241°). the action of Na on an alcoholic solution of C,H,O.CH:CPh.CN, which is formed by condensation of furfuraldehyde with phenyl-acetonitrile (Freund a. Immerwahr, B. 23, 2848). smelling like CH₂Ph₂.

DI-PHENYL-ETHYL-GUANIDINE

C15H17N3 i.e. NHEt.C(NPh).NHPh. Got from phenyl-ethyl-cyanamide and aniline at 100° (Weith, B. 8, 1531). Crystalline.—B',H,PtCl,

PHENYL-ETHYL-HYDANTOIN C11H12N2O2 i.e. CHPh CO.NEt [94°]. Formed from phenyl-hydantoin, alcoholic KOH, and EtI (Pinner, B. 21, 2325). Prisms, v. sol. alcohol, sl. sol. cold water. Decomposed by baryta into ethylamine and phenyl-amido-acetic acid.

Phenyl-ethyl-\psi-hydantoin

CHPh CO.NEt Separates from a dilute alkaline solution of the preceding isomeride on standing. Slender needles, insol. water, nearly insol. alcohol. Decomposed by heating with baryta-water into NH, NH,Et, and a-oxyphenyl-acetic acid.

u-PHENYL-ETHYL-HYDRAZINE C.H., N. i.e. NPhEt.NH,. (230°). Formed by reducing the nitrosamine of ethyl-aniline with zinc-dust, HOAc, and alcohol (Fischer, B. 8, 1642; A. 199, 325; Philips, B. 20, 2485). Formed also from EtBr and NPhNa.NH2 in benzene (Michaelis a. Philips, A. 252, 270). Oil. Reduces Fehling's solution on warming. Oxidised by HgO to diphenyl-di-ethyl-tetrazone NPhEt.N:N.NPhEt [108°]. SOCl₂ forms oily NPhEt.N:SO (Michaelis, B. 22, 2231).—B'HCl. Plates.

Acetyl derivative NEtPh.NHAc. Ethylo-bromide NH2.NPhEt2Br. metric prisms (from alcohol); a:b:c = .822:1:.827. V. e. sol. water, insol. ether. Decomposes at 193°. Insol. KOHAq. Converted by moist Ag₂O into a caustic hydroxide. Yields also (NH₂.NPhEt₂)₂H₅Fe₂Cy₁₂ 2aq (Fischer, A. 190,

Ethylo-chloride NH, NPhEt,Cl. [198°]. Needles, v. e. sol. water. - B',Et,PtCl, Ethylo-iodide NH, NPhEt,I. [145°].

NHPh.NHEt. s-Phenyl-ethyl-hydrazine Formed, together with the preceding isomeride, by heating phenyl-hydrazine with EtBr. The crude product is dissolved in water mixed with NaOHAq, and the ppd. oil extracted with ether. The ethereal solution is freed from phenylhydrazine by HCl and the filtrate oxidised by HgO. On addition of HCl it deposits di-phenyldi-ethyl-tetrazone, and the mother-liquor yields by steam-distillation oily C.H.N:NEt (175°-185°). The NPh:NEt is then reduced by sodiumamalgam (Ehrhardt a. Fischer, B. 11, 613).

Properties.-Oil, sol. alcohol and ether. Readily reduces Fehling's solution and HgO. Zinc-dust and HOAc yield aniline and ethylamine.—B'H₂C₂O₄: needles, v. sol. hot water.
DI-PHENYL-ETHYLIDENE-DIAMINE

Ethylidene-di-aniline, vol. ii. p. 496.

DI - PHENYL - ETHYLIDENE - DI - ETHYL-**DIAMINE** C₁₈H₂₄N₂ i.e. CHMe(NPhEt)₂. Formed from ethyl-aniline and aldehyde (Schiff, A. 140, 95; cf. Schultz, B. 16, 2601). Thick liquid. B'2H2PtCl

PHENYL-ETHYLIDENE DI-ETHYL DI-SULPHONE CH₃.CPh(SO₂Et)₂. [101°]. Formed from benzylidene di-ethyl di-sulphone, MeI, and EtONa (Fromm, A. 253, 154). Needles.

PHENYL-ETHYLIDENE-HYDRAZINE

CH₃.CPh:N.NH₂. (255°). Formed from acetophenone and hydrazine hydrate (Curtius, J. pr. [2] 44,540). Liquid. Yields CH, CPh:N.N:CHPh [59°] and N₂(CPh.CH₃), [121°]. DI-PHENYL ETHYLIDENE DISULPHONE

CH₂.CH(SO₂Ph)₂. [102°]. Got by oxidising CH₃.CH(SPh)₂ or CH₃.C(SPh)₂.CO₂H with dilute (1 p.c.) KMnO, (Escales a. Baumann, B. 19, 2815). Needles or thin lamellæ, insol. water, acids, and alkalis, sl. sol. alcohol and ether.

Not attacked by alcoholic potash at 140°. PHENYL - ETHYLIDENE - DI - THIO - DI -GLYCOLLIC ACID CH,.CPh(S.CH,.CO,H),

[136°]. Got by the action of ZnCl₂ on a mixture of acetophenone and thioglycollic acid (Bongartz, B. 21, 483). Needles, sol. hot water, CHCl3, and HOAc.

PHENYL-ETHYL-IMESATIN v. Di-phenyl-

di-ethyl-diamide of Isatin.

PHENYL ETHYL KETONE C,H10O i.e. C₆H₅.CO.C₂H₅. Propiophenone. [21°]. cor.) (M. a. G.). V.D. 4.64 (obs.). S.G. 2 1.009

(W.).

Formation.--1. By distilling a mixture of calcium benzoate and propionate (Barry, B. 6, 1006). - 2. From BzCl and ZnEt₂ (Freund, A. 118, 20; Kalle, A. 119, 166). -3. By the action of Na on a mixture of BzCl and EtI (Bechi, B. 12, 463).—4. By the action of propionyl chloride on benzene in presence of AlCl, (Morley a. Green, B. 17, 3018; Pampel a. Schmidt, B. 19, 2896).—5. By the decomposition of propylene phenyl-ethyl-ketate by HI or H2SO4 (Morley a. Green).-6. By oxidising CHPhEt.OH (Wagner, J. R. 16, 325).-7. From benzoyl cyanide and ZnEt2, either directly or by oxidising the product (Frankland a. Louis, C. J. 37, 745).—8. From phenylallylene (Körner, B. 21, 277).

Properties.—Tables. Does not combine with $NaHSO_{g}$. Yields benzoic acid on oxidation. Reduced by sodium-amalgam to CHPhEt.OH

(211°).

Oxim CPhEt:NOH. Oil. Phenyl hydrazide. Oil.

Phenyl ethyl diketone C_eH_s.CO.CO.C_eH_s. Propionyl-benzoyl. (239°). Got by distilling its mono-oxim (obtained from ethyl benzoylacetic ether) with dilute H2SO4 (Muller a. Pechmann, B. 22, 2131). Pungent liquid, volatile with steam, m. sol. water.

Reference.—NITRO- and OXY- PHENYL ETHYL KETONE

PHENYL ETHYL KETONE DIBROMIDE v. DI-BROMO-CUMENE

PHENYL-ETHYL KETONE o-CARBOXYLIC ACID C2H5.CO.C6H4.CO2H. Propionyl-benzoic acid. [92°]. Formed by boiling phthalyl-propionic acid with KOHAq (Gabriel, B. 11, 1014; 19, 840). Needles (from dilute alcohol).—AgA'.

Anhydride $C_8H_4 < CO \longrightarrow 0$.

Formed by heating phthalic anhydride with succinic acid and NaOAc (G.). Formed also by distilling the anhydride of the dicarboxylic acid (Reser, B. 18, 3117). Plates (from water). Combines with N_2O_4 forming $C_{10}H_8N_2O_6$ [90°]. $A\ mide$ $C_6H_4(CO.NH_2).CO.C_2H_2$. [159°].

Formed from the anhydride and alcoholic NH.

Isomerides v. Benzoyl-Propionic acid, vol. i. p. 487.

Phenyl ethyl ketone dicarboxylic acid

C₆H₅.CO.CH₂.CH(CO₂H)₂. β-Benzoyl-isosuccinic acid. [180°]. Obtained by saponifying its ether which is formed from ω-bromo-acetophenone and sodium malonic ether (Bischoff, B. 16, 1044; 19, 95; Kues a. Paal, B. 18, 3324). Needles, v. sol. alcohol and ether. Yields \$\beta\$. benzoyl-propionic acid when heated.-Ag2A": needles.—Et,A": oil.

Phenyl hydrazide

C₆H₅.C(N₂HPh).CH₂.CH(CO₂H)₂. [120°]. Phenyl ethyl ketone dicarboxylic acid

C₆H₄(CO₂H).CO.CH₂.CH₂.CO₂H. [137°]. Formed by boiling its dilactone with water or aqueous Vor. ĬV.

alkalis. Small six-sided prisms, re-converted by heat into the dilactone.—CaA".—BaA".—Ag₂A".

Dilactone C.H. CO.O C CH2. CH2. [120°]. A product of the action of phthalic anhydride on succinic acid and NaOAc at 250° (Roser, B. 17, 2770; 18, 804, 3115). Needles, sl. sol. cold water. At 260° it gives off $\rm CO_2$ and changes to C.H. CO.O C:CHMe. HIAq and P at 190° reduce it to $C_6H_4(CO_2H).CH_2.CH_2.CO_2H.CO_2H.Cold NH_3Aq$ forms C_6H_4 C:CH.CH_2.CO_2H [225°]. Boiling alcoholic NH, forms C₁₁H₉NO₃ [c. 205°]. Sodiumamalgam reduces it to phthalyl-propionic acid $C_{11}H_{10}O_4$. Phenyl hydrazine forms $C_{17}H_{14}N_2O_3$ [210°] which gives CaA'_2 aq.

Isomeride v. BENZOYL-SUCCINIC ACID.

Phenyl ethyl ketone tetracarboxylic acid $C_6H_4(CO_2H).CO.C(CO_2H)_2.CH_2.CO_2H$. Formed by the action of chloro-acetic ether on C17H18NaO, which is got from phthalyl-malonic ether and NaOEt (Wislicenus, A. 242, 58). Oil.—Ag, Av.

PHENYLETHYL METHYL KETONE v.

BENZYL-ACETONE.

Diphenylethyl methyl ketone

CH₃ CPh₂.CO.CH₃. [41°]. (311° i.V.). A product of the action of zinc and HCl on an alcoholic solution of acetophenone (Zincke a. Thörner, B. 11, 1989). Prisms (from alcohol).

PHENYLETHYL-METHYL-PYRIDINE C₁₄H₁₅N i.e. C₂H₄Ph.C₅H₅MeN. (290°-295°). S.G. ⁹ 1.0283. Formed by reducing styrylmethyl-pyridine with HIAq at 160° (Bachér, B. 21, 3076). Oil, volatile with steam. - B'2H2PtCl3. [168°]. -B'HHgCl₃ aq. (95°]. Silky needles. -B'C₆H₃N₃O₇. [156°]. Yellow needles. Hexahydride C₁₄H₂₁N. (c. 288°). S.G. ²

9775. Got by reducing styryl-methyl-pyridine in alcohol with Na. Liquid, v. sl. sol. water.

PHENYL-ETHYL-(B)-NAPHTHOTRIAZINE DIHYDRIDE C₁₀H₁₇N₃ i.e. C₁₀H₆</br>
N:CHEt [219°]. Formed by adding propionic aldehyde to benzene-azo-(B)-naphthylamine in alcohol (Goldschmidt a. Poltzer, B. 24, 1006). White

needles, v. sol. alcohol.—B'HCl. [258°].—
B'₂H.PtCl₆. Small yellow crystals.
PHENYL-ETHYL-OXAMIDE C₁₀H₁₂N₂O₂ i.e.

NHPh.CO.CO.NHEt. [170°]. Formed by the action of ethylamine on phenyloxamic ether or of aniline on ethyloxamic ether (Wallach, A. 184, 66; 214, 259). Needles (from alcohol). Converted by PCl_s (2 mols.) into a base which forms the salt (C₁₀H₉N₂Cl)₂H₂PtCl₈.

Di-phenyl-di-ethyl-oxamide v. vol. iii. p. 654. PHENYL - ETHYL - PHENOL v. OXY - DI -

PHENYL-ETHANE.

PHENYL - p - ETHYL - PHENYL - ETHANE C₆H₃.CH₂.CH₂.C₆H₄.C₂H₄. (294°). Formed by reduction of C₆H₃.CH₂.CO.C₆H₄Et with HI and P (Söllscher, B. 15, 1681). Liquid.

Isomeride. Got by the action of zinc-dust on a mixture of ethyl-benzene and α-bromoethyl-benzene (Radziszewski, B. 6. 811; 140). Gives p-benzoyl-benzoic acid on oxidation.

PHENYL-ETHYLPHENYL-ETHYLENE

C.H.:CH:CH.C.H.Et. [90°]. Formed by boiling CH:Ph.OH(OH).C.H.Et[1:4] with dilute H₂SO₄ (Söllscher, B. 15, 1681). Plates.

PHENYL p-ETHYLPHENYL KETONE

CaHa.CO.CaH4Et. (331°) at 720 mm. Prepared by the action of AlCl, on a mixture of BzCl and ethylbenzene (Söllscher, B. 15, 1682; Smith, B. 24, 4029). Liquid, sol. alcohol and ether. Yields two oxims [108°] and [142°], both giving the same acetyl derivative [95°

PHENYL-ETHYL-PHENYL METHANE v.

BENZYL-ETHYL-BENZENE

PHENYL - ETHYLPHENYL - THIO - UREA NHPh.CS.NH.C₆H₄Et. [104°]. Formed from p-ethyl-phenyl-thiocarbimide and aniline in alcohol (Mainzer, B. 16, 2020). Plates.

PHENYL-DI-ETHYL-PHOSPHINE

PEt₂C₆H₅. (222° cor.). S.G. 13 .957. Formed from C₆H₅PCl₂ and ZnEt₂ (Michaelis, B. 8, 493; A. 181, 345). Oil. Absorbs oxygen from air, yielding OPEt₂C₆H₅ [56°]. Combines also with Cl and S forming crystalline C₁₀H₁₅PCl₂ and C₁₀H₁₅PS. — B'2HCl. — B'₂H₂PtCl₆ — B'H₁. — B'EtI. [115°].—B'2Et2PtCl6.—B'MeI. [95°].-B'2Me2PtCl8.

Di-phenyl-ethyl-phosphine PEtPh₂. (298°). Formed from Ph₂PCl and ZnEt₂ (Michaelis a. Link, A. 207, 214). Liquid, sol. alcohol and ether. Yields OPEtPh, on oxidation. EtI yields Ph₂PEt₂I [204°] which gives (Ph₂PEt₂Cl)₂PtCl₄ [218°]. MeI gives Ph₂PEtMeI [181°] S. 1.5 at 22°; 50 at 100°, which gives $(Ph_2PEtMeCl)_2PtCl_4$ [220°] and $Ph_2PEtMe.O.C_0H_2(NO_2)_3$ [86°].

Di-phenyl-ethyl-phosphine oxide PEtPh.O. [121°]. Formed as above, and also by the action of Ag₂O on Ph₃PEtI (Michaelis a. Soden,

A. 229, 317). Prisms, sol. ether and ligroin. PHENYL-DI-ETHYL PROPENYL TRISUL-PHONE PhSO₂CH₂, CMc(SO₂Et)₂. [128°]. Got from EtSH and PhS.CH₂, CO.CH₃, the product being oxidised (Autenrieth, B. 24, 169). Plates.

Di-phenyl ethyl propenyl trisulphone EtSO₂.CH₂.CMe(SO₂Ph)₂. [139°]. Made in like manner from PhSH and EtS.CH₂CO.CII₃. Needles (Autenricth, B. 24, 1513).

TRI-PHENYL-ETHYL-PROPYL-DI-THIO-BIURET C2S2N8Ph3EtPr. The (a)- compound [166°] is formed by the action of di-phenylpropyl-thio-urea on ClCS.NPhEt, while the (B)compound [165°] if got from di-phenyl-ethylthio-urea and CICS.NPhPr (Billeter a. Strohl, B. 21, 109). Both crystallise in needles, v. sol. hot alcohol.

DI - PHENYL - ETHYL - PROPYL - THIO-UREA NPhEt.CS.NPhPr. [66°]. Formed by the action of CSCl₂ on propyl-aniline followed by ethyl-aniline or vice versa (Billeter a. Strohl,

B. 21, 103).

PHENYL-ETHYL-PYRAZOLE C11H12N2 i.e. NPh $<_{N=-\dot{C}H}^{CEt;CH}$. (274°). S.G. 15 1.064. Formed from phenyl-hydrazine and propionyl-acetic aldehyde (Claisen a. Stylos, B. 21, 1148). Oil.

PHENYLETHYL-PYRIDINE CH CH:CH CH, CH, Ph. (289° $[-3^{\circ}].$ cor.). S.G. 2 1.0465. Formed by reducing styrylpridine with HI (Baurath, B. 21, 821). Liquid, sl. sol. water, volatile with steam. - B'2H2PtCl6.

[186°].—B'HAuCl, [150°]. Long yellow needles. —B'HHgCl_s. [149°]. —Hexahydride C₁₈H₁₈N. (288° cor.). S.G. 9874. Got by reducing the preceding base in alcohol with Na. Liquid, smelling like piper-

idine, sl. sol. water. Turns moist litmus blue. Has toxic properties similar to those of conline. Forms an oily nitrosamine. Yields pyridine carboxylic acid on oxidation.—B'HCl. [155°]. $-B'_{c}H_{2}PtCl_{s}$. [189°]. $-B'HAuCl_{4}$. [134°].

TETRA-PHENYL-ETHYL-PYRROLE

NEt<CPh:CPh [221°]. Formed by heating bidesyl with aqueous NEtH2 at 150° (Fehrlin, B. 22, 555). Needles (from chloroform) or plates (from HOAc).

DI-PHENYL-ETHYL-PYRROLE DI-O-CARB-

OXYLIC ACID C20H1, NO. i.e.

 $\text{NEt} < \begin{matrix} \text{C(C}_{0}\text{H}_{4}.\text{CO}_{2}\text{H):CH} \\ \text{C(C}_{0}\text{H}_{4}.\text{CO}_{2}\text{H):CH} \end{matrix}.$ [220°]. Formed by heating C₂H₄(CO.C₆H₄.CO₂H)₂ with ethylamine solution (Baumann, B. 20, 1488). Yellow plates (from alcohol), insol. water. - Ag.A".

PHENYLETHYL-QUINOLINE

C₈H₄ CH:CH N = C.CH₂·CH₂Ph. [c. 30°]. Formed by reducing styryl-quinoline with HI (Heymann a. Königs, B. 21, 1426). Crystals.—Picrate. [c. 130°]. Yellow prisms, sl. sol. alcohol.

PHENYL-ETHYL SULPHIDE Ph.S.Et. (204° i.V.) at 744 mm. S.G. 10 :1.032. Formed by heating PhSNa with EtI in sealed tubes at 120° (Beckmann, J. pr. [2] 17, 457). Obtained also by the action of H₂S at 50° on C_bH₃. SO.OEt (Otto a. Rössing, B. 20, 2275) and by the action of EtSH on C₆H₅.N₂Cl (Stadler, B. 17, 2078). Liquid, with nasty smell.

C.H.S.C.H.SO.H. Sulphonic acidFormed by boiling CaH, S.Na. CaH, SO, Na with

alcohol.—NaA': plates.

Phenyl ethyl disulphide PhS, Et. Formed together with Et,S, by heating C,H,SO,H with EtSH at 100° (Otto a. Rossing, B. 19, 3135; 20, 189). Heavy oil. Decomposed by alcoholic potash into EtSH, PhuS2, CaH, SO2H, and ethane sulphinic acid.

PHENYL ETHYL SULPHONE C,H1,0SO2 i.e. C₆H₅·SO₂·C₂H₅. [43°]. (above 300°). Formed from C₆H₅SO₂Na and EtBr or CH₃·CBr₂·CO₂Na (Otto, B. 13, 1274; J. pr. [2] 40, 535). Formed also from CH, CH(SO2Ph).CO2H by the action of alkalis and by oxidising PhSEt with dilute KMnO₄ (Beckmann, J. pr. [2] 17, 458). Monoclinic plates (Fock, B. 19, 1230), sol. hot water, v. sol. alcohol.

acid C, H, SO, CHMe.CC, H. a-Carboxylic Phenyl sulphono-propionic acid. [116°]. Formed by saponifying its ether, which is got from C, H, SO, Na and CH, CHBr. CO, Et (Otto, J. pr. [2] 40, 548). Groups of needles, v. sol. hot

water.—NaA'.—BaA'2 2aq: leafy aggregates.

Ethylether EtA'. [c. 17°]. Oil.

\$\beta\$-Carboxylic acid C_6H_5.SO_2.CH_2.CH_2.CO_2H. [124]. Formed from benzene sulphinic acid and \$\beta\$-iodo-propionic acid (Otto, \$\bar{B}\$. 21, 95). Plates, sl. sol. cold water, m. sol. ether.

PHENYL-ETHYL-THIAZOLE C, H, NS i.e. N CEt.S (296° cor.). Formed from thiopropionamide and bromo-acetophenone in alcohol (Hubacher, A. 259, 231). Oil.—B',H.PtCl,, [129°].—B'HBr. [70°]. White needles. TRI-PHENYL-DI-ETHYL-DI-THIO-BIURET

NPhEt.C(NPh).S.CS.NPhEt. [158°]. Formed from NPhEt.CS.Cl (2 mols.) and aniline (1 mol.) (Billeter a. Strohl, B. 21, 108). Yellow needles, aniline hydrochloride and potassium sulphosl. sol. alcohol.

PHENYL-ETHYL-THIOCARBAMIC ACID NPhEt.CS.OH.

Ethyl ether EtA. [18°]. (143° at 12 mm.). S.G. 15 1.066. Formed from NPhEt.CS.Cl and NaOEt (Billeter a. Strohl, B. 20, 1629; 21, 104). Crystals.

Phenyl-ether PhA'. [69°]. Formed from the chloride and phenol. Flat needles.

Chloride NPhEt.CS.Cl. [57°]. Formed from ethyl-aniline and CSCl. Prisms (from ligroin). Converted by alcoholinto (NPhEt.CS)20 [143°], S. (alcohol) 7 at 15°.

Phenylethyl-thiocarbamic acid. Phenyl-

ethylamine salt

C₂H₄Ph.NH.CS.SNH₃.C₂H₄Ph [130°]. Formed from phenylethylamine and CS2 (Neubert, B. 19, 1825). Sol. hot water and alcohol.

Phenyl-ethyl-di-thio-carbamic acid

NPhEt.CS.SH.

Ethyl ether EtA'. [67°]. $(305^{\circ}-315^{\circ}).$ Formed by heating NPh:C(NPhEt).SEt with CS, at 160° (Bernthsen a. Friese, B. 15, 568, Formed also from NPhEt.CSCl and 1533).NaSEt (B. a. S.). Prisms (from ether). Forms a crystalline compound with MeI.

Phenylether PhA'. [127]. Formed from NPhEt.CS.Cl and PhSH (B. a. S.). Needles.

PHENYL-ETHYL-THIO-SEMI-CARBAZIDE NHEt.CS.NH.NHPh. [122°]. Formed from ethylthiocarbimide and phenyl-hydrazine (Dixon, U. J. 55, 302). White crystals, v. sl. sol. water. FeCl, gives a red colour changing to blackish-

NHPh.CS.NH.NHEt. [110°]. Isomeride Formed from phenyl-thiocarbimide and ethylhydrazine (Fischer, A. 199, 296). Needles.

D1-phenyl-ethyl-thio-semi-carbazide NPhEt.NH.CS.NHPh. [149]. Formed from u-phenyl-ethyl-hydrazine and phenyl-thiocarbinide (Michaelis a. Philips, A. 252, 273).

PHENYLETHYL-THIOCARBIMIDE C.H.Ph.NCS. Formed from phenylethyl-amine by successive treatment with CS2 and HgCl2

(Neubert, B. 19, 1825). Yellow oil.

ETHÝL PHENYL DITHIOCARBONATE CS(OEt)(SPh). Formed by mixing solutions of C₆H₅.N₂Cl and potassium ethyl dithiocarbonate at 0° (Leuckart, J. pr. [2] 41, 186). Reddishyellow oil, sl. sol. water, slightly volatile with steam. Converted by heating with alcoholic NH3 into phenyl mercaptan and ammonium sulphocyanide.

PHENYLETHYL-THIOHYDANTOÏN

 $C_{\rm H,Ph.N:}C<_{
m NH.CO}^{
m S-CH_2}$ $\mathbf{C}_{11}\mathbf{H}_{12}\mathbf{N}_{2}\mathbf{SO}$

CS N(C₂H₄Ph).CH₂ CO. The hydrochloride B'HCl ĆO [188°] is formed from phenylethyl-thio-urea and chloro-acetic acid (Neubert, B. 19, 1822).

PHENYL-ETHYL-THIOPHENE C12H12S i.e. s<CH:CH Obtained by heating [40°]. CH2Bz.CHEt.CO2Na with P2S3 (Dittrich a. Paal, B. 21, 3457). Small plates. Gives a cherryred colour with isatin and H2SO, and a bluishgreen colour with phenanthraquinone and HOAc.

u-PHENYL-ETHYL-THIO-UREA CaH,2N,S i.e. NPhEt.CS.NH₂. [113°]. Formed from ethylcyanide (Gebhardt, B. 17, 2094). Large pearly prisms (from alcohol).

Benzoyl derivative NPhEt.CS.NHBz. [134°]. Formed from ethylaniline and benzoylthiocarbimide (Dixon, C. J. 55, 305). Pale lemon-yellow prisms, insol. water, sol. alcohol.

NHPh.CS.NHEt. s-Phenyl-ethyl-thio-urea [99°]. Formed from phenyl-thiocarbimide and NH.Et and from ethyl-thiocarbimide and aniline (Weith, B. 8, 1524; Michael a. Palmer, Am. 6, 260). Monoclinic crystals.

Phenylethyl-thio-urea NH_CS.NH.C2H,Ph. [123°]. Formed from phenylethyl-amine hydrochloride and potassium sulphocyanide (Neubert, B. 19, 1822). Plates (from dilute alcohol).

Phonyl-ethyl-\psi-thio-urea NHPh.C(SEt):NH. Formed from phenyl-thio-urea and Etl (Bertram, B. 25, 55). Yields mercaptan on treatment with alkalis.—B'HI. [103°].—B'C6H3N8O7. [196°].

Phenyl-di-ethyl-\(\psi\)-thio-urea

NPhEt.C(SEt):NH. Formed from the preceding
body and EtI (B.).—B'HI.—B'_\(\psi\)-\(\p

 $B'C_0H_3N_3O_7$. [170°]. Phenyl-tri-ethyl- ψ -thio-urea

NPhEt.C(SEt):NEt. (c. 275°). Got from the preceding and EtI (B.). B'C, II, N, O, [c. 96°]. B'2H2PtCl. [135°]

Di-phenyl-ethyl-thio-urea NHPh.CS.NPhEt. [89°]. Formed from phenyl-thiocarbinide and ethyl-aniline (Gebhardt, B. 17, 2090). Crystals.

Di-phenyl-ethyl-ψ-thio-urea

NHPh.C(SEt):NPh. [79]. Formed from diphenyl-thio-urea and EtBr (Rathke, B. 14, 1776) and by the action of mercaptan on C(NPh), in the cold (Will, B. 15, 1308). Needles (from dilute alcohol). Decomposed by heat into C(NPh)₂ and EtSH. Cl passed into a solution of its hydrochloride yields ethane sulphonic acid. _B'HCl. _B'_HPtCl_2aq. - B'HI aq. [157.5°] (Bernthsen, B. 15, 266, 567).

Di-phenyl-di-ethyl-thio-urea CS(NEtPh)2. [75.5°]. Formed from NPhEt.CSCl and ethylamline at 100° (Billeter, B. 20, 1631). White tables (from ligroin) or needles (from alcohol). Di-phenyl-di-ethyl-thio-urea

CS(NH.C,H₄Ph)₂. [84°]. Formed from phenylethyl-amine and alcoholic CS₂ (Neubert, B. 19, 1824). Plates (from alcohol), insol. water.

Acetyl derivative C19H22N2SO.

 $Di-phenyl-di-ethyl-\psi-thio-urea$ NPhEt.C(SEt):NPh. Oil. The hydro-iodide. got by heating di-phenyl-ethyl-ψ-thio-urea with EtI at 130°, is crystalline (B. a. F.).

u-PHENYL-ETHYL-UREA NPhEt.CO.NH2-[62°]. Formed from ethyl-aniline hydrochloride and potassium cyanate (Gebhardt, B. 17, 2095).

s-Phenyl-ethyl-urea NPhH.CO.NEtH. [99°]. Formed from ethyl cyanate and aniline (Wurtz, C. R. 32, 417). Needles (from dilute alcohol). Yields a nitrosamine NPhH.CO.NEt.NO [60°] crystallising in monoclinic prisms (E. Fischer, A. 199, 286).

Phenylethyl - urea NH2.CO.NHC2H4"h. [112°]. Formed from β -phenyl-ethyl-amine and potassium cyanate (Spica, G. 9, 568). Flat prisms, m. sol. cold water.

Phenyl - di - ethyl - urea NHPh.CO.NEt, [85°]. For ned from phenyl cyanate and NHEt. (Gebhardt, B. 17, 3039). Needles.

Di-phenyl-ethyl-urea NHPh.CO.NPhEt. [91°]. Got from phenyl cyanate and ethylaniline (Gebhardt, B. 17, 2093). Prisms.

s - Di - phenyl - di - ethyl - urea CO(NPhEt)2. [79°]. Formed from NPhEt.COCl and ethylaniline at 130° (Michler, B. 9, 712). Crystals.

NPh. CO.NEt2. u-Di-phenyl-di-ethyl-urea [54°]. Formed from NPh2.COCl and NHEt

Diphenyldiethyl-urea NH₂.CO.N(C₂H₄Ph)₂. [109°]. Formed from diphenyldiethylamine hydrochloride and potassium cyanate in aqueous solution (Spica, G. 9, 568). Prisms, sol. hot Aq.

Tri - phenyl - ethyl - urea NPh2.CO.NPhEt. Formed from NPh, COCl and ethylaniline (M.). Formed also from NPhEt.COCl and diphenylamine (Kaufmann, B. 14, 2185).

PHENYL-FORMAMIDE v. vol. ii. p. 568. DI-PHENYL-FORMAMIDINE C13H12N2 i.e. CH(NPh)(NHPh). Di-phenyl-methenyl-diamine.

Mol. w. 196. [138°]

Formation.—1. By heating chloroform with aniline for 12 hours at 190° (Hofmann, Pr. 9, 229).—2. By heating aniline with phenyl-carbamine, with orthoformic ether, or with formic acid (Weith, B. 9, 454; Wichelhaus, B. 2, 116).—3. By passing gaseous HCl into heated formic anilide (Wallach, B. 15, 208).—4. From formic anilide and PCl, (Wallach, A. 214, 233). 5. From CH(NPh)SEt and aniline (Wallach s. Wüsten, B. 16, 146).-6. From aniline and CH(NH2Cl)OEt (Pinner, B. 16, 358).

Properties.—Needles (from ether). Yields

B'HCl and B'2H2PtCl6.

PHENYL-FORMAMIDOXIM C, H, NO i.e. CH(NOH).NHPh. [116°]. Formed from thio-formanilide and hydroxylamine (Muller, B. 22, 2411). Needles, m. sol. water.—B'HCl: needles. —B'₂H₂PtCl₆: yellow needles.

Benzoyl-derivative CH(NOBz)(NHPh).

[145°]. Needles, m. sol. alcohol and ether.
DI-PHENYL-FORMAZIDINE C₁₃H₁₄N₄ i.e.
CH(N₂HPh).N₂H₂Ph. [185°]. Formed from CH(NH2Cl).OEt and phenyl-hydrazine (Pinner, B. 17, 2002). Yellow plates, v. sol. hot alcohol. PHENYL-FORMYLACETIC ETHER

CHO.CHPh.CO₂Et. (145° at 16 mm.). Got, together with an isomeride [71°], by the action of NaOEt on a mixture of formic and phenylacetic ethers (Wislicenus, B. 20, 2931). Liquid. FeCl₃ colours its alcoholic solutions bluishviolet. Phenyl-hydrazine forms oxy-di-phenylpyrazole.

DI-PHENYL-FUMARAMIC ACID v. FUMARIO ACID

PHENYL-FUMARIC ACID C10H8O4 i.e. CO₂H.CPh:CH.CO₂H. [161°]. Got by heating bromo-cinnamic ether with alcoholic KCy for 8 hours at 150° (Barisch, J. pr. [2] 20, 186). Got also by heating CO₂H.CPh(OH).CH₂.CO₂H (Alexander, A. 258, 82). Nodules (from water).-BaA": scales.—Ag,A": amorphous pp.

Di-phenyl-fumaric acid CO₂H.CPh:CPh.CO₂H. [c. 260°]. Formed from its ether, which is got, together with di-phenylnaleic ether, by the action of Na on a-bromophenyl-acetic ether (Rügheimer, B. 15, 1626). Crystals, decomposing at 260° into water and di-phenyl-maleïc anhydride.

Nitrile CN.CPh:CPh.CN. [158°]. Got by adding alcoholic NaOEt to a-chloro-phenylacetonitrile (Michael a. Jeanprêtre, B. 25, 1680). DI-PHENYL FURAZANE CPh:N O. [947].

Formed by heating the (a)-di-oxim of benzil with water at 210° (Dodge, A. 264, 180). Crystals. At 300° it forms di-benzenyl-azoxim. H₂SO₄ and HNO3 give a di-nitro- derivative [220°].

CH:CPh ch:crh>0. DI-PHENYL-FURFURANE

[91°]. (345°). Formed by heating CPh:C.CH(CO₂H).CO.C₆H, or CH₂Bz.CH₂Bz with conc. HClAq at 160° for 3 hours (Kapf a. Paal, B. 21, 1490, 3057). Got also by heating its dicarboxylicacid (W. H. Perkin, jun., a. Schlosser, C. J. 57, 954). Plates, insol. water.

Tetrahydride C₁₆H₁₆O. (321°). Got by

reducing with Na and alcohol. Oil. Octohydride C16H20O. Oil.

Tri-phenyl-furfurane $C_{22}H_{16}O$ i.e.

CPh:CH O. [93°]. Mol. w. 263 by Raoult's method (calc. 296). Formed by boiling CPhBz:CHBz with HIAq (Japp a. Klingemann, C. J. 57,675). Got also from desyl-acetophenone by successive treatment with H₂SO₄ and water (Smith, C. J. 57, 645). Needles (from alcohol). Tetra-phenyl-furfurane v. Lepiden.

DI-PHENYL-FURFURANE CARBOXYLIC

ACID CO₂H.C:CPh O. $[217^{\circ}].$ Formed by boiling CPh:C.CHBz.CO2H with conc. HClAq and alcohol (Kapf a. Paal, B. 21, 1489, 3059). Formed also by heating the dicarboxylic acid (W. H. Perkin, jun., a. Schlosser, C. J. 57, 952).

Needles (from alcohol), insol. water. May be distilled. Br yields $C_{17}H_{12}Br_4O_3$.—NaA': needles. Ethyl ether EtA'. [82°]. Formed by boiling $CH_2Bz.CHBz.CO_2Et$ with alcohol and HClAq. Crystals (from ether). Yields on reduction a tetrahydride which does not react

with AcCl.

Di-phenyl-furfurane dicarboxylic C₁₈H₁₂O₅ i.e. CO₂H.C:CPh O. [238°]. Formed by dissolving di-benzoyl-succinic ether in H2SO4 and boiling the product with alcoholic potash (W. H. Perkin, jun., C. J. 47, 262; 49, 168; 57, 951). Needles, sol. alcohol and ether. FeCl, gives an orange colour in its alcoholic solution. Yields acetophenone on distilling with sodalime. $-Ag_2A''$.

Ethyl ether Et₂A". [86°]. Formed from dibenzoyl-succinic acid and H₂SO₄. Yellow prisms, v. sol. alcohol. Its solution in H₂SO₄

turns violet on heating.

Anhydride C₁₀H₁₀O₄. [255°]. Formed by heating the acid alone or with Ac₂O. Plates, sl. sol. hot alcohol. Its alcoholic solution ex-

hibits violet fluorescence.

PHENYL - FURFURYL - ACRYLIC Nitrile. C₄H₃O.CH:CPh.CN. [43°]. Formed from phenyl-acetonitrile, furfuraldehyde, and NaOEt in alcohol (Frost, A. 250, 159). Yields C4H3O.CHBr.CBrPh.CN [114°] crystallising in orange plates.

Reference. - NITRO-PHENYL-FURFURYL-ACRYLIC

PHENYL - FURFURYL - (β) - NAPHTHO -TRIAZINE. Dihydride $C_{21}H_{15}N_{2}O$ $C_{10}H_{4} < \stackrel{N.CH.C}{N.NPh} \stackrel{H_{8}O}{.}$ [241°]. Formed warming benzene-azo-(\$)-naphthylamine with

furfuraldehyde in alcohol (Goldschmidt a. | Poltzer, B. 24, 1007). Needles. -B'HCl: plates. -B'2H2PtCl6. Light-yellow crystalline pp.

 (β) - PHENYL - (γ) - FURFURYL - PROPYL -AMINE C.H.O.CH.CHPh.CH.NH2. (283°). Formed from phenyl-acetonitrile and furfuraldehyde, the product of condensation of these bodies C4H3O.CH:CPh.CN being reduced by Na and dry alcohol (Freund a. Immerwahr, B. 23, 2850). Oil, with alkaline reaction. verted by nitrous acid into the liquid alcohol C4H3O.CH2.CHPh.CH2OH. Phenyl thiocarbimide forms C,H,O.C,H,Ph.NH.CS.NHPh [113°]. -B'HCl.[176°]. Crystalline, v. sol. water. with potassium cyanate forming C₄H₃O.C₈H₅Ph.NH.CO.NH₂ [101°]. — Mercury double salt: [175°]; needles. — Picrate [152°]. Yellow crystalline powder.

PHENYL-FURIDANE. Dihydride

 $\operatorname{CPh} \leqslant_{\mathrm{O-CH}}^{\mathrm{CH.CH}} > \operatorname{CH}_{2}.$ (250° at 721 mm.). Formed by heating its carboxylic acid at 200° (Perkin, C. J. 51, 731). Oil. HBr forms C_bH_s,CO.C_lH_sBr.

PHENYL-FURIDANE DIHYDRIDE CARB- $CO_2H.C \leqslant^{\operatorname{CH}_2.\operatorname{CH}_2}_{\operatorname{CPh.O}} \searrow \operatorname{CH}_2.$ OXYLIC ACID

Phenyldehydrohexone carboxylic acid. [c. 144°]. Formed by saponifying its ether, which is got by the action of NaOEt on a mixture of trimethylene bromide and benzoyl-acetic ether (Perkin, C. J. 51, 726). Monoclinic prisms (from ether); a:b:c = 2.638:1:3.398; $\beta = 74^{\circ} 44'$. -AgA': needles (from water).

Ethyl ether EtA'. [60°]. Prisms.

PHENYL-GLUTARIC ACID C₁₁H₁₂O₄ i.e. CHPh(CH2.CO2II)2. [138°]. Formed by heating sodium malonic ether with alcohol and cinnamic ether at 100°, saponifying the resulting ether CH(CO_Et)_cHPh.CHNa.CO_Et, and heating the acid at 110° (Michael, J. pr. [2] 35, 352; Am. 9, 116). Needles, sl. sol. water, m. sol. alcohol.—AgA': amorphous pp.

Di-phenyl-glutaric acid CH2(CHPh.CO2H)2. [164°]. Formed by saponifying its nitrile (Zelinsky a. Feldmann, B. 22, 3292). Needles.

Nitrile CH₂(CHPh.CN)₂. [71°]. Formed by heating CH₂Ph.CN with CH₂I₂ and dry NaOH. PHENYL-GLYCERIC ACID v.

PHENYL-PROPIONIC ACID. TRI-PHENYL GLYCERYL TRIKETONE CH_Bz.CHBz.CH_Bz. Tri-benzoyl-propane.

[137°]. Formed from tricarballylic chloride, benzene, and AlCl₃ (Emery, B. 24, 601). Strawyellow needles. Yields a phenyl-hydrazide [57°-60°]

PHENYL-GLYCIDIC ACID C,H,O, i.e. O CHPh CH.CO.H. Phenyl-pyruvic acid? [155°]

Formed by boiling benzoyl-imido-phenyl-propionic acid NBz CHPh with aqueous HCl or with KOH (Plöchl, B. 16, 2817; 19, 3167). Formed also by boiling phenyl-oxalacetic acid with dilute H₂SO₄ (W. Wislicenus, B. 20, 592). Plates, v. e. sol. alcohol and ether. FeCl₂ colours its alcoholic solution green. Sodiumamalgam reduces it to a-oxy-phenyl-propionic acid. Yields an oxim and a phenyl-hydrazide CH_Ph.C(N_HPh)CO.H [1619]. Tolylene-o-diamine forms a quinoxaline (Erlenmeyer, jun., B. 19, 2576; 20, 2465; 22, 1482). Aniline forms C2H2Ph(NHPh)(OH).CO2H.

(β)-Phenyl-glycidic acid. Formed from C₆H₅.CH(OH).CHCl.CO₂H and cold alcoholic potash (Glaser, A. 147, 98). Oil, crystallising at 0°. Decomposes, even at 15°, into phenylacetic aldehyde and CO2. Boiling dilute H2SO4 forms CHPh(OH).CH(OH).CO2H and phenylacetic aldehyde (Erlenmeyer a. Lipp, A. 219, Yields β-oxy-phenyl-propionic acid on reduction with sodium-amalgam.-NaA'.-KA'. -AgA': crystalline powder.

Ethyl ether EtA'. (280° cor.). Oil.

References .- NITRO- and OXY- PHENYL-GLY-CIDIC ACID.

PHENYL-GLYCOCOLL v. PHENYL-AMIDO-ACETIC ACID.

PHENYL-GLYCOL v. DI-OXY-ETHYL-BENZENE. PHENYL-GLYCOLLIC ACID v. Phenylderivative of GLYCOLLIC ACID and MANDELIO

Di-phenyl-glycollic acid v. Benzilic Acid. PHENYL-GLYCOLLIC ACRYLIC ACID v. Carboxy-methyl derivative of Coumaric acid.

PHENYLGLYCOLYL-TROPEINE C10 H21NO3. Homo-atropine. [98°]. Formed by heating tropine mandelate with HClAq (Ladenburg, A. 217, 82). Deliquescent prisms (from ether), m. sol. water. Less poisonous than atropine.— B'HAuCl.—B'HBr.—B'C₆H₂N₃O₂. Yellow plates. TETRA-PHENYL-GLYCOSINE v. GLYCOSINE.

PHENYL-GLYOXAL C.H., CO.CHO. (142° at 125 mm.). Formed by allowing a mixture of its mono-oxim (30 g.) with NaHSO, (120 g. of a 35 p.c. solution) to stand till all is dissolved, and then adding H.SO, and distilling (Müller a. von Pechmann, B. 20, 2904; 22, 2557). Oil. Forms a crystalline hydrate [73°]. Converted by HNO, into phenyl-glyoxylic acid, and by potash into mandelic acid. Ammonia forms $C_{22}H_{19}N_3O$ or $C_{22}H_{17}N_3O$ crystallising in plates [1933] which can be distilled. Hydroxylamine gives C₁₆H₁₃N₈O₃, [219°], sol. alkalis.

Mono-oxim C,H,.CO.CH:NOH. Nitrosoacctophenone. [128°]. Formed from acetophenone, isoamyl nitrite, and alcoholic NaOEt (Claisen, B. 20, 656, 2194; Braun, B. 22, 556). Monoclinic tables (from chloroform), sl. sol. cold water, v. sol. alkalis. Yields C.H.CO.CN on warming with Ac₂O. Boiling NaOHAq forms NaCy and NaOBz. Its acetyl derivative is converted by NaOHAq into CHBz(OH).CO.CO.C₀H₄ [170°] crystallising in minute needles (Söder-

baum, B. 24, 1386, 3034).

Di-oxim C,H,.C(NOH).CH(NOH). phenyl-amphi-glyoxim. [162°] (S.); [168°] (R.). Formed by the action of hydroxylamine on the mono-oxim, or on mono- or di-bromo- acetophenone (Schramm, B. 16, 2183; Strassmann, B. 22, 419; Russanoff, B. 24, 3501). Small needles, v. sol. alcohol, insol. CHCl₃; subliming below 160°. N₂O₄ gives C₀H₅.C₂HN₂O₂ [c. 90°] crystallising in colourless prisms (Scholl, B. 23, 3504). By dissolving in ether and treating with HCl it is converted into an isomeride [180°]. Thin body, called phenyl-anti-glyoxim, is v. sol. Ac2O and yields a di-acetyl derivative [92°]. Phenylanti-glyoxim is only stable in acid solutions being readily re-converted into the original di-oxim [168°]. By treatment of the di-oxim [168°] NaOHAq and CO₂ at -10° there is formed a

glyoxim which yields an oily di-acetyl derivative.—AgC,H,N,O,. Curdy pp.

Phenyl-hydrazide C,H₅.C(N,HPh).CHO.

[143°]. Yellow plates (M. a. P.).

Phenyl-hydrazide C,H, CO.CH:N2HPh. [129°]. Formed by the action of warm dilute KOHAq on the compound got by treatment of sodium benzoylacetic ether with diazobenzene chloride (Stierlin, B. 21, 2123). Yellow plates.

p-Tolyl-hydrazide CHBz:N2H.C6H,Me. Got in like manner, using p-diazotoluene chloride. Yellowish-brown plates, v. sol. ether.

Di-phenyl-di-hydrazide C_6H_5 .C(N_2HPh).CH(N_2HPh). [152°]. heating phenyl-glyoxal or, better, the phenylhydrazide of benzoyl-carbinol with phenylhydrazine hydrochloride and NaOAc at 100° (Laubmann, A. 243, 247). Yellow plates.

Di-phenyl-di-methyl-di-hydrazide C₆H₅.C(N₂MePh).CH(N₂MePh). [151°]. Formed from ω-bromo-acetophenone and phenyl-methylhydrazine (Culmann, B. 21, 2597). Prisms.

Di-phenyl-glyoxal v. Benzil. PHENYL-GLYOXALINE C, H, N, i.e.

 $C_6H_5N <_{\mathrm{CH:N}}^{\mathrm{CH:CH}}$ [13°]. (276°). Formed by warming sulphydro-phenyl-glyoxaline [181°] with dilute HNO₃ or HClAq (Wohl a. Marckwald, B. 22, 576, 1353). Crystalline, miscible with alcohol.—B'HAuCl₄.—B'₂H₂PtCl₆.—Picrate B'C₆H₃N₃O₁. [152°]. Yellow needles (from elabels). Bean Different Properties of the control of th alcohol).—B2AgNO3.—B'MeI. Syrup.

Phenyl-glyoxaline NH CH :CH [148°]. (c. 340°). Formed by distilling its dicarboxylic acid, which is got from di-nitro-tartaric acid, benzoic aldehyde, and NH, (Maquenne, C. R. 111, 742). Plates (from benzene).—B'2H2C2O4. Needles.—B'2H2PtCl6.

Di-phenyl-glyoxaline $CH \leq_{N-CPh}^{NH.CPh}$. [227°]. Formed by the action of formic aldehyde and ammonia on benzil (Japp, C. J. 51, 559). Dimorphous. Crystallises with di-phenyl-methylglyoxaline as (C₃N₂H₂Ph₂)C₃N₂HMePh₂ [218°]. —B'₂H₂PtCl₆: small flat yellow needles.

Tri-phenyl-glyoxaline v. Lophine, vol. i.

PHENYL-GLYOXIM v. Oxim of PHENYL-

GLYOXAL PHENYL-GLYOXYLIC ACID C8H6O3 C₆H,.CO.CO₂H. Benzoyl-formic acid. [66°].

Formation .- 1. From benzoyl cyanide and conc. HClAq in the cold (Claisen, B. 10, 429, 844, 1663; 12, 626, 1505) or HOAc saturated with HCl (Buchka, B. 20, 395).—2. From dioxy-ethyl-benzene (phenyl-glycol) and nitric acid of S.G. 1:38 (Zincke, A. 216, 305).—3. By saponifying its ether, which is got by heating COCl.CO₂Et with HgPh₂ for several hours at 150°; the yield being 40 p.c. of the theoretical (Claisen a. Morley, B. 11, 1596).—4. By oxidising acetophenone with alkaline KMnO, at 0° the yield being 20 p.c. of the theoretical amount (Glückmann, M. 11, 248).

Properties.—Prisms, v. e. sol. water and ether. Decomposed by distillation into benzoic acid and CO, and partly also into benzoic aldehyde and CO2. Phenyl-hydrazine added to its solution in dilute HCl forms a bulky yellow pp.

second isomeride [148°-154°] called phenyl-syn-! of phenyl-hydrazide (Ebers, A. 227, 340). When mixed with benzene (containing thiophene) it gives with conc. H2SO, a red colour changing to violet. On adding water, the layer of benzene becomes crimson.

Reactions. -1. Sodium-amalgam reduces it to mandelic acid.—2. HI and P at 160° reduce it to phenyl-acetic acid .- 3. Phenyl mercaptan and gaseous HCl form C, H, C(SPh), CO, H [142°], sl. sol. benzene (Baumann, B. 18, 891). Phenyl mercaptan alone forms a crystalline powder $C_6H_3.C(OH)(SPh).CO_2H$ [69°].

Salts. NaA'. KA' aq: dimetric tables. NH4A'.—CaA'2aq: flat prisms. —BaA'2.—SrA'2aq. $-\operatorname{CuA'}_2$, $-\operatorname{ZnA'}_2$ 2aq. $-\operatorname{PbA'}_2$, $-\operatorname{PbA'}_2$ aq. $-\operatorname{AgA'}$

Methyl ether MeA'. (248°).

Ethyl ether EtA'. (257°). S.G. 17.5 1.121. Forms a crystalline compound with NaHSO. Reacts in alcoholic solution with hydrazine hydrateforming C₄H₁₂N₄O₂[120°] and C₂₀H₂₀N₂O₄ [135°] (Curtius, J. pr. [2] 44, 566).

n-Propyl ether PrA'. (174° at 60 mm.).

Isobutyl ether. (174° at 38 mm.).

Isoamyl ether. (182° at 40 mm.).

(a)-Amide Coll., CO.CONI... [91°]. Formed from benzoyl cyanide and cold HClAq. On solution and re-ppn. by CO₂ it forms an unstable hydrate C₈H₇NO₂ aq [65°] (so-called (8)-amide).

 (γ) - A mide $(C_8H_7NO_2)_2$. [134°]. Formed by adding an alcoholic solution of the (a)-amide to dilute HClAq (Claisen). Formed also, together with the (a)-amide, by the action of HOAc saturated with HCl upon BzCy (Buchka). Prisms.

Nitrile v. Benzoyl Cyanide. (a) - Oxim C₆H₅.C(NOH).CO₂H. Isonitrosophenyl-acetic acid. [127°]. Formed by the action of hydroxylamine on the Na salt (A. Muller, B. 16, 1617) or on the acid at 0° (Hantzsch, B. 23, 2333; 24, 42). Thick prisms, v. e. sol. water. Yields α-amido-phenyl-acetic acid on reduction — KA'aq: very soluble crystals. -BaA'₂ 1¦aq.—AgA'.

Acetyl derivative. [119°]. Prisms. (β) - OximC,H,.C(NOH).CO,H. [145°]. Formed by the action of HCl on an ethercal solution of the (α) -oxim. Formed also by warming phenyl-glyoxylic acid with hydroxylamine and potash. White needles composed of dimetric prisms.

Acetyl derivative. [125°]. Tables.

Methyl ether of the oxim

CPh(NOH).CO Me. [139°]. Got from the Ag salt and MeI (Muller, B. 16, 2987). Converted by NaOEt and MeI into CPh(NOMe).CO.Me [56°], sol. alcohol, insol. alkalis.

Ethyl ether of the oxim CPh(NOH).CO₂Et. [113°]. Formed from phenylglyoxylic ether and hydroxylamine (Gabriel, B.

16, 519). Needles (from hot water).
Nitrile of the oxim C₆H, C(NOH).CN. [129°]. Formed from phenyl-acetonitrile, amyl nitrite, and alcoholic NaOEt (Frost, A. 250, 163; Meyer, B. 21, 1306). Formed also by the action of hydroxylamine on dibromo-acetophenone in dilute alcohol at 60° (Russanoff, B. 24, 3505).

yellow powder, v. sol. siconol.—C₆H₅.C(NONA).CN; yellow powder, v. sol. water. Yields an acetyl de-rivative [68°].—KA'.—CuA'₂.—Ph₂A'₂O.—AgA'. Phenyl hydrazide C₆H₅.C(N₂HPh).CO₂H₅. [153°]. (F.); [153°-164°] (Von Pechmann, B. 25, 1054). Yellow needles, sl. sol. hot water

Thin plates, v. sol. alcohol.—C.H. C(NONa).CN:

(Fischer, B. 17, 578; Ebers, A. 227, 340). May be reduced to C₆H₅.CH(N₂H₂Ph).CO₂H [158°], and finally to aniline and a-amido-phenyl-acetic

Phenyl-methyl-hydrazide

C₀H₅.C(N₂MePh).CO₂H. [116°]. Plates (from alcohol). Its amide C₀H₅.C(N₂MePh).CONH₂ |156°] is crystalline.

Phenyl-cthyl-hydrazide

 $C_{_0}H_{_5}.C(N_2EtPh).CO_2H.$ [109°]. Formed, together with its amide $C_{_6}H_{_5}.C(N_2EtPh).CO.NH_2$ [111°], by the action of phenyl-ethyl-hydrazine in dilute HOAc on phenyl-glyoxylic acid. Yellow plates, sl. sol. water. Yields ethyl-aniline and benzoic aldehyde when heated with conc. HClAq.

References. -- AMIDO-, NITRO-, and Oxy-

PHENYL-GLYOXYLIC ACID.

PHENYL - GLYOXYLIC o - CARBOXYLIC ACID C₆H₄(CO₂H).CO.CO₂H. [140°]. Formed by oxidising the oxyquinone C16H16O3 or indonaphthene dihydride carboxylic acid with alkaline KMnO₄ (Scherks, B. 18, 378; cf. Zincke, A. 226, 53; 240, 142). Yields CO₂ and phthalic anhydride on heating. May be reduced to phthalide carboxylic acid.—K2A".—BaA" 2aq. -Cu₂A"(OH)₂6aq.—Ag₂A": crystalline pp.

PHENYL-DIGUANIDE C.H.I.N. i.c.

NH:C(NHPh).NH.C(NH₂):NH. [c. 237°]. Formed by the action of alcoholic ammonia and AgNO₃ or HgCl₂ on guanyl-phenyl-thio-urea (Bamberger, B. 13, 1582). Formed also by heating aniline hydrochloride with di-cyan-di-amide in alcohol at 100° (Smolka a. Friedrich, M. 9, 232). Plates, v. sol. water and alcohol; absorbs CO, from air. When boiled with baryta-water it yields NH3, aniline, phenyl-urea [147°], phenylguanidine, guanidine, and urea (Emich, M. 12, 15).- B'HCl: prisms. Yields phenyl-carbamine

when warmed with alcoholic potash and CHCl₃.

Salts. — B'HNO₃. [209°]. — B'₂H₂SO₄. —

Cu(C₄H₁₀N₃)₂ 1½aq: red powder, sol. hot water.

— CuA'₂H₂Cl₂ 1½aq. — CuA'₂H₂SO₃ 1½aq. —

CuA'₂H₂CrO₄aq. — NiA'₂. — NiA'₂H₂Cl₂. — $NiA'_{2}H_{2}SO_{1}I_{2}^{1}aq.-CoA'_{2}I_{2}^{1}aq.-CoA'_{2}H_{2}CI_{2}^{1}aq.$

--CoA'.H.,SO, aq: crimson needles.
(a)-Di-phenyl-diguanide C₁₁H₁,N₅ i.e.
NPh:C(NHPh).NH.C(NH₂):NH. Formed by the action of AgNO3 and aniline on guanyl-phenylthio-urea (Bamberger, B. 13, 1584). White solid, m. sol. water and alcohol, alkaline in reaction.—B'HNO, [231°]. White needles. (B)-Di-phenyl-diguanide

NH:C(NPh₂).NH.C(NH₂):NH. [c. 162°]. Formed by heating diphenylamine hydrochloride with di-cyan-di-amide (Emich, M. 12, 21). Thin needles, with alkaline reaction.—B'HNO3. [203°].

Prisms.— $B'_23H_2SO_4$. Tri-phenyl-diguanide C20H19N5. Formed from guanyl-tri-phenyl-thio-urea, NH3, and AgNO₃. Formed also from di-phenylguanidine and phenyl-cyanamide [78°] in ether (Rathke a. Oppenheim, B. 23, 1672). Prisms (from alcohol). With CS₂ at 100° it yields triphenyl-thio-ammeline C₂,H₁,N₄S. — B'HCl:

prisms, v. sol. alcohol.—B'2H2PtClo.

PHENYL-GUANIDINE C, H, N, i.e. NH:C(NH2)(NHPh). Formed by desulphuration of phenyl-thio-urea in presence of alcoholic NH, (Feuerlein, B. 12, 1602). Formed also by the action of baryta-water on phenyl-diguanide

(Emich, M. 12, 11). Decomposes on standing into phenylcyanamide and ammonia. - Picrate B'C₆H₈N₃O₇. [208°-214°] (Prelinger, M. 13, 99).

Di-phenyl-guanidine C13H13N3 i.e.

NH:C(NHPh)₂. Melaniline. [147°]. S. (90 p.c. alcohol) 9.25 at 21°

Formation .- 1. From aniline and CyCl or CyBr (Hofmann, A. 67, 129; 74, 8; Weith a. Ebert, B. 8, 912). -2. By the action of PbO on a solution of di-phenyl-urea in alcoholic NH, (Hofmann, B. 2, 452, 688; Weith, B. 7, 937; Rathke, B. 12, 772).—3. By boiling phenyl-thiourea with HgCl(NHPh) and alcohol (Forster, B. 7, 294; A. 175, 35).—4. Together with thiourea, by the action of aniline on mercuric fulminate (Steiner, B. 7, 1244; 8, 518).- 5. By combination of di-phenyl-cyanamide with NH, (Weith, B. 7, 10).

Properties. - Monoclinic prisms (Arzruni, I'.

152, 284), sl. sol. cold water.

Reactions .- 1. Ac2O at 100° yields acetylphenyl-urea [183]. At 150° the product is acetyl-di-phenyl-urea [115°] (McCreath, B. 8. 1181).—2. Phenyl thiocarbinide forms, in the cold, NHPh.CS.NPh.C(NH).NHPh [150°], a weak base, sl. sol. benzene.-3. HClAq at 250° forms CO₂, NH₃, and aniline.—4. Cyanogen passed into its alcoholic solution forms C15H13N, which crystallises from alcohol in yellowish needles [154°], decomposed by HClAq into crystalline melanoximide C13H11N,O2, which is split up by alcoholic potash into oxalic acid and diphenylguanidine, and by alcoholic HCl into NH, and di-phenyl-parabanic acid.

Salts. The hydrochloride is gummy. - B'HAuCl₁: golden needles. - B'₂H₂PtCl₀. --B'HBr: stellate groups of needles, v. sol. water. - B'HI. - B'HNO₁. - B',AgNO₃: geodes. - B',H₂SO₄: stellate groups of plates. - B',H₂C₂O₄. -Sulphocyanido: [115°]; m. sol. water.

derivative C27H21N.O. Di-benzoyl[102°]. Monoclinic (McCreath, B. 8, 383).

Tri-phenyl-guanidine $C_{19}H_{17}N_3$ i.e. NPh:C(NHPh)₂. Mol. w. 287. [143°]; [147°] (F. a. W.). S. (alcohol) 4.5 at 0°.

Formation.—1. By adding I to an alcoholic solution of di-phenyl-thio-urea alone or mixed with aniline (Hofmann, B. 2, 453). - 2. From diphenyl-thio-urea and HgCl(NHPh) (Forster, B. 7, 294).—3. From di-phenyl-cyanamide C(NPh), and aniline (Weith, B. 7, 10). -4. Together with phenyl-thiocarbimide by heating di-phenylcyanamide with di-phenyl-urea (Weith, B. 9, 810).—5. By boiling di-phenyl-urea with inverted condenser (Barr, B. 19, 1765).—6. By heating di-phenyl-thio-urea with aniline, with Cu, or with PbCl₂ (Merz a. Weith, Z. [2] 4, 513, 609; 5, 583, 659; Girc d, Bl. [2] 46, 506), or with HgCl, at 145° (Buff, B. 2, 498).—7. By heating di-phenyl-cyanamide (2 mols.) with phenylene-o-diamine (1 mol.) at 210° (Keller, B. 24, 2505). 8. By the action of aniline on CCl, or chloropicrin (Hofmann; Basset, C. J. 18, 31).—9. By the action of PCl₂ (1 mol.) on a mixture of diphenyl-urea (3 mols.) and aniline (3 mols.)—10. By the action of PCl_a (2 mols.) and CO_a (3 mols.) on aniline (9 mols.) at 170° (Merz a. Weith, Z. [2] 6, 160).—11. By fusing di-phenylurea with NaOH (Hentschel, J. pr. [2] 27, 500).

12. From phenyl thiocarbimide by chlorinating

and heating the product NPh:CCl2 with aniline (Sell a. Zierold, B. 7, 1231). The base got is possibly an isomeric tri-phenyl-guanidine.-13. A product of the action of CSCl₂ on CS(NHPh)₂ and benzene (Freund a. Wolf, B. 25, 1464).

Properties.—Trimetric six-sided prisms (from alcohol); a:b:c = .670:1: .558, nearly insol. boiling water. Occurs also in an amorphous form (Giraud, Bl. [2] 46, 505). Decomposed above 250° into aniline and C(NPh)2. KClO, (12 pt.) and HCl give a greenish solution, depositing dark flakes which dissolve in alcohol with violet colour.

Reactions.—1. CS, at 170° forms CS(NPhH), and phenyl thiocarbimide. - 2. H.S passed through it at 170° forms aniline and di-phenylurea (Merz a. Weith, Z. [2] 6, 72).—3. Water at 175° forms aniline and di-phenyl-urea.—4. Conc. KOHAq or conc. HClAq yield aniline and CO₂ on heating.—5. Cyanogen passed into the alcoholic solution forms crystals of C21H17N which is decomposed by HClAq into NH, and the oxalyl-derivative C(NPh) NPh.CO [c. 230°] which is itself decomposed by boiling with alcohol and HClAq into aniline and di-phenylparabanic acid (Hofmann, B. 3, 764; Stojentin, J. pr. [2] 32, 1).—6. Chloro-glyoxylic ether COOLCO_Et forms a carbonyl derivative C(NPh) < NPh > CO which forms the salts B'HCl [190°] and B'HNO, [185°] (Stojentin). — 7. COCl₂ forms the above (?) carbonyl derivative C₂₀H₁₀N₂O, which crystallises from CS₂ in tables [184°] and forms a hydrate B'aq [141°] (Michler a. Keller, B. 14, 2181).

Salts.—B'HOl aq. [242°] (Weith, B. 9, 810); [245°] (F. a. W.). S. 2 at 0°.—B'HNO;: laminæ.—B'₂H₂PtCl₈. S. ·09 at 0°.—B'H₂SO₄: broad needles.—B'HOAc.—B'H₂C₂O₄: laminæ.— B'C, H, N,O,. [178°]. S. . 0082 at 15° (Prelinger, M. 18, 98).

AcetylderivativeC(NPh)(NPhAc)₂.

[131°]. Crystals (MacCreath, B. 8, 383).

Benzoyl derivative C(NPh)(NPhBz)₂.

[185°]. Plates (from alcohol).

(B)-Tri-phenyl-guanidine C(NH)(NHPh)(NPh2). [131°]. Formed heating phenyl-cyanamide (cyanilide) CN.NHPh with diphenylamine at 125° (Weith a. Schröder, B. 8, 912). Regular tablets, nearly insol. water. Its solution in conc. H2SO, becomes violet on heating.

Reactions.-1. Conc. HClAq or KOHAq at 260° forms CO₂, NH₂, aniline, and NHPh₂.—2. CS₂ at 160° forms the sulphocyanide, H₂S, phenyl-thiocarbimide and NHPh₂.

Salts .- B'HCl aq. S. (of B'HCl) 28.4 at

23°.-B'2H2PtCl6: crystalline.

Dicyanide C21H17N3. [173°]. A product of the action of cyanogen as an alcoholic solution of aniline (Hofmann, A. 66, 129; B. 3, 763). Got also by boiling the compounds of cyanogen with di-phenyl-guanidine or (a)-triphenyl-guanidine with aniline hydrochloride (Landgrebe, B. 10, 1593; 11, 973). Dark-brown needles (containing aq) with violet iridescence (from alcohol).—B'HCl Baq: brown needles.

Tetra - phenyl - guanidine C(NH)(NPh₂)₂.

[131°]. Formed by passing gaseous cyanogen

chloride through diphenylamine at 160° (Weith, B.7, 843). Pyramids (from ligroin), insol. water, v. sol. alcohol and ether. Conc. H2SO4 forms a colourless solution turned violet-red by heat. CS₂ at 260° forms tetra-phenyl-thio-urea. Solid KOH at 200°, and conc. HClAq at 840°, yield NH₃, CO₂, and NHPh₂.

Salts.-B'HCl 5aq. Efflorescent tablets.-B',H,PtCl,-B'HNO,

References. - DI-BROMO-, DI-CHLORO-, DI-IODO-,

and Nitro- Phenyl-Guanidines.

DI-PHENYL-GUANIDINE CARBOXYLIC ACID C(NH)(NHPh).NH.C₆H₄.CO₂H. Phenylguanido-benzoic acid. [165°]. Formed by heating m-cyanamido-benzoic acid or cyancarbimidoamido-benzoic acid with aniline (Traube, B. 15, 2120; Griess, B. 16, 836). Crystals (containing aq), nearly insol. alcohol and ether, sol. hot water, aqueous alkalis and acids.—HA'HCl aq.

Reference. — Amido - di - Phenyl - Guanidine

CARBOXYLIC ACID.

PHENYL-GUANIDO-ACETIC ACID

C_pH₁₁N₃O₂ i.e. C(NH)(NHPh).NH.CH₂.CO₂H. [260° uncor.] Formed from phenyl-cyanamide and glycocoll (Berger, B. 13, 9°2). Crystals. DI-PHENYL-DI-GUANIDO-DI-TOLYL-SUL-

S(C,Hc.NH.C(NH).NHPh)2. PHIDE tolyl-di-phenyl-di-guanidine. [153°]. Formed heating di-phenyl-di-uramido-di-tolyl sulphide with alcoholic NH, and HgO (Truhlar, B. 20, 675). Small white needles (from ether-ligroïn). V. sol. alcohol, benzene, and ether.— B"H2Cl2PtCl4: yellow amorphous powder.

Tetra-phenyl-di-guanido-di tolyl-sulphide S(C,H_s,NH.O(NPh),NHPh)_s. Thio-p-tolyl-tetra-phenyl-di-guanidine. [106°]. Formed by heat-ing di-phenyl-di-uramido-di-tolyl sulphide with aniline and HgO (T.). Amorphous powder. PHENYL-HEPTADECYL-THIO-UREA

C₆H₅NH.CS.NHC₁₇H₃₅. [79°]. Formed by boiling heptadecyl-thiocarbimide with aniline and alcohol (Turpin, B. 21, 2491)

PHENYL-HEPTADECYL-UREA C24H42N2O i.e. C₆H₅NH.CO.NHC₁₇H₂₅. [99°]. Formed by heating heptadecylamine hydrochloride with COCl₂ in benzene at 100°, the product being heated with aniline (Turpin, B. 21, 2492).

PHENYL-HEPTANE v. HEPTYL-BENZENE. Di-phenyl-heptane C₆H₁₃.CHPh₂. [14°]. (191° at 13 mm.). Formed from C.H.13.CHCl2, benzene, and AlCl₃ (Auger, Bl. [2] 47, 48; Krafft, B. 19, 2986). Needles.

PHENYL-HEPTENOIC ACID C13H18O2 i.e. Ph.CEt:CEt.CO₂H (?) Oil. Got by passing CO over a heated mixture of NaOEt and sodic cinnamate (M. Schroeder, A. 221, 52)

PHENYL HEPTYL KETONE CARBOXYLIC ACID C_6H_5 .CO.CH₂.CH(C_6H_{11}).CO₂H. Formed by heating phenacyl-isoamyl-malonic acid (Paal a. Hofmann, B. 23, 1504). Plates, insol. water, sl. sol. ligroin. On repeated distillation it yields an oily lactone (310°-320°).

Ethyl other EtA'. (260°). Oil. PHENYL-HEXINENE v. BUTENYL-STYRENE.

DI-PHENYL-HEXOIC ACID. Nitrile. CH₂Ph.CPhPr.CN. [63°]. (830°-840°). Formed from C_sH_s.CHPr.CN, benzyl chloride, and NaOH (Rossolymo, B. 22, 1236). Needles.

TETRA-PHENYL-HEXUNENE

CPh OH.OH CPh. [269°]. Formed by heat-

ing di-benzyl-carbinol with MeI at 265° (Bogdanowska, B. 25, 1273). Needles, insol. alcohol and ether, sol. CHCl.

PHENYL-HEXYL-TRIAZOLE CARBOXYLIC ACID CH₃(CH₂)₅C $\stackrel{\text{N.NPh}}{<_{\text{N:C.CO}_2\text{H}}}$. $[126^{\circ}].$ tained by saponifying its nitrile, which is got from phenyl-hydrazine dicyanide and heptoic aldehyde (œnanthol) (Bladin, B. 25, 186).

very plates, v. sol. alcohol. -CuA'2 aq. -HA'HCl. Amide. [82.5°]. White plates. Thio-amide C₁₄H₁₈N₃.GS.NH₂. Formed from the crude nitrile and alcoholic

ammonium sulphide. Yellow plates, sol. alcohol. DI-PHENYL-HEXYL-TRICYANIDE

(265° at [44°]. 15 mm.). Got from heptoyl chloride, benzonitrile, and AlCl, at 70° (Krafft a. Hanssen, B. 22, 808).

PHÉNYL HEXYL KETONE C,H,CO.C,H, (267°) at 740 mm. Formed by the action of AlCl₃ on a mixture of benzene and heptoyl chloride (Auger, Bl. [2] 47, 50; Krafft, B. 19, 2987). Leaflets. Yields an oxim [55°].

PHENYL-HEXYL-(\beta)-NAPHTHOTRIAZINE. $\label{eq:distance} \textit{Dihydride} \ \ C_{2.l}H_{25}N_s \ \textit{i.e.} \ \ C_{10}H_e < \stackrel{N.CH.C_6H_{13}}{\dot{N}.\dot{N}Ph}.$ [176.5°]. Formed by heating benzene-azo- (β) naphthylamine with cenanthol and alcohol (Goldschmidt a. Poltzer, B. 24, 1007). Needles, sol. alcohol.—B'HCl. [226°]. White needles.

-B'2H2PtCl6. [225°]. Granules. PHENYL-HEXYL-PYRAZOLE C15H20N2 i.e. C.H.3.C & CH-CH.N (319°). Formed from

heptoyl-acetic aldehyde and phenyl-hydrazine

(Claisen a. Stylos, B. 21, 1149). Oil.

PHENYL-HEXYL-THIO-UREA C13H20N2Si.e. [53°]. CS(NHPh).NH.CH₂.CHEt₂. Formed from hexylamine and phenyl-thiocarbimide (Freund a. Herrmann, B. 23, 195). Prisms.

PHENYL-HEXYL-UREA C₁₈H₂₀N₂O i.e. (NHPh).NH.CH₂.CHEt₂. [70°]. For CO(NHPh).NH.CH, CHEt, Formed from hexylamine and phenyl cyanate (Freund a. Herrmann, B. 23, 194). Needles.

PHENYL-HOMO-ITAMALIC ACID v. Oxy-

BENZYL PYROTARTARIC ACID

PHENYL-HYDANTOIC ACID

NH2.CO.NH.CHPh.CO2H. [178°]. Got by boiling (a)-phenyl-hydantoin with baryta (Pinner, R. 21, 2326). Prisms, sl. sol. cold water.

Amide NH2.CO.NH.CHPh.CO.NH, [223°]. Formed by adding the compound got from urea and the cyanhydrin of benzoic aldehyde to H2SO4 and pouring into water at 0° (Pinner a. Spilker, B. 22, 697). Prisms (from dilute alcohol). Yields phenyl-hydantoin on heating.

PHÊNYL-HYDANTOÏN C,H,N2O2 i.e.

co<NH .CO. [192°]. Formed by fusing urea with phenyl-amido-acetic acid (Schwebel, B. 10, 2045). Needles, v. sl. sol. water, sol. alkalis.

(a)-Phenyl-hydantoin $CO < NH.CO \\ NH.CHPh$. oxy-phenyl-glyoxaline. [178°]. Formed by heating the cyanhydrin of benzoic aldehyde with mandelic nitrile at 100° (Pinner, B. 21, 2321). Formed also from phenyl-uramido-acetic ether and KOH (Kossel, B. 24, 4150). Needles (from HClAq). KOH and MeI yield

C₂H₇MeN₂O₂ [162°], crystallising in needles.— KA': needles, v. sol. water.

Acetyl derivative C.H.AcN.O. [145°]. Small needles (from benzene), insol. water.

 $C(NH) < NH.CO \\ CHPh.$ Phenyl- ψ -hydantoïn [above 300°]. Got by dissolving (a)-phenylhydantoin in alcoholic potash and then adding HCl. Crystalline.—KA': almost insol. water.

Acetyl derivative C(NH) < NAc.CO O-CHPh. White prisms (Pinner a. Spilker, B. 22, 698). Decomposes at about 290°.

PHENYL-HYDRAZIDO-ACETIC ACID

C_aH₃NH.NH.CH₂.CO₂H. $[159^{\circ}].$ Formed by reducing the phenyl-hydrazide of glyoxylic acid (Elbers, A. 227, 353). Formed also by reducing phenyl-osotriazole carboxylic acid with sodiumamalgam (Pechmann, A. 262, 288). Hexagonal tables, sol. hot water. Re-oxidised by Fehling's solution to NHPh.N:CH.CO.H.

u-Phenyl-hydrazido-acetic acid NH₂.NPh.CH₂.CO₃H. [131°]. Got by saponifying its ether [127°], which is formed by heating phenyl-hydrazine with chloro-acetic ether at 100° (Reissert a. Kayser, B. 24, 1519). Needles.

PHENYL-HYDRAZIDO-BUTYRIC ACID C₁₀H₁₄N₂O₂ i.e. CHEt(N₂H₂Ph).CO₂H. Formed by reducing the phenyl-hydrazide of ethyl-glyoxylic acid (Japp a. Klingemann, A. 247, 217). Needles (from MeOH). Softens at 165%.

Phenyl-hydrazido-isobutyric acid

 $CMe_2(N_2H_2Ph).CO_2H.$

Anhydride N₂HPh CO [175°]. Got by gently heating the nitrile with H2SO, and pouring into water (Reissert, B. 17, 1458). Crystals, v. sol. alcohol.—B'HCl: crystals.

Nitrile CMe₂(N₂H₂Ph).CN. [70°]. Got by heating the cyanhydrin of acetone with phenyl-hydrazine at 100°. Needles, v. sol. alcohol.

Imide (CMe₂(N₂H₂Ph).CO)₂NH. [117°]. Got by passing HCl into an alcoholic solution

of the nitrile.—B"H,Cl,: crystals.

Phonyl-hydrazido-butyric acid $NH_2.NPh.CHMe.CH_2.CO_2H.$ [111°]. Formed from potassium β-bromo-butyrate, phenylhydrazine and NaOAc in aqueous solution (Lederer, J. pr. [2] 45, 87). Plates (from alcohol), v. sol. hot Aq. Converted by conc. H2SO4 into oxy-phenyl-methyl-pyrazole dihydride.

PHENYL - β - HYDRAZIDO - CROTONIC ETHER CH₂.C(NH.NHPh):CH.CO₂Et. Formed from acetoacetic ether (1 mol.) dissolved in ether by adding phenyl-hydrazine (Nef, A. 266, 70; cf. Knorr, A. 238, 147). Colourless needles, v. e. sol. ether, sl. sol. ligroin. Turns yellow in air. Split up by conc. HClAq into phenyl-When heated hydrazine and acetoacetic ether. in vacuo at 200° it changes to oxy-phenyl-methyl-pyrazole. Conc. H₂SO₄ followed, after 10 minutes, by water forms methyl-indole carboxylic ether [134°]. HgO oxidises it to the azocompound CH, C(N,Ph):CH.CO,Et crystallising from light petroleum in red needler, [51°]. LcCl forms a di-acetyl derivative CH₃.C(NAc.NAcPh):CH.CO₂Et (245°-250°

150 mm.). PHENYL-HYDRAZIDO - METHYL-QUINOL-INE C. H. MeN(N2H2Ph). [197°]. Formed from (Py. 3) methyl-quinoline and phenyl-hydrazine (Ephraim, B. 24, 2820).

PHENYL - HYDRAZIDO - METHYL - THI-AZOLE. $Dihydride C_{10}H_{13}N_3S$ i.e.

CHMe.S C.NH.NHPh. Phenyl-propylene-ψ-thio-semicarbazide. [93°]. Formed by heating phenyl-allyl-thio-urea with conc. HClAq at 100° (Arenarius, B. 24, 269). Yellowish plates (from ligroin).—B'HCl. [202°]. Concentric crystals.—Pierate. [167°].

PHENYL-HYDRAZIDO-PHENYL-ACETIC ACID C₁₄H₁₄N₂O₂ i.e. NHPh.NH.CHPh.CO₂H. [158°]. Got by reducing the phenyl-hydrazide of phenyl-glyoxylic acid (Elbers, A. 227, 345). Crystalline, v. sl. sol. cold water, v. sol. alcohol.

PHENYL-HYDRAZIDO-PROPIONIC ACID C₉H₁₂N₂O₂ i.e. NH₂NPh.CHMe.CO₂H. [187°]. Formed by boiling its amide with NaOHAq (Reissert, B. 17, 1454; 20, 3110; 22, 2924). The ether is got by heating lactic ether with phenyl-hydrazine. Needles. Yields phenyl-a-amido-propionic acid on reduction.

Ethyl ether EtA'. [116°].—EtA'HCl.
Nitrile. [58°]. Formed by heating
CH₃.CH(OH).CN with phenyl-hydrazine at 100°.
Amide. [124°]. White crystals.

Phenyl-hydrazido-propionic acid NHPh.NH.CHMe.CO₂H. [172°]. Formed by reducing the phenyl-hydrazide of pyruvic acid with sodium-amalgam (Fischer a. Jourdan, B. 16, 2244; Japp a. Klingemann, B. 20, 3284; A. 247, 211). Needles (from MeOH). Re-oxidised by cold Fehling's solution to NHPh.N:CMe.CO₂H.

PHENYL-HYDRAZIDO-QUINOLINE C.H.N.NH.NHPh. [191°]. Formed from (Py. 3)-chloro-quinoline and phenyl-hydrazine (Ephraim, B. 24, 2818). Needles, v. sol. chloroform. Reduced by HI and P to (Py. 3)-amido-quinoline.

PHENYL-HYDRAZINE $C_6H_8N_2$ i.c. NHPh.NH₂. [23°]. (241·5° i.V.) at 750 mm. S.G. $\frac{22\cdot7}{4}$ 1·097 (E. Fischer, A. 236, 198). H.C.v. 805,240. H.C.p. 806,300. H.F. 34,200 (from diamond) (Petit, A. Ch. [6] 17, 168).

Formation.—(E. Fischer, B. 8, 589, 1005, 1641; 9, 111, 880; A. 190, 67).—By the action of zine and HOAc on diazo-benzene anilide or diazo-benzene diethylamide.

Preparation.—1. A cold solution of 71 pts. of sodium nitrite in 50 pts. of water is slowly added to a well-cooled mixture of 10 pts. of aniline and 200 pts. of HCl (30 p.c.). diazo-benzene chloride solution so prepared a cold solution of 45 pts. of stannous chloride in 45 pts. of HCl is now added, when the mixture immediately becomes a thick magma of phenylhydrazine hydrochloride, which is filtered off, basified, and distilled (Meyer a. Lecco, B. 16, 2976).—2. Aniline (28 g.) is dissolved in K2CO3 (21 g.) and water (500 c.c.) by passing in SO₂. The solution is mixed with a cold solution of NaNO₂ (25 g.) exactly neutralised by HOAc. After two hours the product is warmed till dissolved, acidified by HOAc, and reduced with zinc-dust and HClAq (Reychler, B. 20, 2463). By adding diazo-benzene nitrate to a cold aqueous solution of KHSO, heating the resulting C_eH₅.N₂.SO₃K with excess of KHSO₃, and decomposing the C₆H₅.NH.NH.SO₃K, thus formed, by heating with HClAq.

Properties.—Oil, solidifying to monoclinic tables; v. sl. sol. hot water, almost insol. KOHAq. Miscible with alcohol, ether, benzene, and CHCl. Volatile with steam. Reduces Fehling's solution in the cold, N, aniline, and benzene being formed. When heated for a long time at 150° it is gradually split up into aniline, NH₃, benzene, and N (Reissert a. Kayser, B. 23, 3703). Reduces SeO₂ to Se (Hinsberg, A. 260, 42). Gives benzene and N when boiled with CuSO₄. It is very poisonous.

Reactions.—1. H₂O₂ forms benzene and diazo-

benzene imide PhN, (Wurster, B. 20, 2633).— 2. Yellow HgO added to its ethereal solution forms aniline, N, and HgPh₂ (Fischer, A. 199, 320). HgO added to a solution of the sulphate forms diazo-benzene sulphate and imide.- SOCl₂ in ether forms in the cold NPhH.N:SO, which crystallises in yellow prisms [105°]; v. sol. ether, volatile with steam. Thionyl-phenyl-hydrazine is hardly attacked by HCl, but is converted by NaOHAq into phenyl-hydrazine and Na₂SO₃. Br forms NPhBr.NBr₂. AcCl forms diazo-benzene chloride, S, and HOAc (Michaelis, B. 22, 2228; A. 270, 115).—4. SO₂ passed into an ethereal solution of phenyl-hydrazine forms NHPh.NH₂SO₂ as a crystalline pp. SO₂ passed into an alcoholic solution of phenyl hydrazine forms (NHPh.NH₂),SO,; v. sol. water and alcohol, insol. ether, crystallising in tables which shrink together at 70°, but decompose at a higher temperature (Michaelis a. Ruhl, B. 23, 474). SO₂ passed into a benzene solution at 75° forms NHPh.N:SO, which when heated with phenylhydrazine gives Ph₂S₂, benzene, and N.--5. Dry CO2 forms solid phenyl-hydrazine phenyl-carbazate (NHPh.NH.), CO. or N, PhH., CO.O. N, PhH,, which is crystalline; sl. sol. water and ether, decomposed by hot water .- 6. CS, forms the corresponding N₂PhH₂.CS.S.N₂PhH₄ [97°], which at 110° splits up into H₂S and di-phenyl-thiocarbazide. COS forms crystalline COS(N₂H₃Ph)₂ (Fischer, B. 22, 1935).—7. Zinc-dust and conc. HClAq slowly form aniline and NH_s.—8. Excess of phenyl-hydrazine acts upon iodine, forming HI, aniline, and PhN, (Fischer, B.10, 1335). Excess of I forms iodo-benzene, N, and HI (E. von Meyer, J. pr. [2] 36, 115). This reaction may be used for the volumetric estimation of phenylhydrazine.—9. Sulphur above 80° forms aniline, H₂S, N, Ph₂S₂, Ph₂S, PhSH, benzene, and NH₃. 10. Nitrous acid forms PhN₃. KNO₂ added to the hydrochloride at 0° forms a yellow crystalline pp. of NPh(NO)NH, -11. Alcoholic potash and chloroform form, on heating, C14H12N4 [180°], mol. w. 236. This body crystallises in white needles; sol. alcohol, scarcely affected by HClAq at 200°, and yielding crystalline C₁₄H₁₁N₄.NO₂ and C₁₄H₁₁N₄.SO₃H (Ruhemann a. Elliott, *C. J.* 53, 850).—12. EtBr forms NPhEt.NH₂, NPhH.NHEt, and NPhEt, Br. NH, -13. Aqueous diazobenzene nitrate added to a cold solution of the hydrochloride ppts. PhN₃, aniline nitrate remaining in solution.—14. Reacts with aldehydes and ketones, forming phenyl-hydrazides; e.g. (Fischer, B. 17, 572) CH₃·CHO+NPhH.NH₂-H₂O+CH₃·CH:N.NHPh. Thus a solution of phenyl-hydrazine hydrochloride (2 pts.) and crystallised sodium acetate (3 pts.) in water (20 pts.) gives a pp. of the phenyl-hydrazide when added to an aqueous solution of an aldehyde or ketone, either in the cold, or, in the case

of aromatic compounds, on warming. These phenyl-hydrazides are decomposed by acids into the original components, and, in alcoholic solution, by sodium-amalgam and HOAc into aniline and a base. The phenyl-hydrazides of ketones are converted by ZnCl₂ into derivatives of indole. The phenyl-hydrazides of aldehydes and ketones are described under the aldehydes and ketones from which they are derived .- 15. Diketones containing the group CO.CHX.CO yield derivatives of phenyl-pyrazole. Saturated y-diketones in ethereal solution usually form a mono- or di- phenyl-hydrazide; in HOAc the product is the anhydride of the mono-phenyl-hydrazide, a derivative either of CH. CH:N or of CH:CH N.NHPh (Klingemann, A. 269, 108) .- 16. Acetoacetic ether forms a phenyl-hydrazide, which at 140° changes to oxyphenyl-methyl-pyrazole NPh $<_{
m N}^{
m CO.CH_2}$ by loss of alcohol. Acetyl-propionic acid forms the com- $CH_3.\tilde{C}(\bar{N_2}H\bar{P}h).CH_2.CH_2.CO_2H$ pound Benzoylacetic ether forms two compounds [132°] and [165°] (Bender, B. 21, 2495).—17. Reacts with sugars as with aldehydes; thus a solution of glucose gives CH₂(OH).(CH(OH))₄.CH:N₂HPh [145°] on heating with phenyl-hydrazine, while galactose gives C₈H₁₂O₅(N₂HPh) [158°] (Fischer, B. 20, 824). On heating these compounds with aqueous phenyl-hydrazine acetate at 100° 'osazones' are formed, e.g. phenyl-glucosazone CH₂OH.(CH.OH)₃.C(N₂HPh).CH:N₂HPh [204°] and the isomeric phenyl-galactosazone [c. 191°]. 18. By heating with phenyl-hydrazine hydro-chloride and NaOAc arabinose is converted into $C_{17}H_{20}N_4O_3$ [158°]; formose into $C_{18}H_{22}N_4O_4$ [c. 144°]; acrose (the product of the action of NaOHAq on glyceric aldehyde) into C₁₈H₂₂N₄O [c. 217°] and an isomeride [159°].—19. Phenyl hydrazine acetate reacts with the product of oxidation of alycerin with Na₂CO₃Aq and Br forming glycerosazone C13H16N4O crystallising from benzene in yellow plates [131°] (Fischer a. Tafel, B. 20, 1089, 3386).—20. Erythrite, after oxidation by dilute HNO_s, yields with phenylhydrazine hydrochloride and NaOAc the compound C₄H₆O₂(N₂HPh)₂ [167°].—21. Mannite on oxidation with HNO, yields levulose (mannitose) and mannose, which with phenyl-hydrazine forms $C_{12}H_{18}N_2O_b$ [195°-200°] (Fischer, B. 21, 1805).—22. The acetate reacts with the product of oxidation of dulcite by bromine and Na₂CO₃Aq with formation of the 'phenyldulciteosazone' C₁₈H₂₂N,O, [206°].—23. Isodulcite heated with phenyl-hydrazine hydrochloride and sodium acetate at 100° forms 'phenyl-isodulcite-osazone' C₁₈H₁₂N₄O₃ or C₆H₁₀O₄(N₂HPh)₂[180°]. Solutions of isodulcite and phenyl-hydrazine form, on mixing, C₆H₁₂O₄(N₂HPh)[159°] (Fischer a. Tafel, B. 20, 2574; Raymann, Bl. [2] 47, 760).—24. Sorbin heated with aqueous phenyl-hydrazine hydrochloride and NaOAc at 100° forms the osazone C₁₈H₂₂N₄O₄ [164°].—25. Milk sugar heated with aqueous phenyl-hydrazine hydrochloride and NaOAc forms C₂₁H₂₂N₄O₆ [200°] and C₂₄H₃₂N₄O₈ [224°].—26. Maltose forms C₂₄H₃₂N₄O₆ [191°].—27. The acetate gives with glyoxylic, phenyl-glyoxylic, and pyruvic acids the corresponding phenyl-hydrazides [137°], [158°], and [192°] respectively.—28. Phenyl-

hydrazine in HOAc (50 p.c.) gives with a dilute solution of gluconic, galactonic, and arabinose carboxylic acids compounds C₈H₁₁O₈.N₂H₂Ph [200°], [203°], and [216°] respectively (Fischer a. Passmore, B. 22, 2730). These compounds are m. sol. hot water. Saccharic acid gives a similar compound C₆H₁₁O₅N₂H₂Ph [165] which, as well as the compounds from mucic and metasaccharic acid, is almost insoluble. The compound from mucic acid melts at 240°. All these compounds contain the group CO.NH.NHPh. Compounds containing this group are also obtained by the action of phenyl-hydrazine and dilute HOAc at 100° on formic, succinic, malic, tartaric, cinnamic, and gallic acids, but not from glycollic, lactic, and glyceric acids. The compounds got from succinic, malic, gallic, and tartaric acid melt at 218°, 223°, 187°, and 240° respectively. Acetyl-phenyl-hydrazine is produced when aqueous phenyl-hydrazine acetate is heated at 130°; and a similar reaction occurs with benzoic, glyceric, and lactic acids. Compounds containing the group CO.NH.NHPh give a reddish-violet colouration with H₂SO₄ and a little FeCl₃, and they are readily decomposed by KOHAq and by baryta into phenyl-hydrazine and a salt of the acid (Fischer, B. 22, 2728).-29. Glucose carboxylic lactone yields C,H,O,N,H,Ph [172°], while mannose carboxylic acid gives an isomeric body [223°].- 30. The acetate gives with rhamnose carboxylic acid the compound C₇H₁₃O₇N₂H₂Ph [c. 210°], while 31. Malonic acid heated at 100° with phenylhydrazine and dilute HOAc forms C₁₃H₁₈N₄O₃, [143°] and CO₂H.CH₂.CO.N₂H₂Ph [154°], which when heated with water and phenyl-hydrazine at 200° yields $CH_2 < {CO \atop CO} > N_2HPh$ [128°]. — 32. no-propionic ether. a-Oxy-butyric acid at 160° forms $C_{10}H_{11}N_{1}O_{2}$ [152°], while mandelic acid yields $C_{11}H_{11}N_{1}O_{2}$ [182°] (Reissert a. Kayser, B. 22, 2924). — 33. γ -Oxy-valeric lactone at 100° forms C.H.O.N. 1177 [79°] (Wishiony) [79°] (Wislicenus, B. 20, 402).—34. Benzoyl-carbinol gives N₂HPh:CPh.CH₂OH [112°] and the 'osazone N₂HPh:CPh.CH:N₂HPh [192°] A. 243,(Laubmann, 35. Boiling aqueous itaconic acid forms the acid NHPh.N CH. CH.CO.H [194°] (Scharfenberger, A. 254, 150).—36. Cinnamic acid forms C₁₃H₁₄N₂O [183°], which on distillation yields C₁₃H₁₂N₂O [251°] (Knorr, B. 20, 1107).—37. Acrolein reacts forming phenyl-pyrazole dihydride NPh<_CH_2.CH_ [52°] (274°) (Fischer a. Knoevenagel, A. 239, 194).-38. Mesityl oxide yields $NPh < \begin{array}{l} N = CMe \\ CMe_2.CH_2 \end{array}$ phenyl-tri-methyl-pyrazole (F. a. K.) .- 39. Chloro-acetone in alcoholic solution at -18° forms $C_{24}H_{26}N_6$ [158°] which yields an acetyl derivative $C_{24}H_{25}AoN_6$ [220°] (Bender, s-Tetra-chloro-acetone forms B. 21, 2496). C_{1.}H_{1.2}N₄ (126°) converted by fuming HNO₄ into C_{1.}H_{1.9}N₄O₃ [235°) and by SnCl₂ to two usses C_{1.8}H_{1.4}N₄ (77°) and [193°] (Levy a. Witte, A. 252, 343; Zincke a. Kegel, B. 22, 1478).—40. Tetrabromo-di-benzyl-ketone heated with alcoholic phenyl-hydrazine at 150° forms C,Ph2(N2Ph), [70°] (Bourcart, B. 22, 1369).-41. Di-bromo-acetophenone forms N.HPh:CPh.CH:N2HPh [148°], crystallising from alcohol in yellow prisms (B.). 42. Bromo-acetyl-propionic ether mixed with an alcoholic solution of phenyl-hydrazine forms CH₃.C(N₂HPh).CH:CH.CO₂Et [117·5°], which on reduction with tin and HCl forms methyl-indole carboxylic acid (Bender). Bromo-acetyl-propionic acid forms CH3.C(N2HPh).CH:CH.CO2H [157°] (Decker, B. 21, 2937). — 43. Di-chloromaleic imide at 100° forms C₄O₂(N₂H₂Ph)₂NH crystallising from acetone in orange-red needles (Ciamician, B. 22, 2495).—44. Amides react thus: N₂H₃Ph + R.CO.NH₂= NH₈ + R.CO.N₂H₂Ph (Just, B. 19, 1201).—45. Carbanic ether forms, on heating, di-phenyl-carbazide CO(NH.NHPh)₂ [151°] (Skinner a. Ruhemann, B. 20, 3372).— 46. Reduces alloxan to alloxantin, N and benzene being formed (Pellizzari, G. 17, 254). In this reaction the first product is the compound CO NH.CO C(OH).N₂H₂Ph (Skinner a. Ruhemann, C. J. 53, 550).—47. Parabanic acid with phenyl-hydrazine hydrochloride and NaOAc yields C₁₅H₂₀N₄O₄ [170°], which on boiling with water forms NH₂.CO.NH.CO.CO.N₂H₂Ph [215°] (S. a. K.).—48. *Urea* heated with phenylhydrazine hydrochloride at 160° forms phenylurazole NPh $<_{
m CO\ NH}^{
m NH.CO}$ [263°]. — 49. Reduces nitroso-aniline to p-phenylene-diamine and $C_{12}H_{12}N_1O$ (Fischer, B. 21, 2610). -50. The acctate (2 mols.) acting on nitroso-dimethylaniline (3 mols.) forms C11H16N4O [103°]. An ethereal solution of phenyl-hydrazine converts nitroso-di-methyl-aniline into di-methyl-amidobenzene-azo-di-methyl-aniline and phenylene-di-methyl-diamine. In alcoholic solution there is formed NHPh.C₆H₄.NMe₂ [130°] which yields a nitrosamine $C_{14}H_{15}N_{3}O$ [116°] (O. Fischer, B. 21, 2609).-51. Di-nitroso-acctone (1 mol.) mixed with phenyl-hydrazine (1 mol.) in hot alcoholic solution forms CH(NOH).C(N₂HPh).CH(NOH) [145°]. Di-nitroso-acetone boiled with excess of phenyl-hydrazine acetate forms C1,H12N4 crystallising in yellow plates [122°].-52. Diazoacetic ether at 100° gives off N and forms a colourless compound [136°]. — 53. p-Diazobenzene sulphonic acid forms the compound N3C8H4.SO3N2H4Phaq crystallising in plates (Griess, B. 20, 1528). - 54. Di-acetyl-succinic ether fused with phenyl-hydrazine at 100° forms CO₂Et.CH(CMe:N₂PhH).CHAc.CO₂Et [91°] and CO₂Et.CH(CMe:N₂PhH).CH(CMe:N₂PhH).CO₂Et which on heating with toluene at 180° gives dioxy-di-phenyl-di-methyl-dipyrazyl (q.v.). — 55. Dioxims form additive compounds; e.g. glyoxim yields C.H. (NOH) 2N. H. Ph [110°], di-phenylglyoxim yields C₂Ph₂(NOH)₂N₂H₃Ph [150°], while the di-oxim of (β)-naphthoquinone forms C₁₀H₆(NOH)₂N₂H₃Ph [138°] (Polonowsky, B. 21, 182).-56. Phloroglucin kept for five days with an alcoholic solution of phenyl-hydrazine in the cold forms $C_0H_8(OH)(N_2H_2Ph)_2[144^\circ]$ which may be oxidised by FeCl₃ to $C_0H_3(OH)(N_2Ph)_2[177^\circ]$ (Baeyer B. 22, 2891).—57. Thio-acetoacetic ether forms S(CH<CMe:N), [183°], together with the compounds NHPh.N:C<CO-NPh $\theta_2(CH < \frac{CO - NPh}{CMe: N})_2$ and di-oxy-di-phenyl-dimethyl-dipyrazyl (Sprague, C. J. 59, 332).-58. Cyanogen passed into an emulsion of phenylhydrazine in cold water forms the dicyanide (v. infra). -59. Cyanogen passed into an alcoholic solution forms PhN₂H₂.C(NH).C(NH).N₂PhH₃ [225°], crystallising in white plates (from alcohol). The same compound is formed by the action of phenyl-hydrazine on cyananiline. gives a rose-red colour with alcoholic NH3. Conc. H₂SO₄ is coloured indigo-blue. Dry HCl forms the hydrochloride C14H18N62HCl [188]. Dilute HCl forms PhN.NH2.CO.C(NH).NHNHPh [180°].—60. Cyananiline in chloroform solution forms, after boiling, matted needles of the compound $C_{21}H_{21}N_{1}Cl_{2}$ [200°-212°]. This body is perhaps (PhH₂N₂·CHCl.C₁,H₁,N₈)2HCl (Senf, J. pr. [2] 35, 533).—61. PCl₈ in ether forms amorphous NHPh.N:P.NH.NHPh converted by water into (NHPh.NH) POH, a yellowish-white powder [92°] which reduces Fehling's solution in the cold (Michaelis a. Oster, A. 270, 126). This body is split up by acids into phenylhydrazine and H₃PC₃, but is attacked by alkalis with difficulty. Benzoic aldehyde produces NPhH.N:CHPh and H₃PO₃.—62. POCl, forms PO(NH.NHPh), [196°], easily decomposed by acids and alkalis (Michaelis, A. 270, 135).— 63. PSCl, forms, in like manner, PS(NH.NHPh), [154°].—64. AsCl, forms(N2PhH3), AsCl, an amorphous powder decomposed at 236°.—65. BCl₃ forms, in like manner, (N₂l'hH₃)₃BCl₃, v. sol. water, decomposed at 242°.—66. C₆H₅PCl₂ in ether forms NHPh.N.PC, H, [152°] crystallising from EtOAc in plates (Michaelis a. Oster, A. 270, 129). It reduces hot Fehling's solution, and is split up by acids into PhPO2H2 and phenyl-hydrazine.

Salts.—B'HCl. Plates, m. sol. water. Got by ppg. an alcoholic solution of the base by HCl, and washing the pp. till it is colourless.—B'HBr. Needles (from water).—B'₂H₂SO₂. Plates, v. sol. water.—B'C₂H₃N₃O₂. Yellow needles.—B'₂H₂C₂O₄. Plates.—Benzene sulphinate; [131°] (Escales, B. 18, 895).—Dinitro-(a)-naphthol sulphonate: lightbrown needles, sl. sol. hot water (Richardson, C. N. 58, 39).—Sulphocyanide: sparingly soluble plates. Changes at 160°-170° into phenyl-thio-semicarbazide.—B'H₃PO₃. [118°] (Michaelis, A. 270, 127, 133).—B'₂H₃PO₃. [121°].—B'C₆H₅PH₂O₂. [135°].

Sodium derivative C.H.NNa.NH. Formed by dissolving 8g. of sodium in 70g. of warm phenyl-hydrazine. The H evolved reduces some of the phenyl-hydrazine to NH, and aniline; when the reaction is complete the aniline and excess of phenyl-hydrazine are distilled off by heating the product in an oil-bath to 200°-210° (not higher) in a current of H. It is a yellow hygroscopic powder. When finely divided it sometimes takes fire in the air. Decomposed by water into phenyl-hydrazine and NaOH. Readily reacts with alkyl haloids, giving unsymmetrical phenyl-alkyl-hydrazines (Michaelis, B. 19, 2448; A. 252, 266).

Formyl derivative NHPh.NH.CHO. [145°]. Formed by heating formamide with phenyl-hydrazine at 180°. White scales.

Acetyl derivative NHPh.NHAc. [128.53]. Formed by heating phenyl-hydrazine with HOAc or acetamide (Fischer, A. 190, 129; Just. B.

19, 1201). Formed also from the sodium-derivative and Ac.O at 0° (Michaelis a. Schmidt, A. 252, 301). Six-sided prisms, sl. sol. cold water.

derivative NPhAc.NHAc. Di-acetyl [108°]. Formed from NPhNa.NH₂ and AcCl in ether. Tables or needles, v. e. sol. alcohol. Completely reduces hot Fehling's solution.

Chloro-acetyl derivative NHPh.NH.CO.CH₂Cl. [115°]. Formed from the base and CH, Cl. COCl in ether (Gattermann, B. 25, 1081).

Propionyl derivative NHPh.NH.CO.Et. [158°] (Freund a. Goldschmidt, B. 21, 2461).

n-Buturul derivative. [114]. Scales, v. sol. ether (Michaelis a. Schmidt, A. 252, 308).

Isovaleryl derivative NHPh.NH.CO.CH,Pr. [101°]. Formed from the base and aceto-valeric anhydride (Autenrieth, B. 20, 3190). Plates, v. sl. sol. cold ligroïn.

Hexoyl derivative. [117°]. Needles.

s-Benzoyl derivative NHPh.NHBz.

s-Benzoyl derivative [169°]. Formed from phenyl-hydrazine by adding BzCl in ether at 0° (Fischer, A. 190, 125), or by heating with benzamide (Just, B. 19, 1203). Formed also by warming phenyl-hydrazine with BzCy (Hausknecht, B. 22, 329). Prisms, sl. sol. hot water. Yields NPhMe.NHBz on methylation. Reduces HgO. Phthalic anhydride at 180° forms an addition-compound $C_{21}H_{16}N_2O_4$ [172°], crystallising from alcohol in prisms. It is probably $CO_2H.C_6H_4.CO.NPh.NHBz$. At 250° it splits up into benzoic acid and (B)-phthalyl-

 $phenyl-hydrazine \ C_{e}H_{4};C_{2}O_{2} \diagdown | \\ NPh$

NPhBz.NH2. derivative u-Benzoyl [70°]. A product of the action of BzCl in benzene on NPhNa.NH2 at 0° (Michaelis a. Schmidt, B. 20, 43, 1718; A. 252, 310). Needles (from water). Reduces hot Fehling's solution but not HgO. Conc. HCl at 150° gives HOBz and phenyl-hydrazine. HNO₂ forms in acid solution benzanilide and N₂O, but in neutral solution it forms N₃Ph and HOBz. Reacts like phenylhydrazine with aldehydes and ketones; thus acetone yields NPhBz.N:CMe₂ [115.5°], acetophenone forms NPhBz.N:CMePh [124°], and benzoic aldehyde gives NPhBz.N:CHPh [122°]. Salts.—B'HCl. [202°]. Needles, m. sol. HClAq. water.—B'₂H₂SO₄. [191°]. Needles, sl. sol. cold water.—B'₂H₂SO₄. [191°]. Needles, sl. sol. cold water.—B'HNO₃. [145°].—B'C₆H₄N₃O₇. [122°].

Di-benzoyl derivative NPhBz.NHBz.

[178°] (Fischer, A. 190, 128; M. a. S.). Prisms, v. sl. sol. water, sol. alkalis. On methylation it yields NPhBz.NHBz.—NPhBz.NBzNa: plates.

Benzoyl - acetyl derivative NPhBz.NHAc. [153°]. Got from NPhBz.NH2 and Ac₂O. Needles (from dilute alcohol).

o-Nitro-benzoyl derivative NHPh.NH.CO.C_eH₄,NO₂. [198°]. Formed from phenyl - hydrazine and C_eH₄(NO₂)CN (Hausknecht, B. 22, 328). Yellowish-red needles.

o-Amido-bensoyl derivative. [170°]. Got by warming phenyl-hydrazine with an

heating phenyl hydrazine (1 mol.) with m-amidobenzoic acid (1 mol.) at 165° (Pellizzari, G. 16, 200). Scales, insol. cold alkalis, sol. acids.

Amido-toluyl derivative [8:6:1] C_eH₃Me(NH₂).CO.N₂H₂Ph. [198°]. Formed from methyl-isatoic acid and phenyl-hydrazine (Panactivitch J. pr. [2] 33, 68). Prisms (from chloroform), sl. sol. ether.

Phenyl-acetyl derivative $N_2H_2Ph.CO.CH_2Ph.$ [169°]. Got by heating phenyl-acetic acid with phenyl-hydrazine at 130° (Bülow, A. 236, 196). Needles (from alcohol), sl. sol. water.

Cinnamoyl derivative N₂H₂Ph.CO.C₂H₂Ph. [258°]. Got by heating cinnamic acid with phenyl-hydrazine at 190° (Knorr, B. 20, 1108). Forms, on distillation, a compound $C_{15}H_{12}N_2O$ [258°], crystallising in needles with green fluorescence.

Oxalyl derivative C₂O₂(N₂H₂Ph)₂. [278°]. Described in vol. iii. p. 655.

Ethyl-malonyl derivative CHEt(CO.N.H.Ph)₂. [233°]. Got by heating ethyl-malonic amide with phenyl-hydrazine at 220° (Freund a. Goldsmith, B.21, 1242). Needles (from HOAc). Converted by COCl, into C,9H,6N,O, [113°].

Succinyl derivative $C_2H_4:C_2O_2:(N_2H_2Ph)_2.$ [218°]. Formed from succinyl chloride (1 mol.) and phenyl-hydrazine (2 mols.). Formed also by heating C2H4:C2O2:S with phenyl-hydrazine and HOAc (Zanetti, Rend. Accad. Linc. [4] 5, i. 225). Plates (from HOAc) (Freund, B. 21, 2456; Fischer a. Passmore, B. 22, 2734). When succinic anhydride is heated with phenyl-hydrazine at 160° there is formed C_2H_4 : \tilde{C}_2O_2 :N.NHPh [155°], which yields a nitrosamine C₂H₄:C₂O₂:N₂Ph.NO [84°] (Hötte, J. pr. [2] 35, 293).

(a) - $\dot{P}hthalyl$ derivative C_0H_4 : C_2O_2 : N.NHPh. [179°]. Formed from phenylhydrazine and phthalyl chloride in ether (Pickel, A. 232, 233). Formed also by heating phenylhydrazine (1 mol.) with phthalimide (1 mol.) at 120° (Pellizzari, G. 16, 203; Just, B. 19, 1204); or with phthalic anhydride (1 mol.) at 150° (Hötte, J. pr. [2] 33, 99; 35, 268). Prepared by boiling an alcoholic solution of phthalic anhydride with phenyl-hydrazine. Colourless needles (from alcohol), insol. water and alkalis. Conc. H.SO₄ forms a violet solution, turning brown. Reactions.—1. Converted by boiling alcoholic potash into N₂H₂Ph.CO.C₆H₄.CO₂H [166°], an acid of which the benzoyl derivative NHPh.NBz.CO.C.H..CO.H [172°] is formed by heating NHPh.NHBz with phthalic anhydride at 180°.—2. Boiling aqueous KOH or HClAq at 150° splits it up into phthalic anhydride and phenylhydrazine.—3. Benzoyl chloride at 160° forms C.H.:C2O2:N.NPhBz [193°].-4. Nitrous acid passed into its solution in diluted HOAc forms the nitramine $C_6H_4:C_2O_2:N.NPh.NO_2$ [148°]; but in boiling HOAc it forms the di-nitro- derivative C14H8N4O6 [182°]. Nitrous acid passed into phthalyl-phenyl-hydrazine suspended in ether forms the nitrosamine C.H.:C.O.:NPh(NO) alcoholic solution of isatoic acid (Meyer, J. pr. [154°].—5. Alcoholic NH, at 80° forms unstable NH, CO.C. H., CO.NH.NHPh [146°], which decomposes at 170° into NH, and (\$\beta\$)-phthaly!-NHPh.NH.CO.C. H., NH, [151°]. Formed by

(B) - Phthalyl derivative

 $C_0H_4:C_2O_2 < \stackrel{NH}{NPh}$. [210°]. Formed at the same time as the (a)-isomeride by heating phenylhydrazine with phthalimide or phthalic anhydride, being the chief product when the temperature is above 163° (Pellizzari, G. 16, 204; 17, 278; Hötte, J. pr. [2] 35, 284). Monoclinic tablets (from benzene) or prisms (from alcohol), sol. alkalis. Ammoniacal AgNO, ppts. C, H4:C2O2:N2PhAg.

Reactions.-1. NaOMe and MeI at 100° form C₆H₄:C₂O₂:N₂PhMe [125°], which is decomposed by conc. HClAq into phthalic acid and NHPh.NHMe.—2. KOEt and EtI give C₈H₄O₂:N₂PhEt [106°].—3. BzCl at 200° forms

C₈H₄O₂:N₂PhAc [122°].

Phthalyl derivative C_sH₄(CO.NH.NHPh)₂. [191°]. Formed by heating phenyl-hydrazine (2 mols.) with phthalic anhydride (1 mol.) at 150° (H.). White tables. At 170° it gives the (β)-phthalyl derivative [210°].

Nitrosamine NH2.NPh.NO. Formed by adding NaNO, to an aqueous solution of phenylhydrazine hydrochloride at 0° (Fischer, A. 190, 92). Pale-yellow plates. Very poisonous. Converted by hot alcoholic potash into diazobenzene-Alcoholic HCl acts in like manner. imide. Exhibits Liebermann's reaction.

Dicyanide C_aH_aN₄ i.c. NH₂.NPh.C(NH).CN? Separates as colourless flakes when cyanogen is passed into an emulsion of phenyl-hydrazine and cold water (Bladin, B. 18, 1544). Monoclinic laminæ, v. sol. alcohol and ether. Decomposes above 160°. Yields a formyl derivative C₈H₂(CHO)N₄ [193°].

Reactions. -1. By heating with Ac₂O it is converted into C₁₀H₈N₄ [108°], which on saponification yields an acid C₁₀H₉N₃O₂ [177°], which on heating loses CO2, leaving only C2H2NO3 (c. 240°).-2. By heating with propionic anhydride it is converted into C₁₁H₁₀N₄[38°].—3. By treatment with nitrous acid, followed by boiling with water, it yields C₈H₅N₅, crystallising in needles [56°], v. sl. sol. water.—4. Acetic aldehyde in alcohol and HCl give CH3.CH:N.NPh.C(NH).CN [96.5°], crystallising in pale-yellow plates, v. e. sol. alcohol and ether, insol. water; converted by alcoholic AgNO₃ into phenyl-methyl-triazole carboxylic nitrile [109°] (Bladin, B. 25, 184).-5. Benzoic aldehyde forms the corresponding CHPh:N.NPh.C(NH).CN [129.5°], crystallising from alcohol in prisms, oxidised by FeCl, to C₁₈H₁₀N, [156°] (Bladin, B. 22, 796). -6. Pyruvic acid in alcohol forms the above phenyl-methyl-triazole carboxylic nitrile on warming gently (Bladin, B. 19, 2598).—7. Boiling acetoacetic ether forms the compound CO₂H.CH₂CMe:N.NPh.C(NH).CN [209°], the oily ether of this acid, and phenyl-methyltriazole carboxylic nitrile N NPh.C.CN [109°] (Pladin, B. 25, 190). The acid forms the salts KA' and NH, A', and is converted by nitrous acid CO₂H.C(NOH).CMe:N.NPh.C(NH).CN [218°], and by potash into the compound CO₂H.CH₂.CMe:N.NPh.CO.CN [188°], from CO.H.CH. CMe: N.NPh.CO.CN [188°], from which potassium nitrite and HClAq form CO.H.C(NOH).CMe: N.NPh.CO.CN [209°].

References.—Amido-, Iodo-, Nitro-, and Oxy-PHENYL-HYDRAZINE

u-Di-phenyl-hydrazine NPh2.NH2. [34·5°]. (220° at 40 mm.) (Stahel, A. 258, 243). Formed by reducing di-phenyl-nitrosamine NPh, NO in alcoholic solution with zinc-dust and HOAc (E. Fischer, A. 190, 174). Monoclinic tables (from ligroin), partially decomposed on distillation into diphenylamine and NH₃. Conc. H₂SO₄ forms a deep-blue solution. Decomposed by nitrous acid into NPh2.NO and N2O. Reduces HgO, forming in the cold crystalline tetraphenyl-tetrazone NPh, N:N.NPh, [123°], but if the temperature rises the products are N and diphenylamine. Benzoic aldehyde NPh...N:CHPh [122°]. With salicylic aldehyde, furfuraldehyde, glucose, mannose, galactose, and rhamnose it forms di-phenyl-hydrazides, melting at 139°, 90°, c. 162°, c. 155°, 157°, and 134° respectively.

Salts.—B'HCl. Needles, v. sl. sol. cold water. $-B'_{2}H_{2}SO_{4}$: needles, m. sol. dilute $H_{2}SO_{4}$.

FormylderivativeNPh₂.NH.CHO. [116.5°]. Formed by boiling the formyl derivative of phenyl-hydrazine with ammoniacal CuSO, (Gattermann, B. 25, 1075). Needles, sl. sol. hot water and ligroïn, v. sol. alcohol.

Acetyl derivative NPh_x.NHAc. [185°]. Formed by boiling NHPh.NHAc with HOAc and

Cu(OAc)₂ (Tafel, B. 25, 413).

Propionyl derivative. [178°]. Needles. Benzoyl derivative NPh, NHBz. [189°] (G.); [192°] (Fischer, A. 190, 178). Necdles (from acetone), m. sol. alcohol.

Phthalyl derivative C,H,O,:N.NPh, [155°]. Formed by heating di-phenyl-hydrazine with phthalic anhydride at 160° (Hötte, J. pr.

[2] 35, 271).

s-Di-phenyl-hydrazine NHPh.NHPh. drazo-benzene. Mol. w. 184. [131°]. H.C. 1,598,000 (from diamond) (Petit, A. Ch. [6] 17, 163). S. (alcohol) 5 at 16° (Moltschanowsky, C. J. 42, 905). Prepared by passing H₂S into a solution of azobenzene Ph.N2.Ph in alcoholic NH_3 (Hofmann, Pr. 12, 576). Formed also by reducing nitro-benzene with sodium-amalgam or zinc-dust (Alexejeff, Z. 1868, 497; Glaser, A. 142, 364). Laminæ, sl. sol. water, m. sol. alcohol and other. Yields aniline and azobenzene on distillation.

Reactions.-1. HClAq converts it into the isomericdi-p-amido-diphenyl (benzidine). H2SO4 acts in like manner. -2. Readily oxidised to azobenzene, even by moist air. -3. Nitrous acid, passed into an alcoholic solution at 0°, yields a nitroso- compound which, when warmed, violently decomposes into NO and azobenzene (Baeyer, B. 2, 683).—4. On heating with BzCl it yields di-benzoyl-di-p-amido-diphenyl; formic acid gives di-formyl-di-p-amido-diphenyl (Stern, B. 17, 379); and phthalic anhydride yields diphthalyl-di-amido-diphenyl (Bandrowski, B. 17, 1181).—5. Boiling benzoic aldehyde forms azobenzene, but benzoic aldehyde and ZnCl, gives C₁₂H₈(N:CHPh)₂ (Clève, Bl. [2] 45, 188). Benzoic aldehyde, heated with hydrazobenzene at 120°-150°, forms' benzhydrazoïn' CHPh $\stackrel{
m NPh}{\sim}$ The o-nitro- derivative CaH4(NO2).CHN2Ph2 of this body melts at 66° .- 6. Furfuraldehyde forms

 $C_4H_3O.CH < \stackrel{NPh}{NPh}$ [59°] (Cornelius a. Homolka, B. 19, 2240).—7. Phenyl cyanate (2 mols.) and some benzene at 150° forms the compound ${\bf NHPh.CO.NPh.NPh.CO.NHPh}$ [220] (Goldschmidt a. Rosell, B. 23, 490).

Acetyl derivative NHPh.NPhAc. [159°]. Formed from s-di-phenyl-hydrazine and Ac2O in the cold (Stern, B. 17, 380). Needles (from alcohol). Decomposed by heat into acetanilide

and azobenzene.

Di-acetyl derivative NPhAc.NPhAc. [105°]. Formed from s-di-phenyl-hydrazine and Ac O (Schmidt a. Schultz, B. 12, 485; A. 207, 326). Yellowish trimetric crystals (from alcohol); a:b:c=67:1: 56, not affected by dilute HClAq.

Chloro- derivatives

 $C_aH_bNH.NHC_aH_bCl.$ [90°].— $N_2H_a(C_aH_bCl[1:3])_2$ [94°].— $N_2H_a(C_aH_bCl[1:4])_2$ [122°] (Claus a. Heumann, B. 13, 1181, 19, 1688; Laubenheimer, B. 8, 1624).

Bremo-derivative C.H. NH.NH.C.H.Br. [115°]. Formed from benzene-azo-p-bromobenzene and alcoholic ammonium sulphide (Janowsky a. Erb, B. 20, 364). Tables. Converted by H2SO4 into bromo-benzidine

Di-bromo- derivatives N2H2(C6H4Br.)2. o. [82°]. m. [109°]. p. [130°] (J. a. E.; Gabriel,

B. 9, 1406; Werigo, A. 165, 192).

Iodo- derivatives v. vol. iii. p. 41. Nitro-derivatives v. vol. iii. p. 607. Oxy-derivatives v. vol. iii. p. 743.

Chloro-nitro-derivative NHPh.NHC, H3Cl(NO2) [1:3:6]. [135°-140°]. Formed from chloro-di-nitro-benzene and phenyl-hydrazine in alcohol (Willgerodt a. Ferko, J. pr. [2] 37, 355). Red prisms.

NHPh.NH.CoH,.NHAc [146°] is got by treating C.H.,N.C.H.,NHAC with alcoholic ammonium sulphide (Schultz, B. 17, 463). Yellowish plates (from dilute alcohol). Re-oxidised to the

azo- compound by air.

Di-m-amido-derivative N₂H₂(C₀H₁,NH₂)₂. [141°] (Gebek, A. 251, 193). Got by treatment of m-nitro-aniline with alcoholic potash and zinc-dust (Graeff, A. 229, 341), or with sodium-amalgam (Haarhaus, A. 135, 164). Golden needles (from alcohol). Yields C₁₁H₂Br₂N₄ [150°] and C₁₂H₁₂Ac₂N₄ [247°].— B"H₂Cl₂.—B"H₂PtCl₈.—B"2HNO₃.—B"H₄SO₄.— B"H₂Cl₂O₄. Prisms, v. sl. sol. water.

Di-p-amido-derivative. [145°]. Got by reducing N2(CeH1.NO2)2 with alcoholic ammonium sulphide (Gerhardt a. Laurent, A. 75, 74; Lermontoff, B. 5, 232). Yellow crystals. Yields quinone on oxidation. — B"H₂Cl₂. - B"2HNO₃.

Di-diphenyl-hydrazine v. Hydrazo-diphenyl. PHENYL-HYDRAZINE o-CARBOXYLIC ACID C, H, N2O2 i.e. NH2.NH.C, H4.CO2H. Hydrazido-benzoic acid. Formed from o-amidobenzoic acid by diazotising and treating the product with Na₂SO₃ (E. Fischer, B. 13, 679; A. 212, 333). Needles, sol. hot water. Reduces Fehling's solution in the cold. — HA'HCl: needles, sol. hot water.

Anhydride C,H_eN₂O. [242°]. Formed by heating the acid at 220°. Monoclinic plates (from alcohol); a:b:c = 1.072:1:.664; $\beta = 75^{\circ} 18'$.

Sublimes in white needles.—NaC, H, N,O xaq: silvery plates. — B'HCl: white needles. B'HgCl₂: needles (from water). Yields an acetyl derivative C,H,Ac,N2O [112°].

Phenyl-hydrazine m-carboxylic acid

N2H3.C6H1.CO2H. [186°]. Formed by the action of tin and HClAq on CaH, (CO2H).N:N.SO3K, which is got by warming the nitrate of m-diazobenzoic acid with aqueous K2SO3 (Griess, B. 9, 1657; Roder, A. 236, 164). Plates, sl. sol. hot Aq.

Reactions.-1. Acetone forms C10H12N2O2 [150°], which gives EtA' [91°].—2. Benzoic aldehyde forms CHPh:N.NH.C.H .. CO.H [172°].-3. Pyruvic acid reacts with formation of CO₂H.CMe:N.NH.C₀H₄.CO₂H [208°], which forms whetstone-shaped crystals (containing aq), and gives EtA' [102°].-4. Glucose, heated with the hydrochloride and NaOAc, gives C20H22N4O8 [208°]. - 5. Phenyl-thiocarbimide forms the compound C₁₁H₁₃N₃SO₂ [205°]. Salts.—HA'HCl.—BaA₂' 4aq. Nodules.

Phenyl-hydrazine p-carboxylic acid

N₂H₃.C₆H₄.CO₂H. [220°-225°]. Formed, in the same way as its isomerides, from p-amido-benzoic acid (Fischer, A. 212, 337). Needles or plates (from water).—HA'HCl: sl. sol. cold water.

s-Di-phenyl-hydrazine di-o-carboxylic acid $C_{14}H_{12}N_2O_4$ i.e. $CO_2H.C_6H_4.NH.NH.C_6H_4.CO_2H.$ Mol. w. 272. [205°]. Formed by reducing o-azoxy-benzoic acid with sodium-amalgam (Griess, B. 7, 1609; Homolka, B. 17, 1904). Small colourless plates. Oxidised by moist air to o-azobenzoic acid.

s-Di-phenyl-hydrazine di-m-carboxylic acid. Hydrazo-benzoic acid. Formed by boiling m-azobenzoic acid with FeSO, and NaOHAq (Strecker, A. 129, 141). Insol. water, sl. sol. hot alcohol. In alkaline solutions it absorbs O, being oxidised to azohenzoic acid. Boiling conc. HClAq forms azobenzoic acid and amidobenzoic acid. Yields orange-red crystals of BaA". On warming with SnCl₂ and HClAq at 100° it yields di-amido-di-phenyl dicarboxylic acid and two other acids. One of these C11H10N2O3 crystallises from hot water in yellowish needles and forms BaA', 7aq and HA'HCl. The other acid C14H12N2O4 crystallises in yellow needles melting above 290°, and forms NaA'4aq, BaA'₂2aq, HA'HCl, HA'HBr, and (HA'₂)₂H₂SO₄ (Kusseroff, B. 23, 913).

s-Di-phenyl-hydrazine di-p-carboxylic acid C, H, N, O,. Formed by boiling p-azobenzoic acid with FeSO, and NaOHAq (Reichenbach a. Beilstein, A. 132, 137; Billfinger, A. 135, 152). Needles (from alcohol), easily oxidised in alkaline

solution.

s-Di-phenyl-hydrazine tetra-carboxylic acid $[1:4:2]C_6H_3(CO_2H)_2.NH.NH.C_6H_3(CO_2H)_2$ [2:1:4]. Formed by reducing azoxyterephthalic acid in alkaline solution with sodium-amalgam (Homolka a. Löw, B. 19, 1092). White crystalline pp., v. sol. alcohol and ether.

PHENYL-HYDRAZINE m-SULPHONIC ACID C₀H₄(N₂H₂).SO₃H. Formed from m-amidobenzene sulphonic acid by diazotisation and subsequent reduction by SnCl, at 0° (Limpricht, B. 21, 3409). Crystals (containing 2aq), sl. sol. cold water, nearly insol. alcohol. Converted in alcoholic solution by nitrous N,C,H,SO,H. acid into

Phenyl-hydrazine p-sulphonic acid

[1:4]C.H.(N.H.).SO.H.

Formation.—1. From diazobenzene p-sulphonic acid by warming with aqueous K,SO₃ followed by HCl (E. Fischer, A. 190, 74).—2. By heating phenyl hydrazine sulphate with ClSO, H at 160° (Limpricht, B. 18, 2196).—3. Together with alcohol, by heating phenyl-hydrazine ethyl-sulphate at 160°.—4. By heating phenyl-hydrazine (1 pt.) with H₂SO₄ (6 pts.) at 100° (Gallinek a. Richter, B. 18, 3172).

Properties.—Needles (containing $\frac{1}{2}$ aq), sl. sol. water. By boiling with CuSO, it is neatly oxidised to N and benzene sulphonic acid. With acetoacetic ether and dilute (50 p. c.) HOAc at 155° it forms $CMe < \stackrel{N-N.C_6H_4.SO_3H}{CH_2.CO}$ (Möllen-

hoff, B. 25, 1849), which crystallises with aq. Salts. - NaA' 12aq. - NH, A'. - BaA'2 5aq. -

PbA'₂ 2aq.—ZnA'₂ 4aq: minute tables.

Phenyl-hydrazine v-sulphonic acid. Potassium salt. C.H.,NH.NH.SO,K. Formed from diazobenzene nitrate and K.,SO, (Römer, Z. 1871, 481) and also by heating phenyl-hydrazine with K₂S₂O₇ (Fischer, A. 190, 97). Crystals (containing aq), sl. sol. cold water and alcohol. Oxidised by HgO to C₆H₃.N₂.SO₃K. Split up by boiling HClAq into phenyl-hydrazine and KHSO.

Phenyl-hydrazine disulphonic acid NH₂.NH.C₆H₃(SO₃H)₂. Two acids of this formula are got by further sulphonation of the mand p-sulphonic acids respectively (L.). They both yield BaA" and BaH2A"2.

s-Di-phenyl-hydrazine m-sulphonic Formed by heating C,H,NH.NH.C,H,SO,H. the disulphonic acid with water at 200° for 3 days (Limpricht, B. 11, 1048). Yellow needles (containing $2\frac{1}{2}$ aq). — KA' 4aq. — PbA'₂ 3aq. — BaA'₂ 4aq: yellow plates, v. sol. water. Chloride C₁₂H₁₁N₂SO₂Cl. [240°].

Di-bromo- derivative

C.H.Br.NH.NH.C.H.Br.SO.H. Formed as a byproduct in the oxidation of CaH2Br2(NH2)SO3H [1:3:6:4] by KMnO₄ (Limpricht, B. 18, 1425).-KA' aq: long needles.

s-Di-phenyl-hydrazine di-m-sulphonic acid $N_2H_2(C_6H_4.SO_3H)_2$. S. .079 at 22°. Formed by reduction of nitro-, azo-, or azoxy-benzene-sulphonic acid (Limpricht, B. 11, 1048; 21, 3409; 23, 1057; Brunnemann, A. 202, 344; Mahrenholz a. Gilbert, A. 202, 337; Neumann, B. 21, 3419). White needles, v. e. sol. water, almost insol. alcohol and ether. Readily reduces ammoniacal AgNO, and Fehling's solution. HClAq immediately forms benzidine disulphonic

acid.—K₂A" aq.—BaA" 2aq.

Amide NH(O₆H₄.SO₂NH₂).NH.C.H₄.SO₂NH₂).

[248°]. Formed by reducing N₂(C₆H₄.SO₂NH₂)2 with zinc-dust and NH₃Aq or HOAc (Limpricht a. F. Meyer, A. 268, 132). White needles (from 50 p. c. acetic acid), almost insol. water, ether, and toluene. Reduces ammoniacal AgNO, forming a mirror. HClAq converts it into the amide of diamido-diphenyl (benzidine) disulphonic acid, m-amido-benzene sulphonic amide and N₂(C₆H₄.SO₂NH₂)₂. — N₂H₂(C₆H₄.SO₃NHK)₂ 1½aq. Thin needles or plates quickly oxidised by air.— $N_3H_2(C_6H_4)SO_5NHNs)_2$ $2\frac{1}{2}$ aq : needles.

s-Di-phenyl-hydrazine di-p-sulphonic acid N₂H₂(C₆Ĥ₄.SO₅H)₂. Obtained, in the same way as the preceding acid, from amido-benzene p-sulphonic acid.—BaH₂A"₂.—BaA": crystalline.

Isomeride .- V. BENZIDINE TETRASULPHONIC ACID, vol. i. p. 174. DI-PHENYL-HYDRAZINE v-DI-THIO-

CARBOXYLIC ACID NPh2.NH.CS.SH. [c. 109°]. Formed by dissolving NPh2.NH2 in CS2 (Stahel, A. 258, 249). Golden prisms, v. sol. alcohol, insol, water.

s-Di-phenyl-hydrazine di-m-thiosulphoniq acid N2H2(C8H4.SO2SH)2. Formed, together with N₂(C₆H₄.SO₂.SH)₂, by adding N₂(C₆H₄.SO₂Cl)₂ to a cold saturated solution of Ba(SH)2 (Limpricht, B. 18, 1470; A. 220, 354). White amorphous pp., becoming sticky on boiling with water. Oxidised by $KMnO_4$ to $N_2(C_6H_4.SO_3H)_2$.—BaA" 2aq: minute needles, sl. sol. cold water.

TRI - PHENYL - HYDRAZOÏN is hydrazoïn, v. s-Di-phenyl-hydrazine, Reaction 5. DI-PHENYL-HYDROQUINONE C18H14O2 i.e. C₆H₂Ph₂(OH)₂ [5:2:4:1]. [219°]. Formed by reducing di-phenyl-quinone with aqueous SO₂ (Müller a. Pechmann, B. 22, 2131). Colourless cubes (from dilute alcohol).

PHENYL-IMESATIN v. ISATIN. PHENYL-IMIDO-DIACETIC ACID

NPh(CH₂.CO₂H)₂. [150°-155°]. A product of the action of aniline on chloro-acetic acid (P. Meyer, B. 14, 1325; Hausdörfer, B. 22, 1796). Formed also by heating phenyl-amido-acetic acid with chloro-acetic acid and NaOAcat 120°-130° (H.). Needles, v. sol. hot water and alcohol. NPhH₃A'. Needles (from alcohol). Decomposes at 151°.

Mono-anilide

CO2H.CH2.NPh.CH2.CO.NHPh. [218°]. Formed together with the dianilide by heating the acid with aniline at 175°. Needles. Its ether CO2Et.CH2.NPh.CH2.CO.NHPh [122°] is got by heating the anilide of phenyl-amido-acetic acid (q.v.) with ether and NaOAc at 140°.

Di-anilide NPh(CH₂.CO.NHPh)₂. [218°].

Needles, v. sl. sol. alcohol.

TRI - PHENYL - TRI - IMIDO - BENZENE $C_{24}H_{21}N_s$ i.e. $C_6H_5(NHPh)_5$. [242°]. Formed by warming $C_6H_6Cl_6$ with aniline (Mohr, M. 11, 22). Golden plates, v. sol. chloroform, insol. water and alcohol. Conc. H₂SO₄ gives a dark-blue colour changed to violet by potash.

PHENYL - IMIDO - BENZYL - MALONIC

ETHER O₆H₈.C(NPh).CH(CO₂Et)₂. [75°]. Formed, together with (O₆H₅.C(NPh))₂C(CO₂Et)₂, by the action of sodium malonic ether on the compound C.H. CCl:NPh (Just, B. 18, 2624). Decomposes at 150° into alcohol and oxyphenyl-quinoline carboxylic ether.

DI-PHENYL IMIDO-DI-ETHYL DISUL-PHONE NH(CH₂.CH₂.SO₂.C₆H₅)₂. [78°]. Got by heating C₂H₄(SO₂.C₆H₅)₂ with NH₃Aq is sealed tubes (Otto, J. pr. [2] 30, 324). Triclinic plates (from alcohol).—Nitrate: [190°].—B'HCl. [193°].—B',H,PtCl,: plates, sl. sol. water. PHENYL-IMIDO - METHENYL-o-AMIDO-

 $C_{\bullet}H_{\bullet} < NH > C:NPh.$ PHENOL

Formed by heating C₆H₄< O >CO with aniline at 210° (Von Chelmicki, J. pr. [2] 42, 440). Needles, insol. water, sl. sol. cold alcohol.

DI-PHENYL-DI-IMIDO-NAPHTHOL v. (β)-

NAPHTHOQUINONE, Reaction 5.

PHENYL-IMIDO-DINAPHTHYL . PHENYL DINAPHTHYLENE AMINE.

PHENYLIMIDO-PHENYL-METHYL-THI-NPh < C(NPh).S CMe = CHAZOLE DIHYDRIDE

[138°]. Formed from di-phenyl-thio-urea and chloro-acetone (Traumann, A. 249, 51). Yellowish prisms, insol. water, v. sol. ether.

DI - PHENYL - DI - IMIDO - DI - PHENYL -OIAZTHIOLE TETRAHYDRIDE C.H. N.S i.e. S<C(NPh).NPh . [181°]. Got by oxidising diphenyl-thio-urea with H₂O₂ (Hector, B. 23, 357). PHENYL-IMIDO-PROPIONIC ACID CH₂.C(NPh).CO₂H. Anilpyruvic acid. [122°].

Formed from aniline and pyruvic acid in ether (Böttinger, B. 16, 1924; A. 188, 336; 263, 126). Crystals, v. sol. water. Decomposed by boiling water. Br yields C₁₈H₁₂Br₂N₂O₂ [264°].

Isomeride v. IMIDO-PHENYL-PROPIONIO ACID. PHENYL-IMIDO-SUCCINIC ACID

CO₂H.CH₂.C(NPh).CO₂H. [150°]. Formed by the action of dilute (1 p.c.) KMnO₄ on 'pyranil-pyroic' acid (Anschütz, B. 22, 738; of. Reissert, B. 21, 1942). Decomposed on fusion.

PHENYL-IMIDO-THIAZOLE DIHYDRIDE

 $C(NPh) < S - CH \\ NH.CH$ [124°]. Formed from CH₂Cl.CHCl.OEt and phenyl-thio-urea (Näf, A. 265, 125). White needles, sol. alcohol. Yields a nitrosamine C₂H₁(NO)N₂S [58°]. Di-phenyl-imido-thiazole dihydride

C(NPh) < CH CH. [105°]. Formed from dichloro-di-ethyl oxide and di-phenyl-thio-urea.

DIPHENYLINE v. DI-AMIDO-DIPHENYL. PHENYL-INDAZINE C13H10N2 i.e.

 $C_eH_i < N$ NPh. [84°]. (345° uncor.). Formed by reducing o-nitro-benzyl-aniline with tin and HClAq (Paal, B. 23, 2640; 24, 961). Needles (from alcohol). Not affected by distillation over red-hot zinc-dust. Reduced to a dihydride [98°] by adding Na to its alcoholic solution. Yields a methylo-iodide B'MeI [188°].

PHENYL-INDOLE C.H. CH NPh CH. (327° Formed by heating its carboxylic acid [176°] (Fischer a. Hess, B. 17, 567; Pfülf, A. 239, 221). Oil, v. sol. alcohol. Its solution in HClAq colours pine-wood intensely violet.

Phenyl-indole $C_6H_4 < \begin{array}{c} CH \\ NH \end{array} > CPh$.

phenanthridine. [186°]. (above 360°). Formation.—1. By boiling ω-bromo-acetophenone or phenyl-ω-amido-acetophenone with aniline (Möhlau, B. 14, 173; 15, 2480; 18, 165). 2. By heating the phenyl-hydrazide of acetophenone or of phenyl-acetic aldehyde with ZnCl2 at 180° (Fischer, B. 19, 1565; 21, 1071; A. 236, 135).—8. By passing benzylidene-o-toluidine through a red-hot tube (Etard, Bl. [2] 39, 531). 4. By the action of zinc-dust and NH, Aq on $C_6H_4(NO_2).CH_2.CO.C_6H_5$ (Pictet, B. 19, 1065).

Properties. - Colourless plates, insol. water. May be sublimed. Colours pine-wood violetblue. Yields B'C.H.N.O, [127°]. Nitrous acid forms a nitroso-derivative $C_eH_4 < \frac{C(NO)}{NH} > CPh$ [c. 258°], which yields B'HCl, B'HNO, and NaC, H,No, and may be reduced to amidophenyl-indole [174°].

Dihydride C,4H,1N. [46°]. Formed by

boiling phenyl-indole with zinc-dust, alcohol, and HCl. Colours pine-wood orange. Yields a nitrosamine.

 $C_{\bullet}H_{\bullet} < \stackrel{CPh}{\sim} CH.$ Phenyl-indole [89°]. Formed by the action of alcoholic HCl on the phenyl-hydrazide of phenyl acetic aldehyde (Fischer, B. 21, 1811). White plates (from ligroïn), v. sol. alcohol. Converted by heating for fifteen minutes with ZnCl, into the preceding isomeride [186°]. Yields a picrate [107°] and a nitrosamine $C_{\bullet}H_{\bullet} < \stackrel{CPh}{N(NO)} > CH$ [61°] which gives Liebermann's reaction (Ince, A. 253, 87).

Di-phenyl-indole C_eH, CPh CPh. [123°]. Formed by heating the phenyl-hydrazide of deoxybenzoin with ZnCl₂ (Fischer, B. 19, 1566; A. 236, 136). Colourless crystals, v. sol. alcohol. Does not give the pine-wood reaction. Its picrate crystallises in dark-red needles.

Di-phenyl-indole $C_8H_4 < \begin{array}{c} CH \\ NPh \end{array} > CPh$. (above 360°). Got by heating the u-di-phenyl-hydrazide of acetophenone with ZnCl, at 175° (Pfülf, A. 239, 223). Oil. Colours acidified pine-wood bluish-violet.

PHENYL-INDOLE CARBOXYLIC ACID

 $C_{15}H_{11}NO_2 \quad i.e. \quad C_{5}H_{4} < \stackrel{C(CO_2H)}{N(C_{5}H_{5})} > CH.$ [176°]. Formed by heating the di-phenyl-hydrazide of pyruvic acid with HOAc and HClAq (E. Fischer, B. 17, 567). Needles, v. sl. sol. water, v. sol. alcohol. NaOCl in the cold forms phenyl- visatin C,H, CO,NPh>CO. [134°] (Pfülf, A. 239, 222).

INDOXAZINE $C_{\bullet}H_{\bullet} < CH > N$. PHENYL [84°]. Got from o-bromo-benzophenone and an

alkaline solution of phenyl-hydrazine (Cathcart a. V. Meyer, B. 25, 1498). Large crystals. PHENYL-INDULINE C₂₄H₁₇N₈ i.e.

-Ç:N CH:CH-C(NPh).CH:C.NPh C₆H₄. [231°]. Mol. w. (by Raoult's method) 320-370 (calc. 347). Formed from amido-phenyl-induline [152°] by treatment with nitrous acid (Fischer a. Hepp, A. 262, 257; 266, 263). Reddish tables, with blue reflex; v. sol. benzene, insol. ligroin. Heated with HOAc (5 pts.) and HClAq (25 pts. of 20 p.e.) at 165° for six hours it forms: (a) C₂₄H₁₈N₃O crystallising from benzene in steel-blue plates [218°] forming a brownish-red solution in HClAq, and converted by conc. HClAq at 200° into a compound C₁₈H₁₄N₂O₂; (b) the compound C₁₈H₁₄N₂O₂ [c. 280°] which crystallises from benzene or alcohol in brownish-yellow prisms or plates, and is both base and phenol. Its hydrochloride crystallises from conc. HClAq in prisms with green lustre; (c) benzolindone C₁₈H₁₂N₂O; (d) a body which dissolves in benzene with fiery brown fluorescence, and is perhaps carbazole-fluor-indine (Fischer a. Hepp, A. 266, 249).

Benzolindone C18H12N2O i.e. CH:CH.C:N CO.CH.C.NPh>C.H. Formed by heating phenyl-induline with HOAc and HClAq at 160°-200° (Fischer a. Hepp, A. 266, 253). Lustrous brown plates, insol. alkalis, sol. dilute HClAq, forming a rose-coloured solution. Its

solution in conc. H₂SO₄ is green. phenazine on distillation with zinc-dust.

PHENYL IODIDE v. IODOBENZENE.

PHENYL IODOMETHYL SULPHONE

 $\rm C_{e}H_{s}.SO_{2}.CH_{2}I.~[64.5^{\circ}].~Formed from benzene sulphinic acid, NaOEt, and CH_{2}I_{2}~at~100^{\circ}$ (Michael a. Palmer, Am. 6, 253). Prisms (from alcohol). NaOEt at 75° gives phenyl-methyl sulphone and products of oxidation (Otto, B. 21, 654).

PHENYL-ITACONIC ACID

CHPh:C(CO₂H).CH₂.CO₂H. [172°]. Formed by the action of Na or NaOEt on phenyl-paraconic acid (Fittig, A. 256, 68). Prisms, sl. sol. cold Yields phenyl-isocrotonic acid [179°] and y-oxy-phenyl-butyric lactone [37°] when boiled with diluted H.SO₄ (1:1). Br gives bromophenyl-paraconic acid C₁₁H₂BrO₄ [99°] and an isomeride [144°].—BaA" 2½aq.—CaA".—Ag.A".

Ethyl ether Et₂A". (315° uncor.). Oil.

Di-PHENYL-KETONE v. BENZOPHENONE.

Di-phenyl diketone v. Benzil.

Di-phenyl triketone C₀H₃.CO.CO.CO.CO.C₀H₄. [70°]. (248° at 40 mm.). Formed by distilling CBz₂Br.OAc in vacuo. Formed also by adding aqueous NaNO2 to a solution of the mono-oxim in HOAc, and boiling (Pechmann, B. 22, 852; 23, 3375). Hygroscopic golden needles (from ligroin), v. sol. all solvents. Give a blue colour with H₂SO₄ and benzene that contains thiophene.

Reactions .-- 1. Alkalis decompose it, forming benzoïn, benzoic acid, and CBzPh(OH).CO2H [118°].—2. Phenyl-hydrazine forms a phenylhydrazide C₂₁H₁₆N₂O₂ [c. 135°], benzene-azo-triphenyl-pyrazole $C_eH_s.N_2.C < CPh.NPh CPh.N$ [157°].

and a third compound [223°].—3. Aniline gives rise to CPh(NPh).C(OH)2.COPh [100°] and C(OH)₂(CPh:NPh)₂ [148°]. Hydrate Bz₂C(OH)₂. [90°]. Mono-oxim Bz₂C:NOH. [146°]. Formed

from Bz₂CH₂ and nitrous acid.

Tri-oxim (C₆H_s.C(NOH))₂C:NOH. [186°]. Formed from the triketone and hydroxylamine. Di-phenyl tetraketone

C_eH_a.CO.CO.CO.CO.C_eH_a. The hydrate [88°] of this body is formed by the action of conc. HNO_e on phenyl-glyoxal (Abenius a. Söderbaum, B.

DI-PHENYL-KETONE CARBOXYLIC ACID

v. BENZOYL-BENZOIC ACID.

Di-phenyl ketone dicarboxylic acid v. Benzo-

PHENONE DICARBOXYLIC ACID.

Di-phenyl diketone o-carboxylic acid C₆H₅.CO.CO.C₆H₄.CO₂H. Formed by oxidising deoxybenzoin o-carboxylic acid with alkaline KMnO₄ (Graebe, B. 21, 2003; 23, 1344). Occurs in a yellow modification [141.5°] and a white variety [125°-130°]. The yellow modification is nearly twice as sol, alcohol and chloroform as the white variety, and is converted into the white variety by dissolving in alkali and ppn. by acids. The white modification changes into the yellow one at 115°-125°.

Dicarboxylic acid v. DIPHTHALYLIC ACID. DI-PHENYL-KETOXIM v. Oxim of Benzo-

PHENONE. PHENYLLACTIC ACID v. OXY-PHENYL-PRO-PIONIC ACID.

PHENYL-LACTIMIDE v. vol. i. p. 180.

PHENYL-LUTIDINE-CARBOXYLIC ACID o. PHENYL-DI-METHYL-PYRIDINE-CARBOXYLIC ACID.

PHENYL-LUTIDONE v. OXY-PHENYL-DI-

METHYL-PYRIDINE.

PHENYL-MALEÏC ACID C₁₀H₈O₄ i.e. CO₂H.CPh:CH.CO₂H. Formed by dissolving its anhydride in water and extracting with ether (Alexander, A. 258, 77). Prisms, m. sol. water. Begins to lose water below 100°.

Anhydride CH .CO O. [119.5°]. Formed by distilling phenyl-malic acid. Needles, insol.

water, v. sol. chloroform and ether.

Di-phenyl-maleïc acid CO₂H.CPh:CPh.CO₂H. Stilbene dicarboxylic acid. Formed by saponification of its nitrile. Its ether is formed, together with di-phenyl-fumaric ether, by the action of Na on a-bromo-phenyl-acetic ether in ether (Reimer, B. 14, 1800; Rügheimer, B. 15, 1626). The free acid splits up at once, on liberation from its salts, into H₂O and anhydride. On distillation with lime it yields s-diphenyl-ethylene. H_2A'' : needles, v. sol. water. —Ag,A'': amorphous pp.—AgHA'': needles. Ethyl ether Et, A''. [54°]. Prisms. Imide $C_{16}H_{10}O_2(NH)$. [213°]. Needles,

almost insol. water, v. sol. alcohol.

[175°]. Phenylimide $C_{16}H_{10}O_2(NPh)$. Formed by heating the anhydride with aniline at 120° (Anschütz a. Bendix, A. 259, 63). Nitrile CN.CPh.CPh.CN. Dicyanostilbene.

[158°]. Formed by heating a-bromo-phenylacetonitrile alone or with alcoholic KCy (Reimer,

B. 14, 1798). Plates (from benzene).

Isonitrile. $[2\dot{4}2^{\circ}].$ Occurs in small quantity in the preparation of the nitrile by heating the product of bromination of phenylacetonitrile. Golden needles (from alcohol). Yields an acid [222°] on saponification.

Anhydride CPh.CO O. [155°]. (236° at 15 mm.). Formed by adding HCl to solutions of salts of the acid, or by heating di-phenylfumaric acid at 260° (Reimer, B. 13, 742). Trimetric needles (from dilute alcohol), a:b:c. = .693:1:.385, insol. water, sol. CS₂. May be sublimed. Its alkaline solution reduces AgNOs. Not changed by NaOHAq into the fumaroid form (Delisle, A. 269, 91). NaOHAq (15 p.c.) at 100° forms a polymeride melting much above 250°, insol. water and other solvents.

Reactions.-1. Reduced in alcoholic solution by zinc-dust and HCl to a mixture of the two diphenyl-succinic acids.—2. Fuming HNO, gives rise to an amorphous di-nitro- derivative (C₀H₄(NO₂))₂C₄O₂. -3. p-Tolyl-acetic acid forms C₄Ph₂O₂:CH.C₇H, [165°], crystallising in yellow needles.-4. Phenyl-acetic acid and NaOAc at 225° form CPh.CO——>O [176°], which may be reduced by HI and P to benzyl-di-phenylmaleïde CPh.CH(CH₂Ph) O [128°] (Gabriel a. Cohn, B. 24, 3228, 3854). The compound C₄Ph₂O₂(CHPh) combines with Br, forming C,Ph,BrO,(CHBrPh) [154°], which splits up at 155°-160° into HBr and C,Ph,O,(CBrPh) [165°]. The compound C,Ph,HO,(CH,Ph) is converted by potash into CO₂H.C.Ph₂.CH(OH).CH.Ph [174°], and by an alcoholic solution of NEtH, into CO(NHEt).C2Ph2.CH(OH).CH2Ph [196°].

compound C₄Ph₂O₂(CHPh) is converted by KOH into C₄Ph₂O₂(OH).CH₂Ph [185°], and in benzene reacts on PhSNa in ether, forming acetonyl-solution by HNO₂ into C₄Ph₂O₂(OH).CH(NO₂)Ph phenyl-sulphide PhS.CH₂CO.CH₃ [35°] (269°) [125°] (which yields an acetyl derivative [166°]), and the compound C4Ph2O2(NO2).CH(NO2)Ph [146°], which is converted by boiling alcohol into C₄Ph₂O₂:C(NO)Ph [177], a body that may be reduced by HI and P to iso-benzyl-diphenylmaleïde [118°]. Benzyl-di-phenyl-maleïde is converted by alcoholic NH, at 180° into CPh.CH(CH,Ph) NH [170°], which yields a nitrosamine [136°

PHENYL-MALIC ACID v. OXY-PHENYL-SUC-CINIC ACID.

PHENYL-MALONAMIC ACID v. MALONIC

DI-PHENYL - MALONIC ACID. Ethylether of the nitrile CPh₂(CN).CO₂Et. [59°]. Formed from CPh₂Cl.CO₂Et and HgCy₂ at 125° (Bickel, B. 22, 1537). Yellowish tables.

Reference.—DI-NITRO-PHENYL-MALONIC ETHER. PHENYL-MANDELIC ACID v. Phenyl derivative of Mandelic acid.

PHENYL-MELAMINE v. vol. ii. p. 322 PHENYL MERCAPTAN C.H.SH. Thio-

phenol. Phenyl-sulphydrate. Mol. w. 110. (168°) (L.); (172°) (F. a. C.). S.G. 2± 1·078.

Formation.—1. A product of the dry distilla-

tion of sodium benzene sulphonate (Kekulé, Z. 1867, 194; Stenhouse, A. 149, 248; Pr. 17, 62). 2. Together with Ph₂S and diphenylene sulphide by heating benzene with S and AlCl, at 75° (Friedel a. Crafts, C. R. 86, 884; A. Ch. [6] 14, 437).—3. By distilling phenol (10 mols.) with P₂S_a (1 mol.), the yield being 6 p.c. of the phenol used (Kekulé, Z. [2] 3, 193, 306; Beckmann, J. pr. [2] 17, 456).—4. From Ph₂S₂ and K₂S in alcohol (Otto a. Rössing, B. 19, 3129).—5. By distilling C.H.SO2Na with NaSH (Stadler, B. 17, 2080).

Preparation. - Phenyl ethyl dithiocarbonate, or a mixture of diazobenzene chloride solution and potassium ethyl dithiocarbonate (xanthate) is boiled with alcoholic potash until the solution remains clear on addition of water. The alcohol is then distilled off and the residue dissolved in water, mixed with a little H₂SO₄ and zincdust (to prevent oxidation), and distilled with steam. The distillate is extracted with ether. The yield is 75 p.c. of the theoretical (Leuckart, J. pr. [2] 41, 187).

Properties.-Oil, v. sol. alcohol and ether. Smells like garlic. Attacks the skin. Conc. H₂SO₄ forms a violet solution, changing on warming through cherry-red to blue (Baumann,

H. 5, 321; cf. Brunner, B. 4, 984).

Reactions.—1. Readily oxidised to Ph. S. 2. CH, CCl, and dilute (20 p.c.) NaOHAq form CH₃.C(SPh)₄, crystallising in white plates; v. sol. hot alcohol (Laves, B. 25, 353).—3. Acctoacetic ether and dry hydrochloric acid form CH₃.C(SPh)₂.CH₂.CO₂Et [58°], crystallising from alcohol in plates. Not attacked by boiling HClAq, but decomposed by alkalis into alcohol, PhSH, and CH₃.C(SPh):CH.CO₂H [177°] (Escales a. Baumann, B. 19, 1790).—4. NH₂.COCl forms NH₂.CO.NH.CO.SPh [218°] (Gattermann, A. 244, 43).—5. Acctone and HCl in the cold give Me₂C(SPh)₂ [56°] (Baumann, B. 19, 2804), which is oxidised by KMnO₄ and H₂SO₄

S.G. z 1.244 (Delisle, A. 260, 252). This body crystallises in prisms, sl. sol. hot water, v. sol. alcohol; combines with KHSO, yields a phenylhydrazide [82 5°], and is converted by oxidation into Ph.SO₂.CH₂.CO.CH, [57°]. Acetonylphenyl sulphide may be converted by successive pheny sulpinde hay be converted by successive treatment with HCy, alcoholic hydrochloric acid, and alcoholic potash into the acid PhS.CH₂:CMe(OH).CO₂H [97°], which yields CaA'₂ aq, BaA'₂ aq, and AgA', and is oxidised by KMnO₄ to Ph.SO₂:CH₂:CMe(OH).CO₂H [121°].—
7. Chloral forms CCl₂:CH(OH)(SPh) [53°], decomposed by best into its composed by the c composed by heat into its components (Baumann, B. 18, 886).—8. Combines with pyruvic acid, forming CH₂.C(OH)(SPh).CO₂H [87°], but in presence of gaseous HCl it forms the 'mercaptol' CH₃.C(SPh)₂.CO₂H [117°] which is not attacked by boiling HClAq or alcoholic potash. -9. Phenylglyoxylic acid forms C_eH_o.C(OH)(SPh).CO₂H converted by HCl gas into C_eH_o.C(SPh)₂.CO₂H [148°] (Escales a. Baumann, B. 19, 1789).—10. β-Acetylpropionic acid does not combine directly, but in presence of hydrochloric acid it forms CH₂.C(SPh)₂.CH₂.CH₂.CO₂H [69°], which is not attacked by alkalis, but is decomposed by dilute acids into PhSH and acetyl-propionic acid (E. a. B.).—11. CH₂CO.CH₂SPh forms CH₃C(SPh)₂CH₂SPh [55°], crystallising in cubes; v. sol. alcohol, insol. water; converted by oxidation in CH₃C(SPh)(SO₂Ph).CH₂SO₂Ph [157°], crystallising in needles, sl. sol. cold alcohol, split up by alkalis into Ph.SO₂CH₃, PhSH, and Ph.SO₂H (Autenrieth, B. 24, 170).—12. CBr₂(SO₂Et)₂ and NaOHAq form PhŚ.CH(SO₂Et)₂ [86°], crystallising in tables; v. sl. sol. hot water, v. sol. alkalis; converted by alcoholic NaOH and MeI into PhS.CMe(SO,Et), [113°], crystallising in white needles; m. sol. hot alcohol (Fromm, A. 253, 166; Laves, B. 25, 361). — 13. NaSPh reacts with isobutylene bromide, forming NaBr, Ph₂S₂, and isobutylene (Otto, B. 23, 1052)

Salts.-NaSPh. When Na is dissolved in an ethereal solution of PhSII containing phenol, a pp. of NaSPh is got while NaOPh remains in solution.—Hg(SPh)₂: colourless needles (from alcohol).—ClHgSPh: plates.—Pb(SPh)₂: yellow crystalline pp., split up on distillation into PbS and Ph₂S.—Cu(SPh)₂: pale-yellow powder.—

AgSPh: pale-yellow crystalline pp.

Methyl ether PhSMe. (188°). Formed from Pb(SPh)2 and MeI (Obermeyer, B. 20,

Ethyl ether PhSEt. (204°) at 744 mm. Formed from NaSPh and EtI at 120° (Beckmann, J. pr. [2] 17, 457), and by the action of PhSH on PhN₂Cl (Stadler, B. 17, 2078). Liquid with unpleasant smell.

Allyl ether v. Phenyl allyl sulphide. Acetyl derivative PhSAc. (229°). Formed from phenyl mercaptan and AcCl (Michler, A. 176, 177). Oil.

References. — Amido-, Bromo-, Chloro-, Chloro-, Nifro-, and Oxx- Phenyl-

MERCAPTAN.

PHENYL-MERCAPTURIC ACID C, H, NSO i.e. NHAc.CMe(SPh).CO2H. [143°]. Formed by reducing bromo-phenyl-mercapturic acid with sodium-amalgam (Baumann, B. 15, 1732; 18, 258; H. 5, 835; 8, 190). Tetrahedra, m. sol. Levorotatory in alcoholic solution. alcohol. Solutions of its salts are dextrorotatory. Decomposed by dilute H.SO, into HOAc and phenylcystein.—BaA', Saq: needles.

References .- Bromo- and Chloro- Phenyl-

MERCAPTURIC ACID.

PHENYL-MESITYL-CARBINOL v. PHENYL-TRI-METHYL-PHENYL-CARBINOL.

PHENYL-MESITYL-CARBINOL CARBOXY-LIC ANHYDRIDE v. TRI - METHYL-PHENYL-PHTHALIDE.

PHENYL MESITYL KETONE v. PHENYL TRI-METHYL-PHENYL KETONE.

PHENYL-METHACRYLIC ACID v. PHENYL-CROTONIC ACID.

PHENYL-METHACRYLIC ALDEHYDE

CHPh:CMe.COH. (150° at 100 mm.). Formed by adding NaOHAq to a mixture of benzoic aldehyde and propionic aldehyde (Miller a. Kinkelin, B. 19, 525). Oil. Reduces ammoniacal AgNO₃. Combines with bisulphite. Yields a phenyl-hydrazide [137°].

PHENYL-METHANE v. Toluene.

Di-phenyl-methane C₁₃H₁₂ i.e. CH₂Ph₂. Mol.

w. 168. [25°]. (c. 263°).

Formation.—1. Together with C.H. (CH2Ph)2, by boiling benzyl chloride with benzene and zinc-dust (Zincke, B. 4, 298; C. J. 24, 508, 688). 2. By the action of H2SO, and HOAc on a mixture of benzene and benzyl alcohol (Meyer a. Wurster, B. 6, 963).-3. By heating benzophenone with zino-dust (Staedel, B. 6, 1401; 7, 1480; cf. Barbier, C. R. 79, 812), or by reducing it with HI and P (Graebe, B. 7, 1624), or Zn and H₂SO₄ (Zincke, B. 10, 1473).—4. From benzene, CH₂Cl₂, and AlCl₃ (Friedel a. Crafts, Bl. [2] 41, 324; A. Ch. [6] 11, 264).—5. From CH₂(OMe)₂, benzene, HOAc, and H₂SO₄ (Baeyer, B. 6, 221).-6. By distilling barium di-phenylacetate with soda-lime (Jena, A. 155, 86).

Preparation.—By adding AlCl₃ (35 g.) to a mixture of benzyl chloride (100 g.) and benzene (500 g.) (Friedel a. Crafts, A. Ch. [6] 1, 478; Bl.

[2] 83, 337).

Properties.—Prisms, v. sol. alcohol and ether, smelling like oranges. According to Reissert (B. 23, 2242) it melts at 23°. Oxidised by chromic acid mixture to benzophenone. Yields diphenylene-methane when passed through a red-hot tube (Graebe, B. 7, 1623). Prolonged treatment with Cl in presence of I forms CCl, and C.Cl. (Ruoff, B. 9, 1048).

Tri-phenyl-methane C₁₉H₁₆ i.e. CHPh₂. [92°].

(330°) (Schwarz); (355°) (K. a. F.).

Formation.—1. By heating PhCHCl₂ with
HgPh₂ at 150° (Kekulé a. Franchimont, B. 5, 907).-2. By heating Ph₂CH(OH) with benzene and $P_{5}O_{5}$ at 135° (Hemilian, B. 7, 1203).—3. By heating (β)-benzpinacolin $C_{28}H_{20}O$ with sodalime at 300° (Thörner a. Zincke, B. 9, 65).—4. Together with a little CH₂Ph₂ by adding AlCl₃ a mixture of chloroform and benzene (Friedel a. Crafts, J. pr. [2] 16, 233; A. Ch. [6] 1, 496). 5. With other products, by the action of AlCl. on benzene mixed with CH2Cl2, with CCl4, with PhCCl₁, and with CPhHBr. CPhHBr (Magatti, B. 12, 1468; Schwarz, B. 14, 1526; Anschütz, A. 285, 208).—6. By heating benzoic aldehyde (100 g.) with benzene (147 g.) and ZnCl₂ (100 g.) at 250° for 7 hours (Griepentrog, B. 19, 1876; A. 242, 329).

Preparation.—AlCl₂ (500 g.) is slowly added to a mixture of benzene (1400 g.) and chloroform (400 g.). The mixture is finally heated on the water-bath, water is carefully added, and the oil dried and fractionally distilled. It is finally purified by conversion into the compound with benzene (Allen a. Kölliker, A. 227, 108; cf. Fischer, A. 194, 252; Schwarz, B. 14, 1516). If the benzene contains toluene the product will contain a homologue (Hanriot, Bl. [3] 1, 773).

Properties.—Plates (from alcohol), v. sol. benzene, ether, and CHCl_s, m. sol. cold alcohol. Crystallises from benzene in efflorescent rhombohedra CHPh_sC_eH_e [76°]. Oxidised by chromic acid to tri-phenyl-carbinol [159°]. AlCl, at 120° decomposes it, but below 80° it forms CH₂Ph₂ (Friedel a. Crafts, C. R. 100, 692). On exhaustive chlorination with SbCl, it yields CCl, and C₆Cl₈ (Merz a. Weith, B. 16, 2876). Na has no action at 300°, but K at 200° forms a red substance (CKPh₃). K at 250° forms a dark mass, whence HClAq liberates C₁₉H₁₄ [148·5°] (350°) (Hanriot a. Saint-Pierre, C. R. 108, 1119; Bl. [3] 1, 774). The potassium derivative CKPh, absorbs CO, at 200°, yielding CPh, CO, H [264°]. Benzyl chloride acting on CKPh, forms the compound CPh₃.CH₂Ph [140°]. BzCl forms C₂₀H₁₈O [172°] which may be reduced by HI and P to

C₂₆H₂₀ [234°]. References. — Amido-, Bromo-, Chloro-, Methyl-amido-, Nitro-, Nitro-amido-, Oxyamido-, and Oxy- Phenyl-methane.

DI-PHENYL-METHANE DI-O-CARBOXY-

LIC ACID CH₂(C₆H₄·CO₂H)₂. [255°]. Formed by reducing the lactone of CH(OH)(C₆H₄·CO₂H)₂ with HI and P (Graebe a. Juillard, A. 242, 253). Crystals (from alcohol). Yields anthraquinone on heating at 280°.—BaA" 6aq. S. 4.7.

Methyl ether Me₂A". [44°].

Isomeride v. BENZYL-ISOPHTHALIC ACID.

Di-phenyl-methane tricarboxylic acid CO₂H.CH(C₆H₄.CO₂H)₂. [220°]. S. 095 at 25°. Got by heating the lactone of the oxyacid CO₂H.C(OH)(C₆H₄.CO₂H)₂ with HI and P at 170° (Graebe a. Juillard, A. 242, 285). Crystals (containing aq). At 270° it forms a red compound C16H,O, [261°].

Methyl ether Me, A'". [145°]. Tri-phenyl methane o-carboxylic acid

CHPh₂.C₆H₄.CO₂II. [162°].

Formation.-1. By the action of NaOH and zinc-dust on di-phenyl-phthalide (Baeyer, B. 12, 644; A. 202, 52).—2. By heating phenylphthalide with benzene and AlCl, (Gresly, A. 234, 242).—3. By saponifying the nitrile, which is obtained from CHCl₂.C₆H₄.CN, benzene, and AlCl, (Drory, B. 24, 2572)

Properties.—Needles (from alcohol), insol. water. May be sublimed. Yields tri-phonylmethane when heated with Ba(OH)2. CrOs in

HOAc oxidises it to diphenyl-phthalide. Nitrile CHPh₂.C₆H₄.CN. [89°].

Tri - phenyl - methane - p - carboxylic acid. [162°]. Obtained by saponifying the nitrile, which is obtained by Sandmeyer's reaction from p-amido-tri-phenyl-methane [84°] (Fischer a Fränkel, A. 241, 364; Otto a. Fischer, B. 24, 729). Needles, v. sol. alcohol and ether.—AgA'.

An isomeric acid melting above 360° was obtained by Oppenheimer (B. 19, 2029) by oxidising the corresponding aldehyde with Ag.O.

Tri-phenyl-methane dicarboxylic acid $CHPh_2.C_6H_3(CO_2H)_2$ [1:2:5]. [280°]. Formed by reduction of the lactone of tri-phenyl-carbinol dicarboxylic acid by zinc-dust and NaOH (Hemilian, B. 16, 2375). Needles (from HOAc). Conc. H₂SO₄ forms a greenish-yellow solution, which on heating changes through green, blue, and violet to purple.—BaA"4aq.—Ag₂A": pp.

Tri-phenyl-methane dicarboxylic acid $CHPh_2 \cdot C_6H_3(CO_2H)_2$ [1:3:4]. [278°]. Formed by boiling di-phenyl-phthalide dicarboxylic acid with zinc-dust and NaOHAq (Hemilian, B. 19, 3068). Needles (from dilute alcohol). May be sublimed. Yields CHPh2 when distilled with Ba(OH). Re-oxidised by alkaline KMnO to diphenyl-phthalide dicarboxylic acid.—CaA" 2aq: needles. - Ag2A": curdy pp.

Reference. - OXY-DI-PHENYL-METHANE CARB-OXYLIC ACID.

DIPHENYL - METHANE DISULPHONIC ACID C₁₃H₁₀(SO₃H)₂. [59°]. Got by sulphonation (Doer, B. 5, 796). Deliquescent plates. -K₂A" aq: prisms (from dilute alcohol).-BaA". -CuA": green plates.

Tri-phenyl-methane trisulphonic acid $C_{19}H_{18}(SO_3H)_8$. Formed by warming CHPh₃ with fuming H_2SO_4 (Kekulé a. Franchimont, B. 5, 908; Hemilian, B. 7, 1205).— $Ba_3A'''_28aq$: needles, ppd. by adding alcohol to its aqueous solution.

DI-PHENYL-METHAZINE $C_{14}H_{12}O_2$ *i.e.* CHPh $\langle N \rangle$ CHPh. [93°]. Formed by shaking a dilute solution of hydrazine with benzoic aldehyde (Curtius, J. pr. [2] 39, 44). Light-yellow prisms, v. sl. sol. hot water, v. sol. hot alcohol. Decomposed by boiling into CHPh:CHPh and nitrogen, and by boiling dilute acids into benzoic aldehyde and hydrazine. Reduced in alcoholic solution by sodium-amalgam to di-benzyl-hydrazine, and by excess of Na to benzylamine.

PHENYL-METHENYL-AMIDINE v. BENZ-

Diphenyl-methenyl-diamine is described as DI-PHENYL-FORMAMIDINE.

PHENYL-METHENYL-AZIDINE C,H,N, i.e. NH₂.CH:N₂HPh. [225°]. Formed from phenylhydrazine and HCy (Fischer, B. 22, 1934). Insol. water, v. sol. hot alcohol. The nitrate and hydrochloride are crystalline.

PHENYL METHENYL DI-ETHYL TRISUL-**PHONE** $C_6H_5.SO_2.CH(SO_2.C_2H_5)_2$. [166°]. 2 at 15°. Formed by oxidising Phs.CH(SO₂Et)₂ with alkaline KMnO, (Fromm, A. 253, 167; Laves, B. 25, 362). Snow-white needles, v. sol. alkalis.-KA': prisms.-BaA'2: tables.

TRI-PHENYL METHENYL TRIKETONE v. TRI-BENZOYL-METHANE.

TRI-PHENYL METHENYL TRISULPHONE CH(SO₂Ph)₂. [215°]. Formed by oxidising PhS.CH(SO₂Ph)₂ with alkaline KMnO₄ (Laves, B. 25, 348). Tables, v. sol. alcohol, insol. ether, sm. sol. water.—KA': tablets.—BaA', —AgA'.

DI - PHENYL - DI - METHYL - ACETYLENE TETRAKETONE v. DI-BENZOYL-DI-ACETYL-ETHANE.

PHENYL-METHYL-ACRIDINE C.H., N i.e. $C_eH_e < CPh.C.CH:CMe$ N = C.CH:CH[136°]. heating phenyl-p-toluidine with benzoic acid and ZnCl₂ at 260° (Bonna, A. 239, 62). Needles, sl. sol. water. Its alcoholic solution shows greenish-blue fluorescence. Yields phenylacridine carboxylic acid [253°] on oxidation.— B'HCl: yellow needles.—B'HI.—B'2H2SO3.— B'C_sH₃N₃O₇. Red needles.

Phenyl-methyl-acridine dihydride

 $C_6H_4 < C_{NMe}^{CHPh} > C_6H_4$. [104°]. Formed by the action of MeI on phenyl-acridine dihydride; or by reduction of the methylo-chloride of phenylacridine with zinc-dust (Bernthsen a. Bender. B. 16, 1816). White needles or prisms. On oxidation with nitrous acid it gives the methylohydroxide of phenyl-acridine.

PHENYL-METHYL-ALLYL-PYRROLE

CH:CMe NC₃H₃. [52°]. (278°). Formed by heating the carboxylic acid [158°] (Lederer a. Paal, B. 18, 2595). Large colourless plates. Blue fluorescence. Extremely sol. all ordinary solvents.

PHENYL - METHYL - ALLYL - PYRROLF. $_{\mathrm{CO_2H.C}=\mathrm{CMe}}^{\mathrm{CH:CPh}} > \mathrm{NC_2H_5}.$ CARBOXYLIC ACID [158°]. Obtained by saponification of its ether, which is formed by the action of allylamine upon acetophenone-acetoacetic ether (Lederer a. Paal, B. 18, 2594). Short glistening prisms.

alcohol, ether, benzene, and acetic acid. PHENYL-METHYL-AMIDO-ACETIC C₀H₁₁NO₂ i.e. CH₂(NPhMe).CO₂H. Phenylmethyl-glycocoll. Formed by heating chloroacetic ether with dimethylaniline and saponifying the product by boiling HClAq (Silberstein, B. 17, 2661). Liquid.—HA'HCl: prisms.

CH₂(NPhMe).CO.NH₂. Amide [168°]. Formed by heating CH2Cl.CONH2 with NPhMe2 or NPhMeH. Crystals, sol. hot water. hydrochloride forms easily soluble prisms.

Methylo-chloride CH2(NPhMe2Cl).CO2H. Formed by warming chloro-acetic acid with NPhMe₂ in ether (Zimmermann, B. 12, 2206). Needles. Moist Ag₂O forms an hydroxide.

Amide of the methylo-chloride CH_(NPhMe,Cl).CO.NH,. Formed by heating chloro-acetamide with dimethylaniline in alcohol (S.). Crystals, decomposed by heat into

MeCl and CH₂(NPhMe).CO.NH₂.
PHENYL - METHYL - ω - AMIDO - ACETO-PHENONE C, H, CO.CH, NPhMe. Phenacylmethyl-aniline. [120°]. Formed by the action of NPhMeH or NPhMe, on ω-bromo-acetophenone (Staedel a. Siepermann, B. 13, 842; 14, 983; 21, 2196). Prisms (from alcohol). A solution of its hydrochloride gives a red pp. with dilute HNO₃ (Weller, B. 16, 27). ZnCl₂ forms C₁₅H₁₅N [102°].—B'2H2PtCl6: tables.—B'MeI. Crystals.

-B'McOH. Strongly alkaline.
PHENYL - METHYL - AMIDO - BENZENE NPhMe.C.H.P(OH) PHOSPHINIC ACID [150.5°]. Formed by the action of NaOHAq on the oily chloride NPhMe.C.H.PCl2, which is formed by the action of AlCl, on a mixture of NPh,Me and PCl, (Michaelis, A. 260, 37). Small needles (from water) or plates (from alcohol).—NaHA" 2aq. [265°]. Plates, v. e. sol. water. PHENYL-METHYL-AMIDO-BENZOIC ACID C_{1,1}H₁₃NO₂ i.e. C₆H₄(NPhMe).CO₂H. [184°]. Formed from NPh₂Me and COCl₂ (Michler a. Sarauw, B. 14, 2180). Plates.—BaA'₂: pearly plates.—AgA': white pp.

PHENYL-METHYL-AMIDO-ETHANE SUL-PHONIC ACID NPhMe.C₂H₄,SO₃H. Formed from CH₂Cl.CH₂.SO₃H and methylaniline at 160° (James, *J. pr.* [2] 31, 417). Silky crystals (from alcohol).

PHENYL - METHYL - AMIDO - ETHYL - PHTHALIMIDE C₈H₄O₂:N.C₂H₄.NPhMe. [105°]. Formed from bromo-ethyl-phthalimide and methyl-aniline at 165° (Newman, B. 24, 2199). Greenish-yellow four-sided tables, v. e. sol. C₆H₆.

PHENYL - METHYL - AMIDO-METHENYL-AMIDO-PHENOL C₁₄H₁₂N₂O *i.e.*

 $C_6H_4 < \stackrel{N}{\bigcirc} C.NPhMe$. (above 360°). Formed from $C_6H_4 < \stackrel{N}{\bigcirc} C.SH$ and methyl-aniline (Kalckhoff, B. 16, 1825). Syrup, with blue fluorescence.— $B'_2H_2PtCl_6$: prisms.

PHENYL - METHYL - AMIDO-DI-METHYL-PYBROLE C₁₈H₁₈N₂ i.e. NPhMe.N CMe:CH: (41°). (310°). Formed by heating its dicarboxylic acid at 231° (Knorr, A. 236, 310). Crystalline mass, v. sol. alcohol. Volatile with steam.

Phenyl-methyl-amido-di-methyl-pyrrole dicarboxylic acid C₁₃H₁₆N₂O₄. Formed by saponification of its oily ether, which is formed from diacetyl-succinic ether and phenyl-methyl-hydrazine (K.). Prisms, decomposing at 231°.—AgA': flocculent pp.

PHENYL - METHYL - p - AMIDO - PHENOL. Methyl derivative NPhMe.C₆H₄.OMe. (313°). Formed from phenyl-p-amido-phenol, KOH, and MeI (Philip a. Calm, B. 17, 2433). Oil, insol. NaOHAq.

PHENYL - TETRA - METHYL - TRI-AMIDO-DI-PHENYL-NAPHTHYL-CARBINOL

 $\begin{array}{lll} \textbf{C}_{33}\textbf{H}_{33}\textbf{N}_{3}\textbf{O} & \textbf{i.e.} & \text{NPhH.C}_{10}\textbf{H}_{4}.\textbf{C}(\textbf{OH})(\textbf{C}_{6}\textbf{H}_{4}.\textbf{NMe}_{2})_{2}.\\ \textbf{[95°].} & \textbf{Base of Victoria Blue B.} & \textbf{Formed by heating tetra-methyl-di-amido-benzophenone}\\ \textbf{(10 pts.)} & \text{with phenyl-}(a)-\text{naphthylamine (9 pts.),}\\ \textbf{and POCl}_{8} & \textbf{(7 pts.)} & \text{at } 110^{\circ} & \textbf{(Nathansohn a. Müller, B. 22, 1890).} & \textbf{The base is set free by NaOHAq.} & \textbf{Brick-red powder, v. sol. alcohol and benzene.} & \textbf{(C}_{33}\textbf{H}_{32}\textbf{N}_{3}\textbf{Cl})_{2}\textbf{PtCl}_{4}: & \text{violet needles.} \\ & \textbf{-C}_{33}\textbf{H}_{31}\textbf{N}_{3}\textbf{C}_{8}\textbf{H}_{2}(\textbf{NO}_{2})_{3}\textbf{OH}: & \text{dark-blue pp.} \end{array}$

Phenyl - penta - methyl-tri-amido-di-phenyl-naphthyl-carbinol $C_{34}H_{35}N_3O$ i.e.

NPhMe. $C_{10}H_0$. $C(OH)(C_0H_1$. $NMe_2)_2$. [77°]. The chloride C_3 , $H_{34}N_3$ Cl, 'Victoria Blue 4 R,' formed by the action of methyl-phenyl- (α) -naphthyl-amine on $CO(C_0H_1NMe_2)_2$ in presence of POCl₃ is v. sol. hot water (Nathansohn a. Müller, B. 22, 1892). The carbinol is a brick red pp., v. sol. alcohol. It forms $(C_3,H_{34}N_3C_1)_2$ PtCl₄ and $C_3,H_{32}N_3C_3H_2(NO_2)_3$ OH crystallising in very small dark-blue needles.

PHENYL TETRA METHYL TRI-AMIDODI-PHENYL NAPHTHYL METHANE $C_{32}H_{23}N_3$ i.e. NHPh. $C_{10}H_a$. CH(C_6H_4 .NMe $_2$)₂. [125°]. Formed by reducing the carbinol (v. supra) (Nathansohn a. Müller, B. 22, 1891). Flocculent pp., v. sol. hot alcohol. Yields B''_2H_2 PtCl $_4$ and $B'''C_6H_7N_4O_7$, both being crystalline,

Phenyl-penta-methyl-tri-amide-di-phenyl-naphthyl-methane $C_{34}H_{35}N_3$ i.e. NPhMe. $C_{16}H_{6}$.CH($C_6H_{10}Me_2$). [87°]. Formed by reducing the carbinol (N. a. M.). Amorphous. Vields crystalline B". H. PtCl. and B". C. H. N.O.

Yields crystalline B"',H₂PtCl₈ and B"'C₈H₃N₃O₃.

PHENYL DI - METHYL - AMIDO - TOLYL

KETONE C₈H₃.CO.C₈H₃Me.NMe₂. [67°]. (350°360°). Formed from benzoic acid, di-methyl-qtoluidine, and P₂O₃ (O. Fischer, A. 206, 91).

PHENYL-TETRA - METHYL-DI-AMIDO-DI-

TOLYL-METHANE $C_{22}H_{36}N_2$ i.e. CHPh $(C_8H_3Me.NMe.)_2$. [109°]. Formed by heating di-methyl-m-toluidine with benzoic aldehyde and $ZnCl_2$ (Fischer, B. 13, 807). Prisms.

-B"H₂PtCl₆ 2aq: small yellow crystals.
PHENYL-METHYL-AMINE v. METHYL-ANIL-

Phenyl-di-methyl-amine v. DI-METHYL-ANIL-

Di-phenyl-methyl-amine v. Methyl-di-phenyl-amine.

Tri-phenyl-methyl-amine v. Tri-PHENYL CARBINYLAMINE.

PHENYL-METHYL-ANTHRACENE C₂₁H₁₈
i.e. C₈H₃Me<CPh C₆H₄. [119°]. Obtained
by distilling phenyl-methyl-anthranol with zincdust (Hemilian, B. 16, 2367). Yellow pointed
crystals. Dissolves in alcohol and ether, with a
strong greenish-blue fluorescence. By CrO₃ in
acetic acid it is oxidised to phenyl-methyloxanthranol.

PHENYL-METHYL-ANTHRANOL $C_{2l}H_{16}O$, i.e. $C_{6}H_{3}Me < C_{(OH)}C_{6}H_{4}$. [157°]. Obtained by heating di-phenyl-tolyl-methane carboxylic acid with $H_{2}SO_{4}$ (Hemilian, B. 16, 2365). Yellow glistening tables. Sol. ether, boiling alcohol and acetic acid, insol. cold alkalis, but dissolves on boiling to a yellow solution. On oxidation it gives phenyl-methyl-oxanthranol.

PHENYL-DI-METHYL-ARŠINE v. vol. i. p. 320.

DI-PHENYL-DI-METHYL-AZIMETHYL-ENE N₂(CMePh)₂. [121°]. Formed by the action of acetophenone on hydrazine hydrate or on CPhMe:N.NH₂ (Curtius, J. pr. [2] 44, 542).

PHENYL-METHYL-BIAZOLE DIHYDRIDE OCH₂. NPh [140°]. Formed by the action

of tin and HCl on O CCl₂.NPh (122°), which is made by the action of COCl₂ on acetyl-phenylhydrazine (Freund a. Kuh, B. 23, 2838). Yellowish plates.

PHÉNYL-METHYL-TRIAZOLE C_pH_pN₃ i.e. N.NPh CH. Formed by heating its carboxylic acid to 180° (Bladin, B. 19, 2602). Oil. B'₂H_pPtCl₆ aq. [124°]. Lemon-yellow tables. PHENYL-METHYL-TRIAZOLE CARBOXY-

LIC ACID N.NPh C.CO₂H. [177°]. Formed by saponifying its nitrile, which is got by the action of Ac₂O on phenyl-hydrazine dicyanide and also by gently warming phenyl-hydrazine dicyanide with an alcoholic solution of pyruvic acid (Bladin, B. 18, 1544; 19, 2598; 22, 1749; 25, 185).

Salts. — CuA', 1½aq. — PbA', 2½aq. — AgA' 1½aq. — HA'HCl: tables (from HClAq).

Ethyl ether EtA'. Oil.

Nitrile. [109°]. Prisms. Amide C₂N₈PhMe.CO.NH₂. [170°]. Prisms,

m. sol. water and alcohol. Amidoxim C2N2PhMe.C(NH2):NOH. [210°]. Formed from the nitrile and hydroxylamine in alcohol. Plates (from alcohol), v. sl. sol. water. FeCl₃ colours its alcoholic solution deep red. Yields B'HCl and B'2H2PtCl₆ [201°]. C₂N₃PhMe.C(NH₂):NOAc [148°] and C_2N_3 PhMe.C < N > CMe [105.5°]. BzCl forms

DI-PHENYL-DI-METHYL-TETRAZONE

NPhMe.N:N.NPhMe. [137°]. Formed by oxidising phenyl-methyl-hydrazine (Fischer, A. 190, 152; Tafel, B. 18, 1744). Monoclinic plates. With I in CHCl₃ it forms C₁₄H₁₆N₁I₄, a black crystalline pp. DI-PHENYL-DI-METHYL-DI-TRIAZYL

N.NPh $C.C \leqslant NPh.N$ C.Me.N[223°]. Formed by heating the cyanide of phenyl-hydrazine with Ac₂O (Bladin, B. 21, 3063). Prisms (from alcohol).—B"H₂Cl₂.—B"H₂PtCl₃ aq. Plates.
DI-PHENYL-METHYL-BENZAMIDINE

CaHs.C(NMe).NPh2. Formed by heating diphenyl-benzamidine with MeI at 130° (Bernthsen, A. 192, 17). Syrup.—B'₂H₂PtCl₆: yellow pp. PHENYL p-METHYL-BENZYL KETONE

C₆H₄.CO.CH₂.C₆H₄Me. [94°]. Formed from ptolyl-acetic chloride, benzene, and AlCl₂ (Strassmann, B. 22, 1231). Yields an oxim [109°]. PHENYL-METHYL-BENZYL-UREA

NHPh.CO.NH.CH₂C₆H₄Me. [131°]. from m-methyl-benzylamine and phenyl cyanate (Brömme, B. 21, 2700).

TRI-PHENYL-METHYL-BROMIDE Bromo-tri-phenyl-methane.

PHENYL-METHYL-CARBAMIC ACID.

(244°). Ethyl ether NPhMe.CO.OEt. Formed from methylaniline and Cl.CO₂Et (Geb-

hardt, B. 17, 3042). Yellow oil.

Chloride NPhMe.COCl. [88°]. Got by passing COCl2 into a solution of methylaniline in benzene (Michler a. Zimmermann, B. 12, 1165). Tables (from alcohol), insol. water. Reacts with toluene and aluminium chloride, forming NPhMe.CO.C₆H₄Me [70°] while AlCl₂ and o-, m-, and p- xylene give rise to the compounds NPhMe.CO.C, H, Me, melting at 78°, 54°,

and 74° respectively (Lellmann, B. 24, 2114).

Phenyl ether NPhMe.CO.OC₈H₃. [58°]. Formed from the chloride and KOPh in alcohol (Lellmann a. Benz, A. 24, 2108). Large crystals.

Nitro-phenyl ethers C.H.(NO2)A'. The o-, m-, and p- nitro-phenyl ethers melt at 110°, 105°, and 70° respectively. They yield on reduction the corresponding amido-phenyl ethers melting at 103°, 94°, and 104° respectively. p-Tolyl ether C₆H₄MeA'. [62°].

PHENYL-METHYL-SEMI-CARBAZIDE

NPhMe.NH.CO.NH₂. [133°]. Formed from phenyl-methyl-hydrazine hydrochloride potassium cyanate (Fischer, A. 190, 164). Crystalline mass, m. sol. hot water. Yields a nitrosamine NPhMe.N(NO).CO.NH2 [77°].

B. 13, 2172). Pyramids. Decomposed by HCl into the parent substances.

Di-phenyl-di-methyl-carbazide

CS(NH.NPhMe)₂. [c. 168°]. Got by heating phenyl-methyl-hydrazine with CS₂ (Stahel, A. 258, 250).

PHENYL-METHYL-CYANIDE v. Nitrile of PHENYL-ACETIC ACID.

Di-phenyl-methyl-cyanide v. Nitrile of Di-PHENYL-ACETIC ACID.

Di-phenyl-methyl dicyanide CieHiaN, i.e. $N \stackrel{CMe.N}{\stackrel{CPh.N}{\stackrel{}}} CPh.$ [110°]. (227° at 15 mm.).

Formed from benzonitrile, AcCl, and AlCl, (Krafft a. Hansen, B. 22, 803; 23, 2382). Needles, sol. alcohol and ligroin. Decomposed by diluted H2SO, into benzoic soid, acetic scid, and NH₂. Oxidised by alkaline KMnO₄ to C₃N₃Ph₂CO₂H [c. 192°] which on heating yields C₃N₂Ph₂H [75°] (205° at 9 mm.).

Salts.—B'HCl: needles.—B'2H2PtCl6

DI - PHENYL - METHYLENE - p - AMIDO-BENZOIC ACID CPh₂:N.C₆H₄,CO₂H. [240°]. Formed from Ph₂CCl₂ and p-amido benzoic acid (Hantzsch a. Krafft, B. 24, 3522). Short prisms. PHENYL-METHYLENE-AMINE CH,:NPh

or CH₂<NPh>CH₂? [140°]. Formed by adding aniline to a solution of formic aldehyde (Wellington a. Tollens, B. 18, 3309). Formed also from trioxymethylene and aniline (Kolotoff, J. 1885, 777). Long slender needles. composed by water at 100° into trioxymethylene and aniline.

Phenyl-trimethylene-diamine

NHPh.C₃H₆,NH₂. (282° cor.). S.G. § 1.0356. A product of the action of Na on an alcoholic solution of phenyl-pyrazole (Balbiano, G. 18, 354; 19, 688; Rend. Accad. Linc. [4] 4, ii. 44). Formed also by boiling phenyl-y-amido-propyl-phthalimide with HClAq (Goldenring, B. 23, 1169). Oil. Sulphide of carbon forms NHPh.C3H6.NH.CS.SNH2Ph.C3H6NH2, orystallising from alcohol in white plates. -B"2HCl: tufts of needles, v. e. sol. water. When heated it gives off NH₃ and NPh:C₃H₆, which forms (C₃H₁₁N)₂H₂PtCl₆.—C₃H₁₁N,2C₃H₃N₃O₅. Greenish crystals, decomposing at 195°.—Succinate [100°].

Phenyl-di-trimethylene-triamine

NPh(CH₂.CH₂.CH₂.NH₂)₂. Formed by heating its phthalyl derivative with HClAq at 240° (Goldenring, B. 23, 1170). — B"3HCl. — B₂"3H₂PtCl₈.
Di-phthalyl derivative

 ${
m NPh}({
m CH}_2{
m CH}_2{
m CH}_2{
m N}:{
m C}_4{
m H}_1{
m O}_2)_2$ [145°]. Formed from γ -bromo-propyl-phthalimide and aniline. Di-phenyl-methylene-amine CPh $_2$:NH.

Formed by heating CPh2Cl2 with carbamic ether at 130° (Hantzsch a. Krafft, B. 24, 3517). Colourless liquid.—B'HCl: snow-white powder.

Di-phenyl-methylene-diamine CH2(NHPh)2. [49°]. Formed by mixing a large excess of an aqueous solution of aniline with formic aldehyde solution (Pratesi, G. 14, 351; cf. Lermontoff. B 7, 1255). Four-sided tables, sol. alcohol. Its hydrochloride is unstable.—B"H,PtCl.

Di-phenyl-trimethylene-diamine Phenyl-di-methyl-semi carbazide

NMe_x.NH.CO.NHPh. [108°]. Formed from trimethylene bromide (Hanssen, B. 20, 781).

Phenyl-di-methyl-semi carbazide

CH₂(CH₂.NHPh)₂. Formed from aniline and trimethylene bromide (Hanssen, B. 20, 781).

Thick brown liquid.—B"H₂SO₄. [156°]. Needles. Nitrosamine CH2(CH2.NPh.NO)2. DI-PHENYL-METHYLENE-ANILINE

NPh:CPh₂. [109°]. (above 360°). Formed from CPh₂Cl₂ and aniline (Pauly, A. 187, 199). Yellow tables (from ether). Split up by HClAq into aniline and benzophenone.

TETRA - PHENYL - DI - METHYLENE -**TETRAZONE** $N_2(N:CPh_2)_x$ Formed by shaking a benzene solution of $NH_2.N:CPh_2$ with HgO (Curtius a. Rauterberg, J. pr. [2] 44, 200). Red Decomposes readily into nitrogen and CPh2:N.N:CPh2. Gaseous HBr forms CPh2HBr

DI-PHENYL-TRIMETHYLENE-DI-CARB-AMIC ACID.

Ethyl ether CH2(CH2.NPh.CO2Et)2 [56°]. Formed from the chloride and NaOEt (Hanssen,

B. 20, 783). Crystalline. Chloride CH₂(CH₂NPh.COCl)₂. [102°]. Formed from CH₂(CH₂NPhH)₂ and COCl₂ in

benzene. Needles.

PHENYL-TRIMETHYLENE DICARBOXYLIC

ACID CHPh CH.CO.H. [175°]. Got by saponifying its ether. Prisms, m. sol. water, v. sol. alcohol and ether (Buchner a. Dessauer, B. 21, 2646; 25, 1152). When heated in vacuo it yields an anhydride C₁₁H₈O₈ [134°]. —NaHA": needles.

Di-methyl ether Me,A". [63°]. Ethyl ether Et,A" (257° at 120 mm.). Formed by distilling CO₂Et.CH<N.CHPh N.CH.CO₂Et

in vacuo. Thick oil.

Phenyl-trimethylene-tri-carboxylic acid C₃H₂Ph(CO₃H)₃. [188°]. Got by saponifying its ether (B. a. D.). Crystals containing 4aq.

Methyl ether Me₃A'''. [47°]. (210° at

20 mm.). Formed from NaOMe, methyl malonate, and methyl di-bromo-phenyl-propionate.

PHENYL METHYLENE ETHYL DIKETONE C.H. CO.CH. CO.C.H. Propionyl-acetophenone. (277°). S.G. 15 1.081. Formed from acetophenone, propionic ether, and NaOEt (Stylos, B. 20, 2181). Liquid.

DI-PHENYL-METHYLENE-HYDRAZINE [98°]. (c. 227° at 55 mm.). CPh,:N.NH, Hydrazide of benzophenone. Formed by heating benzophenone (\bar{b} g.) with hydrazine hydrate (1.8 g.) and alcohol (1 g.) at 150° (Curtius a. Lauterberg, J. pr. [2] 44, 192). Prisms (from alcohol). Reduces alcoholic AgNO, forming a mirror. Reduces Fehling's solution on warming. Boiling dilute H2SO4 splits off hydrazine.

Reacts readily with ketones.—B'HCl. [183°].

Acetyl derivative. [107°]. Prisms.

Benzoyl derivative. [116.5°]. Prisms. Tetra-phenyl-di-methylene-hydrazine

CPh2:N.N:CPh2. Di-phenyl-ketazine. [162°]. Formed by boiling CPh2:N.NH2 with iodine in alcohol. Yellow prisms, v. sl. sol. hot water, sol. alkalis. Decomposed by boiling dilute H_2SO_4 into benzophenone and $N_2H_4H_2SO_4$.

Phenyl - trimethylene - hydrazine C,H12N2 [53°]. Got from phenyl-hydrazine and trimethylene b.omide (Marckwald, C. C. 1888, 1410). Converted by trimethylene bromide on further heating into (PhN₂)₂(C₃H₆)₃ [226°].

DI-PHENYL METHYLENE DIKETONE $CH_2(CO_*C_6H_5)_2$. Benzoyl-acetophenone. Dibenzoyl methans. [81°]. (over 200°).

Formation.—1. By boiling di-benzoyl-acetic acid with water (Baeyer a. Perkin, jun., B. 16, 2134; C. J. 47, 240).—2. By the action of malonyl chloride and AlCl, upon benzene (Auger,

A. Ch. [6] 22, 348).

Preparation.—By adding dry sodium ethylate to a mixture of acetophenone and benzoic ether, the reaction taking place with spontaneous heating. The crystalline mass is extracted with cold water, finally with dilute NaOH, and from the solution of the sodium salt the ketone is precipitated by means of CO_2 ; the yield is c. 50 p.c. of the acetophenone used (Claisen, $B.\ 20,\ 655$).

Properties.—Trimetric plates, v. sol. alkalis. Yields products of substitution by treatment

with NaOEt and alkyl iodides.

Reactions.—1. Bromine (1 mol.) in CHCl. forms Bz₂CHBr [93°], which on heating with HOAc and KOAc yields Bz₂CHOAc [94°], which on further bromination gives Bz₂CBr.OAc [102°] which is split up by heat into Ph.CO.CO.CO.Ph and AcBr.-2. Excess of bromine (2 mols.) in CHCl, forms Bz, CBr, [95°] whence HOAc and KOAc form di-phenyl triketone.—3. Amyl nitrite and alcoholic HCl give Bz₂C:NOH [146°], converted by nitrous acid into the triketone (Neufville a. Pechmann, B. 22, 852; 23, 3376).—4. Phenyl-hydrazine forms tri-phenyl-pyrazole (Japp a. Klingemann, B. 22, 2886)

METHYL DI-PHENYL METHYLENE KETONE v. BENZOYL-ACETONE.

PHENYL METHYLENE METHYL SUL-PHONE-KETONE C.H. SO. CH. CO.CH. Acetonyl phenyl sulphone. [57°]. Formed by oxidising PhS.CH.Ac with KMnO₄ (Delisle, A. 260, 262). Formed also from chloro-acetone and C₂H₃.SO₂Na (Otto, B. 19, 1642; 21, 1871; 23, 756; J. pr. [2] 36, 402). Plates, sol. water and alcohol. Yields a phenyl-hydrazide [129°] and an oxim [148°]. Ammonia yields the compound NH:CMe.CH.,SO,Ph [111°]. Phenyl mercaptan yields (PhS),CMe.CH,SO,Ph [104]. Potash yields phenyl methyl sulphone. Br forms a bromo-derivative [96°] and a di-bromo-derivative [114°]

PHENYL-METHYLENE-PHTHALIMIDINE $C_{15}H_{11}NO$ i.e. $C_{6}H_{4} < CO > NPh$. Formed by heating acetophenone o-carboxylic anilide at 230° (Mertens, B. 19, 2372). Prisms, v. sol. alcohol. An isomeride [265°], got by

treating acetophenone o-carboxylic anilide with conc. H₂SO₄, is sl. sol. alcohol.

DI-PHENYL METHYLENE DISULPHONE CH₂(SO₂Ph)₂. [119°]. Formed by oxidation of CH2(SPh)2 which is got by boiling CH2Cl2 with NaSPh (Fromm, A. 253, 161). Needles, v. sl. sol. boiling water.

DI-PHENYL-DI-METHYLENE DI-SUL. PHONE KETONE C15H14S2O5i.e. CO(CH2SO2Ph)2. Got from CHBr.CO.CH, SO, Ph and C.H.SO.Na. Got also from CO(CHCl2)2 and C.H., SO, Na (Otto, B. 19, 1542; 22, 1967). Small crystals (from HOAc). Yields an oxim [137°] and a phenyl-hydrazide [171°]. PhSH forms (PhS),C(CH,SO,Ph), [191°]. DI - PHENYL-METHYLENE-DI-THIO-GLY-

COLLIC ACID CPh2(S.CH2.CO2H)2. [176°]. Formed by heating benzophenone with thioglycollic acid and ZnCl₂ (Bongartz, B. 21, 483). Needles (from dilute HOAc), insol. hot water.

PHENYL-TRIMETHYLENE-THIO-UREA

CS NH.CH. CH. [215°]. Formed by boiling NHPh.C₃H₆.NH.CS.SNPhH₂.C₂H₆.NH. with water (Goldenring, B. 23, 1172). White prisms.

Di-phenyl-methylene- ψ -thio-urea $C_{1_1}H_{12}N_2S$ i.e. S < C(NPh) NPh. [68°]. Formed from di-phenyl-thio-urea and CH_2I_2 (Foerster, B. 21, 1872). Thin needles, v. e. sol. alcohol.— $B'_2H_2PtCl_4$. Crystalline pp.

Di-phenyl-trimethylene-ψ-thio-urea

S<C(NPh)-NPh CH₂.CH₂.CH₂. [123°]. Formed from diphenyl-thio-urea and trimethylene bromide (F.).

Di-phenyl-trimethylene-di-thio-di-urea CH₂(CH₂.NH.CS.NPhH)₂. Formed from phenylthiocarbimide, trimethylene-diamine, and alcohol (Lellmann a. Würthner, A. 228, 236). Plates [115°] or nodules [60°], v. sol. alcohol.

Tri-phenyl-trimethylene-di-thio-di-urea NHPh.CS.NH.CH₂.CH₂.CH₂.NPh.CS.NHPh. [145°]. Formed by heating phenyl-trimethylenediamine (1 g.) with phenyl-thiocarbimide (2 g.) and alcohol (15 c.c.) (G.). Slender needles.

DIPHENYL-METHYLENE-p-TOLUIDINE C₂H₄MeN:CPh₂. (above 360°). Formed from Ph₂CCl₂ and p-toluidine (Pauly, A. 187, 214). Oil, split up by acids into p-toluidine and benzophenone.

PHENYL-TRIMETHYLENE-UREA

CO NPh.CH₂ CH₂ [215°]. Formed by heating phenyl-amido-propyl-urea at 240° (Goldenring, B. 23, 1173). Yellowish plates, sl. sol. hot Aq. Di-phenyl-trimethylene-urea

CO NPh.CH₂ CH₂. [156°]. Formed, together with CH₂(CH₂NPh.COCl)₂ [102°], from di-phenyl-trimethylene-diamine and COCl₂ in benzene (Hanssen, B. 20, 782). Plates.

Tetra-phenyl-trimethylene-di-urea
CH₂(CH₂:NPh.CO.NHPh)₂. [153°]. Formed
from CH₂(CH₂:NPh.COCl)₂ and aniline (H.).
PHENYL TRIMETHYLENYL KETONE

 C_8H_3 -CO.CH $\stackrel{CH_2}{CH_2}$ or $\stackrel{CH:CPh}{CH_2}$ O. Benzoyltrimethylenc. Phenyl-furfurane dihydride (239°) at 720 mm. V.D. 5·27 (calc. 5·04). Formed by heating its carboxylic acid at 200° (Perkin, jun., B. 16, 2140; C. J. 47, 842). Oil with aromatic odour. Alcoholic hydroxylamine forms the oxim in the cold, but by heating the ketone with alcoholic hydroxylamine hydrochloride at 135° there are formed $C_{33}H_{27}N_3O_3$ (a red amorphous powder) and $C_{36}H_{28}N_2O_2$ (a black brittle mass which forms deep-blue B" H_2Cl_2 and B" H_2PtCl_4).

Oxim. [87°]. Colourless plates.

PHENYL TRIMETHYLEÑYL KETONE CARBOXYLIC ACID $\overset{\text{CH}_2}{\text{CH}_2}$ CBz.CO₂H. [149°]. Formed by saponification of the ether. Monoclinic prisms; $a:b:c=2\cdot330:1:2\cdot132$; $\beta=83^\circ4'$. M. sol. ether, sl. sol. water. Decomposed by water at 110° into phenyl oxypropyl ketone CH₂OH.CH₂.CH₂Bz, which when extracted with

M. sol. ether, sl. sol. water. Decomposed by water at 110° into phenyl oxypropyl ketone CH₂OH.CH₂CH₂Bz, which when extracted with ether and dried over K₂CO₃ is found to have split off water, the resulting anhydride being possibly phenyl trimethylenyl ketone (Perkin, C. J. 51, 837).—AgA': flocculent pp.

Ethyl ether Et.A". (282°) at 720 mm. Formed from benzoyl-acetic ether, ethylene bromide, and NaOEt (Perkin, B. 16, 2138; C. J. 47, 841). Oil.

Reference.—Nitro - Phenyl - Trimethylenyl

KETONE CARBOXYLIC ACID.

PHENYL-METHYL-ETHYLENE-DIAMINE NH₂.CH₂.CH₂.NPhMe. (255°). Formed by the action of HCl on C₆H₄:C₂O₂:N.C₂H₄.NPhMe (Newman, B. 24, 2200). Liquid, v. sol. water and alcohol.—B'C₆H₂(NO₂)₃OH. [173°]. Needles-

Di-phenyl-di-methyl-ethylene-diamine C₂H_{*}(NPhMe)₂. The methylo-bromide B"Me₂Br₂, got by heating di-methyl-aniline with ethylene bromide (Hübner, A. 224, 346), crystallises in deliquescent prisms and yields B"Me₂(OH)₂, B"Me₂Cl₂, B"Me₂Cl₂, B"Me₂Cl₂, B"Me₂Cl₂, B"Me₂Cl₂, B"Me₂Cl₂, B"Me₂Cl₂, B"Me₂Cl₂O₇, and B"Me₂(O₂H₂N₃O₇)₂ [124°].

PHENYL-METHYL-ETHYLENE DIKE-TONE v. ACETOPHENONE-ACETONE. Its carboxylic acid is described as ACETOPHENONE-ACETO-

ACETIC ACID.

DI-PHENYL-DI-METHYL-v-ETHYLENE-DI-PYRROLE-DI-CARBOXYLIC ACID

CO₂H.Ç:CMe N.C₂H₄.N CMe:Ç.CO₂H. [181°]. Obtained by saponification of its ether, which is formed by mixing acetophenone-aceto-acetic ether with ethylene-diamine (Paal a. Schneider, B. 19, 8158). Crystalline solid. Sl. sol. alcohol, acetic acid, and conc. HCl, insol. other ordinary solvents. On heating above 180° it evolves CO₂, giving the free base.

Di-ethyl-ether Et₂A": [197°]; plates; distillable under ordinary pressure; sol. alcohol, benzene, chloroform, and acetic acid, insol.

water and petroleum-spirit.

PHENYL-METHYL-ETHYL-0SOTRIAZOLE

NPh N:CMe (270°). Formed by condensation of the oxim-phenyl-hydrazide of CH₂.CO.CO.C₂H₃ (Pechmann, A. 262, 312).

Needles. Yields a di-nitro-derivative [113°].

Oxidised by alkaline KMnO₄ to phenyl-methyl-triazole carboxylic acid.

PHENYL - METHYL - ETHYL - PYRAZOLE

NPh \(\bigcap \text{CEt:CMe} \) (283°). S.G. \(\frac{16}{2} \) 1.0476. Got

by the action of phenyl-hydrazine on

C2H3.CO.CHMc.CHO (Claisen, B. 22, 3276).

Reference.—OXY - PHENYL - METHYL - ETHYL -

PYRAZOLE.

TRI - PHENYL - METHYL - ETHYL - THIO-BIURET C₂S₂N₃Ph₃MeEt. [157°]. Formed from di-phenyl-ethyl-thio-urea and NPhMe.CSCl (Billeter a. Strohl, B. 21, 108). An isomeride [156°] is formed from di-phenyl-methyl-urea and NPhEt.CSCl. Both bodies crystallise in needles, sl. sol. alcohol.

PHENYL - METHYL - ETHYL - THIO - UREA CS(NHMe).NPhEt. Got from methyl thiocarbimide and ethyl-aniline (Gebhardt, B. 17,

3037). Crystalline solid.

Di-phenyl-methyl-ethyl-thio-ure? CS(NPhMe).NPhEt. [49·5°]. Formed from methyl-aniline and NPhEt.COCl (Billeter, B. 20, 1682). Monoclinic crystals.

PHENYL-METHYL-FURFURANE C₁₁H₁₀O, CH:CMe Or Oxyphenyl-pentamethenyl

hydride CH₂·CH₂ CO. [40°]. (c. 238°). Formed by heating phenythronic acid, and prepared by warming acetophenone-acetone with fuming HClAq, the yield being 65 p.c. (Paal, B. 17, 2759; Fittig, B. 21, 2134; A. 250, 222). Needles or prisms, insol. water and alkalis, volatile with steam. Br forms C₁₁H_aBr_sO [210°]. Phenylhydrazine forms a compound [165°–175°], crystallising from alcohol in golden spangles.

tallising from alcohol in golden spangles.

Tetrahydride CH. CHMe O. (c. 230°).

Formed by reduction in alcoholic solution by Na. Oil, miscible with alcohol and ether.

PHENYL-METHYL-FURFURANE CARB-OXYLIC ACID C₁₂H₁₀O₃ i.e. CO₂H.C:CMe HC:CPh>O.

[181°]. Formed by boiling dehydro-acetophenone-acetoacetic ether (cf. vol. i. p. 36) with HClAq (Paal, B. 17, 2762). Needles. Boiling Ac₂O forms an acetyl derivative C₁₄H₁₂O₄ [83°], reconverted by NaOHAq into the parent acid.—KA'.—AgA': crystalline pp.

Phenyl-methyl-furfurane carboxylic acid

C₁₂H₁₀O₃ i.e. CO₂H.C=CPh O or

CO₂H.CH.CPh CH. Phenuvic acid. [145°]. Formed by heating phenythronic acid (Fittig, B. 21, 2184). Its ether is obtained from chloroacetone and sodium benzoyl-acetic ether (Coleax, C. J. 59, 190). Long silky needles.—CaA'₂ 2aq.—BaA'₂aq.—AgA': amorphous pp.

Phenyl-methyl-furfurane dicarboxylic acid $C_{12}H_{10}O_3$ i.e. $CO_2H.CH.CPh$ CCO_2H or $CH_2.CO$

CO₂H.C:CMe O. Phenythronic acid [193°]. Formed by saponifying its ether with boiling baryta (Fittig, A.250, 216). Silky needles, m. sol. hot water, v. sol. alcohol.—CaA" 3aq. S. 5.—BaA" aq.—Ag₂A": crystalline pp.

Mono-ethyl ether EtHA". [112°]. Formed by the action of Ac₂O on a mixture of sodium succinate and benzoyl-acetic ether.— Ca(EtA")₂: crystalline nodules (from alcohol).—Ba(EtA")₂.—AgEtA": white amorphous pp.

Di-ethyl ether EtA". [44.5°].

Di-ethyl ether Et,A". [44.5°]. PHENYL-METHYL-GLYCOCOLL v. PHENYL-METHYL-AMIDO-AGETIC ACID.

PHENYL-METHYL-GLYOXALINE $C_{10}H_{10}N_2$ i.e. CMe < NH.CPh. [159°]. Formed by heating phenyl-methyl-oxazole with alcoholic NH_3 at 230° (Lewy, B. 22, 2195). Needles, v. sl. sol. alcohol. Salts.—B'HCl. [242°]. Needles.—B'_2H_2PtCl_2aq. [212°]. Yellow prisms.

Phenyl-di-methyl-glyoxaline $C_{11}H_{12}N_2$ i.e. CPh < N-CMe [230°-234°]. Formed by heating diacetyl with benzoic aldehyde and alcoholic NH_2 (Wadsworth, C. J. 57, 8). Needles (from benzene). $-B'_2H_2PtCl_3$: needles.

Di-phenyl-methyl-glyoxaline CMe $\sqrt{N-CPh}$. [235°]. Formed by the action of aldehyde and NH, on benzil (Japp, C. J. 49, 466; 51, 557; C. J. Proc. 3, 34). Trimetric crystals; a:b:c=1'207:1:1'929. Crystallises with diphenylglyoxaline (1 mol.), the compound melting at 218°.

PHENYL-METHYL-HYDANTOIC ACID C_cH_b.NH.CO.NH.CH(CH₃).CO_cH [170°]. Formed by the combination of phenyl cyanate with alanine and boiling the product with alcoholic KOH (Kühn, B. 17, 2884). Glistening scales.

PHENYL-METHYL-HYDANTOIN

CHPh CO. NMe [162°]. Formed from phenylhydantoin, KOMe, and MeI (Pinner, B. 21, 2325). Needles, sl. sol. cold alcohol.

s-PHENYL-METHYL-HYDRAZINE

C_eH_sNH.NHMe. Hydrazo - phenyl - methyl. Got by saponification of its dibenzoyl derivative, which is formed by heating NPhBz.NBzH with NaOMe and MeI (Tafel, B. 18, 1740). Obtained also by the action of alkalis on nitroso-antipyrin (Knorr, A. 238, 137). Oil, oxidised by HgO to PhN:NMe. Reduces Fehling's solution and ammoniacal AgNO₂ in the cold.—B'₂H₂SO₄. [180°].—Oxalate: white needles.

Di-benzoyl derivative PhNBz.NMeBz. [145°]. Crystals, v. sol. alcohol and ether.

u-Phenyl-methyl-hydrazine NPhMe.NH₂. (227° i.V.) at 745 mm. Formed by converting methyl-aniline into phenyl-methyl-nitrosamine, and reducing this body (5 pts.) by zinc-dust (20 pts.), HOAc (10 pts.), and water (35 pts.) at 0° (E. Fischer, A. 190, 152; 236, 199).

Properties.—Colourless oil, turning brown in air; m. sol. hot water, miscible with alcohol.

Reactions.—1. Nitrous acid yields NPhMe.NO, water, and N₂O.—2. Diazobenzene nitrate solu-tion forms PhN₂ and methyl-aniline nitrate.— 3. HgO added to its solution in chloroform produces NPhMc.N:N.NPhMe [137°].-4. Reacts with aldehydes and ketones in the same way as phenyl-hydrazine, forming products of condensation; thus benzoic aldehyde forms NPhMe.N:CHPh [104°] (Ebers, A. 227, 340), acetophenone forms NPhMe.N:CMePh [50°], acetone forms $C_{18}H_{11}N_{2}$ (216°) (Degen, A. 236, 152), propionic aldehyde yields $C_{17}H_{11}N_{2}$ (198° at 170 mm.), glucose yields $C_{6}H_{16}O_{5}$:N.NPhMe[171°], while benzil yields Ph.CO.CPh:N.PhMe [56°] and N₂PhMe:CPh.CPh:N₂PhMe[180°](Kohlrausch, A. 253, 16; Fischer, B.22,87).—5. CS₂ at 100° forms CS(NH.NPhMe)₂ [c. 168°] (Stahel, A.258,250).— 6. Nitroso-aniline forms $C_{18}H_{14}N_4O$ [151°], while nitroso-dimethylaniline gives $C_{15}H_{18}N_4O$ [141°] (O. Fischer, B. 22, 622).—7. With PhN:80 it forms PhNMe.N:SO [77°] crystallising from alcohol in yellow needles (Michaelis, A. 270, 120).

Salt.—B'₂H₂SO₄: plates, v. e. sol. water. Acetyl derivative NPhMe.NHAc. [93°]. Prisms, m. sol. hot water. Converted by Na and MeI into NPhMe.NMeAc [68°], which is not decomposed by alkalis, but gives methylaniline when boiled with HClAq.

Benzoyl derivative NPhMe.NHBz. [153°]. Formed by heating NPhH.NHBz with NaOMe and MeI. Needles (Tafel, B. 18, 1743).

Naume and Mel. Needles (Taiei, B. 18, 1743). Sulphonic acid $SO_3H.C_5H_1NMe.NH_2$. Formed by heating with fuming H_2SO_4 (Pfülf, A. 239, 219). Plates.—Na Λ' aq: slender needles.

PHENYL-METHYL-IMESATIN v. Anilide of Methyl Isatin.

DI-PHENYL METHYLIMIDO-DI-ETHYL DISULPHONE NMe(C₂H₄.SO₂Ph)₂. Formed by heating NH(C₂H₄.SO₂Ph) with alcohol and MeI at 115° (Otto, J. pr. [2] 80, 385).—B'HCl [221°]. PHENYL-METHYL-INDOLE C₁₈H₁₈N i.s. C₄H₄CH NMe CPh. [101°]. Formed by heating the phenyl-hydrazide of acetophenone with ZnCl₂ at 130° (Fischer, B. 19, 1565; Degen, A. 236, 155). Formed also by heating CH₂Bz.NMePh with ZnCl₂ at 180° (Staedel, B. 21, 2196; Culmann, B. 21, 2596), and by heating the isomeric C₂H₄CPh NMe CH with ZnCl₂ at 220° (Ince, A. 253, 40). Needles (from alcohol). Colours pinewood, moistened with HCl, dark-red.

Acetyl derivative C_eH₄ CAc CPh. [136°]. Formed by heating CH₂Ac.CPh:N.NMePh with ZnCl₂ at 150°. Needles (from dilute HOAc).

 $\begin{array}{lll} \textbf{Phenyl-methyl-indole} & \textbf{C}_{6}\textbf{H}_{4} \diagdown \begin{matrix} \textbf{CPh} \\ \textbf{NMe} \end{matrix} \\ \textbf{CH}. \\ \textbf{[65°]}. & \textbf{Formed from phenyl-acetic aldehyde and phenyl-methyl-hydrazine} & \textbf{(Ince, A. 253, 38)}. \\ \textbf{Crystals, v. sol. benzene.} & \textbf{Colours acidified pine-wood reddish-violet.} \\ \textbf{-B'C}_{6}\textbf{H}_{4}\textbf{N}_{3}\textbf{O}_{7}. & \textbf{[90°]}. \\ \textbf{Phenyl-methyl-indole} & \textbf{C}_{6}\textbf{H}_{4} \diagdown \begin{matrix} \textbf{CPh} \\ \textbf{NH} \end{matrix} \\ \textbf{CMe}. \\ \end{array}$

Phenyl-methyl-indole C₈H₄ NH CMe. [60°]. Formed by heating the phenyl-hydrazide of benzyl methyl ketone with ZnCl₂ (Trenkler, A. 248, 110). Prisms (from ligroïn).—Picrate: [142°]; v. sol. benzene.

PHENYL METHYL KETONE v. Aceto-PHENONE.

Phenyl methyl diketone C₆H₃.CO.CO.CH₃. Acetyl-benzoyl. (217°). S.G. ¹⁴/₄ 1·1041. S. ·26 at 20°. Formed by distilling its oxim Bz.C(NOH)Me with dilute H₂SO₄ (Von Pechmann, B. 21, 2119; 22, 2129) or with isoamyl nitrite (Manasse, B. 21, 2177). Heavy oil, volatile with steam.

Oxim C_aH_b.CO.C(NOH).CH₃. Nutroso-propiophenone. [114°]. Formed by the action of isoamyl nitrite and HCl or NaOEt on the ketone (Claisen a. Manasse, B. 22, 529). Prepared by the action of nitrous acid on benzoyl-propionic ether (1 mol.) and KOHAq (3 mols.) (P.; Gudeman, B. 22, 562). Needles (from hot water), forming a deep-yellow solution in alkalis. SnCl₂ and HCl yield hydroxylamine and phenyl methyl diketone. Reduced by sodium-amalgam to a base which crystallises from alcohol in needles [108°].

Di-oxim Ph.C(NOH).C(NOH).Me. [236°]. Formed from the mono-oxim and hydroxylamine hydrochloride. Needles (from dilute alcohol).

Phenyl hydrazide C₁₅H_{1,1}N₂O. [145°] Di-phenyl-di-hydrazide CMe(N₂HPh).CPh(N₂HPh). [105°]. Yellow

crystals, v. sol. ether.

Oxim-phenyl-hydrazide

CPh(N₂HPh).CMe:NOH. [202°]. White needles. Phenyl methyl triketone. Mono-oxim C₆H₈.CO.C(NOH).CO.CH₈. [124°]. Formed by the action of nitrous acid on phenyl methylene methyl diketone (Ceresole, B. 17, 814). Needles, v. sol. benzene, sol. alkalis.

Di-oxim C_eH₂.C(NOH).C(NOH).CO.CH₃. [178°]. Formed by neating the mono-oxim with hydroxylamine hydrochloride. Needles, insol.

cold water, sol. alkalis.

PHENYL-METHYL-KETONE CARBOXYLIC ACID v. Benzoyy-acetic acid and Acetophenone carboxylic acid.

PHENYL-METHYL-MALONIC ACID v. BEN-AYL-MALONIC ACID. Tri-phenyl-methyl-malonic ether

CPh₃.CH(CO₂Et)₂. [133·5°]. Formed from CPh₂Br and sodium malonic ether (Henderson, B. 20, 1014; C. J. Proc. 2, 251). Needles (from alcohol). Yields tri-phenyl-propionic acid when heated with alcoholic potash.—NaA'aq: needles.—BaA'₂ aq.—AgA': minute white crystals.

PHENYL - DI - METHYL - NAPHTHALENE TETRAHYDRIDE v. METHRONOL.

PHENYL - METHYL - (β)-NAPHTHOTRIAZ-INE DIHYDRIDE $C_{18}H_{15}N_3$ i.e. $C_{10}H_6 < \stackrel{N.CHMe}{N.NPh}$.

Formed by adding aldehyde to a mixture of benzene-azo-(\$\beta\)-naphthylamine and alcohol (Goldschmidt a. Poltzer, B. 24, 1004). Plates, v. sol. hot alcohol.—B'HCl. [252°].—B'_2H_2PtCl_. Yellow needles.—B'MeI. [244°].—B'_2Me_2PtCl_. [260°]. Yellow needles.

PHENYL - METHYL-(β)-NAPHTHYL-THIO-UREA C₁₈H₁₆N₂S i.e. NPhMe.CS.NHC₁₀H₁, [127°]. Formed from (β)-naphthyl thiocarbimide and methyl-aniline (Gebhardt, B. 17, 2091). Yellow needles.

PHENYL - METHYL - NITROSAMINE v_{\bullet} Nitrosamine of METHYL-ANILINE, vol. iii, p. 285. PHENYL-METHYL-OSOTRIAZOLE

NPh N:CH (242°). S.G. 7 1·1071. Formed by oxidising the diphenyl-hydrazide of pyruvic aldehyde and warming the resulting NPh.N:CH Formed also by the action of NaOH on CH₃·C(N₂HPh).CH:NOAc (Pechmann, B. 21, 2751; A. 262, 279). Oil. Yields a trinitro-derivative [138°], and a sulphonic acid, of which the Na salt crystallises in hexagonal plates.

Phenyl-di-methyl-osotriazole

NPh N:CMe N:CMe [35°]. (351°). Formed by heating the phenyl-hydrazide-oxim of diacetyl with Ac O (P.). Crystalline, insol. water, sol. conc. HClAq, but reppd. by water. Yields a dinitro-derivative [189°] and a tri-nitro-derivative [227°].

PHENYL-METHYL-OSOTRIAZOLE CARBOXYLIC ACID NPh<n:CMe
N:C.CO...H. [198°].
Formed by oxidising phenyl-di-methyl-osotriazole with KMnO4 (Pechmann, A. 262, 308).

Needles, v. sl. sol. water. – AgA': white pp. PHENYL-METHYL-OXAMIDE C₀H₁₀N₂O₂ i.e. NHPh.C.O₂.NHMe. [180°]. Formed from NHMe.C₂O₂.OEt and aniline (Wallach, A. 184, 70). Needles.

PHENYL-METHYL-OXANTHRANOL

C₂₁H_{1e}O₂ i.e. C₆H₃Me COPh(OH) C₆H₄. [195°]. Formed by oxidation of phenyl-methyl-anthranol with K₂Cr₂O₇ and HOAc (Hemilian, B. 16, 2366). Trimetric tables, v. sol. hot alcohol, insol. alkalis. Conc. H₂SO₄ forms a purple solution.

Isomeride [1:\frac{4}{5}] C_oH_1Me \(\begin{align*}\frac{\text{CPh}(\text{CH})}{\text{CO}} \end{align*} \cdot C_oH_4. \]

[213°]. Formed by oxidising di-phenyl-tolyl-methane o-carboxylic acid with K2CO4 and conc. H2SO4 (Hemilian, B. 19, 3064). Prisms, insol. alkalis. Reduced by zinc-dust to a phenyl-methyl-anthranol.

PHENYL-METHYL-OXAZOLE C10H,NO i.e. o<CPh:CH [45°]. (242°). Formed by heating acetamide with bromo-acetophenone at 125° (Blümlein, B. 17, 2578; Lewy, B. 20, 2576; 21, 924). Needles (from alcohol). Yields benzoic soid on oxidation. Na and alcohol form an oily tetrahydride $C_{10}H_{12}NO$ (251°), which yields crystalline $C_{10}H_{12}BZNO$ [140°]. HI and P give oily $C_{18}H_{18}$ (c. 275°). Conc. nitric acid forms $C_{10}H_8(NO_2)NO$ [157°], which may be reduced to $C_{18}H_8(NH_2)NO$ [115°]. Hydroxylamine, phenylhydrazine, and hot zinc-dust have no action.—

 $\begin{array}{ll} B'HCl.-B'_2H_2PtCl_s\,2aq.-B'C_sH_sN_sO_7. & [134^\circ]. \\ \textbf{Phenyl-methyl-oxazole} & O<\begin{matrix}CPh:N\\CMe:CH.\end{matrix} (240^\circ). \end{array}$ Formed by boiling benzamide with chloroacetone (Lewy, B. 21, 2193). Oil with fruity odour.—B'₂H₂PtCl₆ 2aq. [170°]. Yellow needles.

Dihydride $0 < \frac{\text{CPh} = \text{N}}{\text{CHMe.CH}_2}$. (244°). Formed

by the action of KOH on bromo-propyl-benzamide in hot aqueous or alcoholic solution, the product being distilled with steam (Gabriel a. Heymann, B. 23, 2499). Liquid. Partly converted, by evaporation with excess of HClAq (or HBrAq), into β-chloro- (or bromo-) propyl-benzamide [78°] (or [73°]). Boiling with an equivalent of HClAq yields & amido-propyl benzoate.

—B',H,PtCl,...B',H,Cr,O,...B'C,H,N,O,.[167°].

PHÉNYL-TRÍ-METHYL-PHENYL-CARBI.

NOL C₁₆H₁₈O i.e. CHPh(OH).C₆H₂Me₈. Phenylmesitylene carbinol. [34°]. (330°). Formed by reducing benzoyl-mesitylene with sodiumsmalgam (Louise, Bl. [2] 45, 231; A. Ch. [6] 6, 209). Prisms. Boiling diluted H₂SO₄ (1:1) converts it into (C_{1e}H_{1,7})₂O [137°].

Ethyl ether C_{1e}H_{1,7}EtO. [32°]. Prisms.

Acetyl derivative C_{1e}H_{1,4}AcO. [52°].

Benzoyl derivative. [94°]. Crystals. DIPHENYL TRIMETHYLPHENYLENE DI-**KETONE** (C₆H₃.CO)₂C₆HMe₃. [117°]. Formed by heating C₆H₃.CO.C₆H₂Me₃ with BzCl and AlCl₂ (Louise, C. R. 98, 151; A. Ch. [6] 6, 236). Prisms, sol. alcohol and ether.

Di-phenyl tetramethylphenylene diketone C_sMe_s(CO.C_sH_s)₂. [270°]. A product of the action of BzCl and AlCl, on durene (Friedel a. Crafts, C. R. 88, 880). Minute prisms, sol. benzene, almost insol. boiling alcohol.

PHENYL METHYLPHENYL KETONE v. PHENYL TOLYL KETONE.

Phenyl dimethylphenyl ketone v. PHENYL

XYLYL KETONE. Phenyl trimethylphenyl ketone

C₀H₃.CO.C₆H₂Me₅[1:2:4:6]. Benzoyl-mesitylene. Phenyl mesityl ketone. [35°]. (319°). V.D. 8·18. Formed from BzCl, mesitylene, CS2, and AlCl, (Elbs, J. pr. [2] 35, 485; cf. Louise, A. Ch. [6] 6, 200; C. R. 96, 499; Claus, B. 19, 2879). Prisms (from alcohol), sol. acetone and ligroïn. Does not react with hydroxylamine (Smith, B. 24, 4052).

Reactions.-1. Conc. H2SO, or H2S2O, at 100° gives benzoic acid and mesitylene sulphonic acid.—2. Conc. H.SO₄ at 50° forms phenylmesityl ketone sulphonic acid C1.H15O.SO3H.—3. A mixture of fuming HNO, and H2S,O, forms two trin itro-derivatives [188°] and [145°].
4. HNO. (S.G. 1·1) at 200° forms benzoyl-uvitic or phenyl-tolyl-ketone dicarboxylic acid.-5. Potash-fusion gives mesitylene and KOBz.--6. HI reduces it to benzyl-mesitylene.—7. Sodiumamalgam yields the carbinol .- 8. Chromic acid oxidises it to C.H.CO.C.H.Mer.CO.H.

Phenyl tetra-methyl-phenyl ketone

C.H.CO.C.HMe. [119°]. (343°) Formed by heating durene with BzCl and AlCl, at 120° (Friedel, Crafts, a. Ador, C. R. 88, 980). Crystalline. Decomposed by potash-fusion into benzoic acid and durene. Reduced by HIAq and P at 240° to CH, Ph.C, HMe, [60.5°], (310°).

(300°). Isomeride C₆H₅.CO.C₆HMe. [63°]. Formed from BzCl and isodurene (got from toluene, McCl, and AlCl₂). Crystals (Essner a.

Gossin, Bl. [2] 42, 171). PHENYL TRIMETHYLPHENYL KETONE o-CARBOXYLIC ACID C₆H₂Me₃.CO.C₆H₄.CO₂H. [212°]. Formed from phthalic anhydride, mesitylene, and AlCl, (F. Meyer a. Gresly, B. 15, 639; A. 234, 234). Needles (from alcohol).

Phenyl tetramethylphenyl ketone carboxylic

acid v. Tetra-methyl-benzoyl-benzoic acid. PHENYL - TRIMETHYLPHENYL - THIO-UREA NHPh.CS.NHC, H, Me,. [198°]. Formed by heating methyl-thiocarbimide with an alcoholic solution of aniline (Eisenberg, B. 15, 1014).

PHENYL-DI-METHYL-PHOSPHINE PMe₂Ph. (192° cor.). S.G. 11 .977. Formed from PCl₂Ph and ZnMe₂ in benzene (Michaelis, A. 181, 359). Liquid, oxidised by air. Combines with CS₂ forming PMe₂PhCS₂ crystallising in red plates and yielding B'2H2PtCla (Czimatis, B. 15, 2017).

Salts.—B'HCl: crystalline, sl. sol. ether, v. e. sol. alcohol.—B'2HCl. Liquid.—B'2Hcl.,

Methylo-iodide PMe, PhI. [205°].

Bromo-ethylo-bromide $PMe_2Ph(C_2H_4Br)\check{B}r.$ [173°]. Formed PMe.Ph and C.H.Br. (Gleichmann, B. 15, 199). Yields (B'(C₂H₄Br)Br)₂PtCl₄ and the unstable B'C₂H₄Br₆. An alcoholic solution of phenyldi-methyl-phosphine forms (PMe2Ph)2C2H4Br2 [above 800], S. (alcohol) 2 at 21°, which gives unstable $(PMe_2Ph)_2C_2H_4Br_6$ [171°]

Di-phenyl-methyl-phosphine PMePh2. (284°). S.G. 15 1.08. Formed from PClPh2 and ZnMe2 (Michaelis a. Link, A. 207, 210). Liquid. Yields PMe₂Ph₂I [241°], (PMe₂Ph₂)₂PtCl₆ [218°], and PMePh₂O [112°] (Michaelis a. La Coste, B. 18,

2116; A. 229, 316).

PHENYL DI . METHYL-PHOSPHINE $\begin{array}{ll} \textbf{CARBOXYLIC} & \textbf{ACID} & \textbf{C}_{\text{o}}\textbf{H}_{\text{d}}(\textbf{CO}_{2}\textbf{H}).\textbf{PMe}_{2}.\\ \textbf{methylo-chloride} & \textbf{C}_{\text{e}}\textbf{H}_{\text{d}}(\textbf{CO}_{2}\textbf{H}).\textbf{PMe}_{2}\textbf{Cl}, & \textbf{f} \end{array}$ formed by oxidation of the methylo-chloride of p-tolyldi-methyl-phosphine, crystallises in prisms (Michaelis, B. 15, 2018). It yields the compounds B'Me,PtCl, C,H,(CO,H).PMe,OH 2aq, and C,H,(CO,H)PMe,SO,H.

Oxide C,H,(CO,H).POMe, [243°]. Formed

by oxidation of the ethyleno-bromide of tolyl-di-

methyl-phosphine. Colourless prisms.

PHENYL METHYL PROPYLENE KETONE C₆H₃.CO.CHEt.CO.CH₃. (265°-270°). Formed from benzoyl-acetone, Na, and EtI (Claisen a. Lowman, B. 21, 1152). Liquid.

PHENYL - METHYL-PROPYLENE-4-THIO-UREA CHMe.S CNPhMe. (c. 300°). Formed by heating methyl-aniline with allyl thiocarbimide and HClAq at 100° (Prager, B. 22, 2996).

—B'2H_PtCl_s. [184°].—B'C_sH₂N_sO_s. [125°].

TRI-PHENYL-METHYL-PROPYL-DI-THIO-BIURET C₂S₂N₃Ph₃MePr. [110°]. Formed from NPhMe.CSCl and di-phenyl-propyl-thio-urea (Billeter a. Strohl, B. 21, 109). Small yellow pyramids. An isomeride [111°] is formed from NPhPr.CSCl and di-phenyl-methyl-thio-urea.

DI - PHENYL - METHYL - PROPYL - THIO-UREA NPhMe.CS.NPhPr. [56°]. Formed from CSCl₂, propyl-aniline, and MeCl or from CSCl₂, methyl-aniline, and PrCl (Billeter a. Strohl, B. 21, 103). Colourless prisms.

DI-PHENYL-DI-METHYL-PYRAZINE

 $CMe \leqslant_{CPh.N}^{N:CPh} \geqslant CMe.$ [124°]. Formed when phenyl amido-ethyl ketone is allowed to stand with NH₃Aq (Schmidt, B. 22, 3253).—B'HCl: needles.—B'₂H₂PtCl₆.

Di-phenyl-methyl pyrazine dihydride

 $N < CPh.CPh \\ CH_2.CHM_0 > N.$ [112°]. Formed from benzil and propylene-diamine (Strache, B. 21, 2363). Needles, v. sol. benzene, m. sol. alcohol.

Di-phenyl-tetra-methyl-pyrazine dihydride C₂₀H₂₂N₂ i.e. NPh CMe:CMe NPh. [108°]. (281°). Formed by heating \$\beta\$-bromo-\$-acetylpropionic acid (1 pt.) with aniline (3 pts.) at 95°; the yield being 70 p.c. of the theoretical amount (Wolff, B. 20, 429). Satiny plates, v. sol. ether and conc. HClAq.

References .- NITRO- and OXY-DI-PHENYL-METHYL-PYRAZINE.

PHENYL-METHYL-PYRAZOLE

NPh $<_{\text{CH:CH}}^{\text{N=CMe}}$. [37°]. (255°). S.G. 15 1.085.

Formation.—1. By heating oxy-phenylmethyl-pyrazole NPh $<_{\text{CO. CH}_2}^{\text{N=CMe}}$ with zinc-dust (Knorr, A. 238, 199).-2. From acetoacetic aldehyde, phenyl-hydrazine, and HOAc (Claisen, B. 21, 1147; 24, 1888).—3. By heating its carboxylic acid at 210° (Ach, A. 253, 55).—4. From oxy-phenyl-methyl-pyrazole dihydride and P2S, (Knorr a. Duden, B. 25, 766).

Properties.—Needles, v. sol. alcohol and ether; volatile with steam. Oxidised by KMnO₄ to phenyl-pyrazole carboxylic acids [146°].

Salt.—B'H2PtCl.: orange needles.

NPh CH2.CH2 Dihydride [75°]. Formed by reducing the base in alcoholic solution with Na. Flat needles, m. sol. alcohol. Phenyl-methyl-pyrazole NPh N=CH

(262°). Formed by distilling its dicarboxylic acid (Knorr a. Laubmann, B. 22, 178). Got also from acetone, oxalic acid, and phenylhydrazine (Claisen a. Roosen, B. 24, 1890). Yields phenyl-pyrazole carboxylic acid [183°] on oxidation by KMnO..—B'2H.PtCl, 2aq. [147°]. Phenyl-di-methyl-pyrazole

NPh < N = CMe. (273° i.V.). Formed by heating its carboxylic acid (Knorr, B. 20, 1103) and by warming acetyl-acetone with phenylhydrazine (K.; Combes, Bl. [2] 50, 145). Oil, volatile with steam. B' H2PtCl 4aq. Decomposes at 186°.

Methylo-iodide B'MeI. [190°]. Crystals v. sol. water.—B'2Me2PtCl. Decomposes at 220°.

Dihydride NPh CHMe.CH₂. (290° i.V.). Formed by adding Na to a boiling alcoholic solution of the base.

Phenyl-tri-methyl-pyrazole. Dihydride $NPh < \begin{array}{c} N = CMe \\ CMe_2 \cdot CH_2 \end{array}.$ Formed by heating mesityl oxide with phenyl-hydrazine (Fischer a. Knoevenagel, A. 239, 203). Oil, v. e. sol. ether and alcohol, sl. sol. water. Its solution in dilute HCl gives a violet colour with K2Cr2O7-B'2H2PtCla: crystalline pp.

Di-phenyl-methyl-pyrazole C16H14N2 i.e.

NPh \(\frac{\text{N} - \text{CPh}}{\text{CMe:CH}} \). [63°]. (335° or 355°). Formed by heating its carboxylic acid at 250°, and also by warming benzoyl-acetone with phenyl-hydrazine (Knorr, B. 18, 314; Fischer, B. 18, 2135). Insol. water, v. sol. alcohol. Weak base. Yields $C_{18}H_{13}BrO_2$ [75°]. — $B'_2H_2PtCl_8$ aq. — B'MeI. [187°].— $B'_2Me_2PtCl_8$. [241°]. Orange-red needles.

Dihydride C₁₀H₁₆N₂. [109°]. Long prisms (from ether). Its acid solution is coloured in-

tensely red by nitrous acid.

Di-phenyl-methyl-pyrazole C18H14N2 NPh<N-CH (365°). Formed by heating its carboxylic acid, and also, together with its dihydride, by distilling the phenylhydrazide of benzylidene-acetone (Knorr, B. 18, 931; 20, 1100). Weak base.—B'₂H₂PtCl₂: red prisms.—B'MeI. [192°].—B'₂Me₂PtCl₂. [229°].

Dihydride $C_{18}H_{18}N_2$. [114°]. (c. 350°). Prisms. Its solutions fluoresce blue and give a

blue colour with nitrous acid.

Tri - phenyl - methyl-pyrazole. Tetrahy-NPh CHPh. CHPh. [110°]. Formed by the action of Na and boiling alcohol on triphenyl-pyrazole methylo-iodide (Knorr a. Laubmann, B. 21, 1205). Insol. water, v. sol. alcohol. Resinified by conc. HNO.

References .- OXY-AMIDO- and OXY-PHENYL-METHYL-PYRAZOLE.

PHENYL - METHYL - PYRAZOLE CARB. OXYLIC ACID NPh N-C,CO,H [106°].

Formed from acetone, oxalic acid, and phenyl hydrazine (Claisen a. Roosen, B. 24, 1891).

Methylether MeA'. (256° at 109 mm.).

Amide. [146°]. Phenyl-methyl-pyrazole carboxylic acid NPh $<_{C(CO,H):CH}^{N}$. [166°]. Formed by heating NPh CO.C(OH) CH with HClAq at 170° (Ach, A. 253, 54). Needles, sol. hot alcohol.

Phenyl-methyl-pyrazole dicarboxylic acid $C_{12}H_{10}N_2O_4$ i.e. $NPh < N = C.CO_2H$ $CMe: \dot{C}.CO_2H$. Formed by oxidising phenyl-di-methyl-pyrazole carboxylic acid with alkaline KMnO, (Knorr a. Laubmann, B. 22, 177). Needles (from water).

Phenyl-di-methyl-pyrazole carboxylic acid NPh \ \frac{N = CMe}{CMe: C.CO_2H} \cdot [197°]. Got by saponifying its ether, which is obtained from ethylideneacetoacetic ether and phenyl-hydrazine (Knorz, B. 20, 1102). Needles, v. sl. sol. water.—KA'. Ethyl ether EtA'. [68°]. (286° at 260 mm.).

Di-phenyl-methyl-pyrazole carboxylic acid $C_{17}H_{14}N_2O_2$ i.e. $NPh < N = CPh \\ OMe: C.CO_2H$. [205°].

Got from its ether, which is obtained by the action of phenyl-hydrazine on benzoyl-acetoacetic ether (Knorr a. Blank, B. 18, 311). Sol. alkalis and conc. HClAq, insol. water.

 $Ethyl\ ether\ { t EtA'}.$ [122°].

Di-phenyl-methyl-pyrazole carboxylic acid $NPh < N = C.CO_2H$. [194°]. Formed from its ether, which is got by mixing benzylidene-acetoacetic ether with phenyl-hydrazine (Knorr a. Blank, B. 18, 931). Crystalline, v. sol. alkalis and acids.

Ethyl ether EtA'. [110°].

References .- NITRO- and OXY-PHENYL-METHYL-

PYRAZOLE CARBOXYLIC ACID.

PHENYL-DI-METHYL-PYRIDAZINE. Dihydride NPh NH .CMe CII. [82°]. (176° at 730 mm.). Formed by heating the dicarboxylic acid at 220° (Knorr, B. 18, 304, 1568). Volatile with steam. Insol. water and alkalis, sol. conc. HClAq. Colours pinewood, dipped in HClAq, red. H₂SO₄ added to its solution in HOAc containing phenanthraquinone gives a deep-red colour.

Dicarboxylic acid of the dihydride C14H14N2O4. Obtained from its ether, which is formed by mixing acetic acid solutions of diacetyl-succinic ether and phenyl-hydrazine. Needles (from HOAc).—BaA".

Ethyl ether Et₂A". [127°]. Prisms.

Reference.—OXY-PHENYL-METHYL-PYRIDAZINE. PHENYL-DI-METHYL-PYRIDINE

 $N \leqslant CMe:CH > CPh$. [55°]. (287°) at 731 mm. Formed by distilling its dicarboxylic acid with lime (Bally, B. 20, 2591). Prisms (from other).

—B'HCl 3aq: needles.—B'₂H₂PtCl₆4aq: needles. [177°]. Needles. — B'H₂Cr₂O₃. - B'HNO_s. Needles. -B'C, H, N,O, [222°]. Needles.

Methylo-iodide B'MeI. Granules, sl. sol.

water.—B'_Me_PtCl_s. [250°]. Hexahydride C_bNH_sMe_Ph. (274°) at 731 mm. Got by adding Na to an alcoholic solution of phenyl-di-methyl-pyridine. Liquid. not give a pp. with picric acid.-B'HCl.-B'HNO₃. [210°].—B'₂H₂PtCl_s.

References.—Nitro- and [237°]. Plates. OXY-PHENYL-DI-

METHYL-PYRIDINE PHENYL - METHYL - PYRIDINE DICARB-OXYLIC ACID C,4H11NO4 i.e.

CO₂H.C_eH.,C</br>
CH.CH.: OMe
C(CO₂H):N
[201°]. Formed oxidising methyl - (B) - naphthoquinoline by KMnO, in acid solution (F. Seitz, B. 22, 257). Needles (containing aq) or anhydrous FeSO, colours its aqueous solution llow. — Na₂A" 2aq. — ZnA" 1½aq. femon-yellow. -CuA" 1 aq: minute bluish-green plates.

Phenyl-di-methyl-pyridine carboxylic acid N≪CMe:C(CO,H)>CPh. [190°]. Got from its ether, which is formed by heating the monoethyl ether of the dicarboxylic acid (Hantzsch, B.17, 2911). Small prisms (containing 2aq), v. sol. hot water.—CuA'₂.—(HA')₂H₂PtCl₂ aq.

Ethyl ether EtA'. (316°-820°). Thick

liquid. Yields (EtA'), H.PtCl. [196°] and also

EtA'MeI [c. 206°], which is converted by alcoholic potash into C₁₅H₁₅NO₂ [161°], split up by HClAq at 180° into acetic acid and oxy-phenyldi-methyl-pyridine [112°].

Phenyl-di-methyl-pyridine dicarboxylic acid

 $N \leqslant_{\mathrm{CMe.C(CO_2H)}}^{\mathrm{CMe.C(CO_2H)}} \geqslant_{\mathrm{CPh.}}$

HEtA". [180°]. Mono-ethyl ether Formed by boiling the di-ethyl ether with alcoholic KOH (1 mol.) (H.). Cubes (from alcohol).

Di-ethyl ether Et, A". [67°]. Formed by oxidising its dihydride with nitrous acid.

Dihydride of the di-ethyl ether $N \leqslant_{\text{CMe}---\text{C(CO}_2\text{Et)}}^{\text{CHMc.CH(CO}_2\text{Et)}} > \text{CPh.} \quad [157^\circ]. \quad \text{Formed}$

by heating a mixture of benzoic aldehyde, acetoacetic ether, and alcoholic NH₃ (Schiff a. Puliti, B. 16, 1607; Bally, B. 20, 2591). Formed also by mixing benzylidene-acetoacetic ether with β -

imido-butyric ether (Beyer, B. 24, 1666). Colourless crystals, sol. alcohol.

(β)-Phenyl-di-methyl-pyridine dicarboxylic $\text{[1:2]CO}_2\text{H.C}_6\text{H}_4\text{.C} \leqslant^{\text{CMe}}_{\text{C(CO}_2\text{H).N}} > \text{CMe}.$

Formed by oxidising di-methyl-(8)-naphthoquinoline with KMnO₄ (Reed, J. pr. [2] 35, 311). Syrup.—AgA'.

Reference.—Oxy-PHENYL-DI-METHYL-PYRIDINE

CARBOXYLIC ACID.

PHENYLTRI-METHYL-PYRIDYL KETONE CARBOXYLIC ETHER

 $N \leqslant_{CMe:C(CO.Ph)}^{CMe.C(CO.Et)} > CMe.$ Formed by the action of nitrous acid on its dihydride. - B'HNO: white plates. — B'HCl. [192°]. Prisms. -B'2H2PtCl6: needles.

 $Dihydride \text{ NH} < \stackrel{\text{CMe:C(CO}_2\text{Et)}}{\text{CMe:C(CO.Ph)}} > \text{CHMe.}$

Formed from ethylidene-acetoacetic ether and C,H₅.CO.CH₂.C(NH).CH₃ (Beyer, B. 24, 1667). Yellowish plates.

Methyl derivative of the dihydride NMe CMe:C(CO.Ph) CHMe. [87°]. Formed, in like manner, by the action of the methylimide of benzoyl-acetone on ethylidene-acetoacetic ether (B.). Yellowish-white needles.

PHENYL-METHYL-PYRIMIDINE C11H10N2 i.e. CPh $<_{
m N:CH}^{
m N.CMe} >$ CH. [74°-78°]. by distilling the oxy-derivative (from benzamidine and acetoacetic ether) with zinc-dust (Pinner, B. 18, 2850). Colourless needles. -B', H, Cl, PtCl,. [190°].

References. Oxy- and Oxy-AMIDO-PHENYL-METHYL PYRIMIDINE.

PHENYL-METHYL-PYRROLE C, H, N i.e. NH CMe:CH. [101°]. Got by heating acetophenone-acetone CH₂Bz.CH₂Ac with alcoholic NH₃ at 150°; the yield being 70 p.c. of the theoretical (Paal, B. 18, 367). Plates, v. sol. alcohol. May be sublimed. Gives a purple-red colour with isatin and H2SO4. Phenyl-di-methyl-pyrrole C12H12N i.e.

NPh CMe.CH [52°]. (252°). Formed by distilling its dicarboxylic acid at 244° (Knorr, A. 236, 306).

DI-PHENYL-METHYL-PYRROLE $C_1, H_{13}N$ Le. NPh CPh: (OH [84°]. Formed by heating the carboxylic acid [226°] to above its melting-point (Lederer a. Paal, B. 18, 2596). Large tables.

Tetra-phenyl-methyl-pyrrole

NMe CPh: CPh [214°]. Formed by heating by by leaving by leaving by leaving by leaving at 150° (Fehrlin, B. 22, 554). Groups of minute needles, sol. hot alcohol and ether.

Reference.—OXY - TRI - PHENYL - METHYL - PYR-

PHENYL-METHYL-PYRROLE CARBOXY-LIC ACID NHCCMe:C.CO₂H. [c. 190°]. Got

by saponification of its ether, which is formed by allowing acetophenone-acetoacetic ether to stand with excess of NH₂Aq (Lederer a. Paal, B. 18, 2593). Flat needles (from HOAc).

Ethylether EtA'. [120°]. Needles.

Phenyl-di-methyl-pyrrole carboxylic ether C₁₈H₁₇NO₂ i.e. NMe CMe:C.CO₂Et. [112°]. Formed from acetophenone acetoacetic ether and NMeH₂Aq (L. a. P.). Plates (from alcoholether), v. e. sol. alcohol.

Phenyl-di-methyl-pyrrole m-carboxylic acid CO₂H.C₃H₄.N

CMe:CH. [135°]. Formed by heating acetonyl-acetone with m-amido-benzoic acid and alcohol (Paal, B. 19, 558). Flat crystals, sl. sol. water, v. sol. alcohol.

Phenyl-di-methyl-pyrrole di-carboxylic acid ${\rm CO_2H.CH_2.N} < {\rm CMe:C.CO_2H.}$ [152°]. Got by saponifying its ether. Small needles (from alcohol), sl. sol. water.

Ethyl ether Et₂A". [131°]. Got by allowing acetophonone-acetoacetic ether to stand with glycocoll and HOAc (Paal a. Schneider, B. 19, 3160). Needles (from HOAc), sl. sol. water.

Phenyl-di-methyl-pyrrole dicarboxylic acid NPh CMe:C.CO₂H. Gotfrom its ether. Powder, decomposing at 224° into CO₂ and phenyl-dimethyl-pyrrole.—CaA".—CaH₂A"₂: prisms.

Ethyl ether Et₂A". [38°]. (280°) at 500

Ethyl ether Et₂A". [38°]. (280°) at 500 mm. Formed by warming di-acetyl-succinic ether with aniline and HOAc (Knorr, Λ. 236, 305). Crystals.

Di-phenyl-methyl-pyrrole carboxylic acid

NPh CPh:CH (226°]. Obtained by saponification of its ether, which is formed by boiling acetophenone-aceto-acetic ether with an acetic acid solution of aniline (Lederer a. Paal, B. 18, 2595). Small needles (from acetic acid). Ethyl ether A'Et: [100°]; prisms.

Di-phenyl-methyl-pyrrole di-o-carboxylic acid NMe< $C(C_aH_+,CO_2H)$:CH (C_aH_+,CO_2H) :CH (C_aH_+,CO_2H) :CH (C_aH_+,CO_2H) :CH (C_aH_+,CO_2H) and methylamine in alcohol at (C_aH_+,CO_2H) and methylamine in alcohol at (C_aH_+,CO_2H) :CO (C_aH_+,CO_2H) . Deepyellow plates, insol. water, sl. sol. ether.

Di-phenyl-methyl-pyrrole dicarboxylic acid CO₂H.C_eH_e.N CMe:C.CO₂H. [210°]. Formed by saponifying its ether, which is got from acetophenone-acetoacetic ether, m-amido-benz-

oic acid, and HOAc (Paal a. Schneider, B. 19, 3162). Needles (from dilute alcohol).

Éthyl ether EtA'. [160°]. Néedlos. Reference.—Oxy-Phenyl-Methyl-Pyrrole Carboxylic acid.

PHENYL-METHYL-QUINAZOLINE DIHY.

DRIDE C₁₈H₁₄N₂ i.e. C₈H₄<N-CMe (60°). Formed by reduction of the actyl derivative of o-nitro-benzyl-aniline with tin and HClAq (Paal a. Krecke, B. 23, 2638). Rosettes of white needles, insol. water, m. sol. ligroin. The tin double chloride melts at 122°.—B'HCl 2aq. [257°].—B'₂H₂PtCl₈. Flat orange needles. Decomposes at 223°.

Reference.—OXY - PHENYL - METHYL - QUINAZ-OLINE.

(Py. 1,3) - PHENYL-METHYL-QUINOLINE C_eH₁ < CPh:CH (100°). Formed by boiling an alcoholic solution of o-amido-benzophenone with acetone and KOHAq (Geigy a. Königs, B. 18, 2406; 19, 2428). Obtained also from paraldehyde, acetophenone, HCl, and aniline, and by heating benzoyl-acetone-anilide C_{1e}H_{1s}NO with H₂SO₄ on the water-bath (Beyer, J. pr. [2] 33, 420; B. 20, 1771). Tables (from ether).—B'H₂SO₄. [236°].—B'₂H₂PtCl_e2aq. [225°]. Its acid solutions fluoresce blue. By heating with phthalic anhydride and ZnCl₂ at 150° it is converted into the phthalone C₂;H_{1s}NO₂ [270°], which crystallises from alcohol or HOAc in sparingly-soluble orange-red crystals.

(Py. 3,1)-Phenyl-methyl-quinoline C_{1e}H_{1s}N i.e. C_eH₄<N=CMe:CH Flavoline. [65°]. (374°). V.D. 7·7 (calc. 7·6). Formed by distilling flavenol with zinc-dust. Prepared by adding a little dilute (10 p.c.) NaOHAq to a solution of equimolecular quantities of o-amido-acetophenone and acetophenone, and then heating on the water-bath (Fischer, B. 15, 1503; 16, 68; 19, 1037). Thick tables. Yields flavaniline on nitration and reduction. Salts.—B'HCl 2aq.—B'HT4PtCl_e: reddish-yellow needles.—B'MeI. [185°] (Bernthsen a. Hess, B. 18, 34). Converted by KOHAq into a strongly alkaline solid 'methyl-flavolinium hydrate.'—B'₂Me₂PtCl_e.

(Py. 3,2)-Phenyl-methyl-quinoline $C_{\rm e}H_4 < \begin{array}{c} {\rm CH:CMe} \\ {\rm N=CPh.} \end{array}$ [53°]. (above 800°). Formed by heating a mixture of CHPh:CMe.CHO, aniline, and conc. HClAq at 200° (Miller a. Kinkelin, B. 19, 527). Trimetric prisms, v. sol. alcohol and ether.— $B'_2H_2PtCl_6$: orange plates.— $B'C_6H_3N_3O_7$. [202°]. Large yellow plates.

(B. 2)-Phenyl-(Py. 4)-methyl-quinoline tetrahydride C_{1e}H₁,N i.e. CPh:CH.C.CH₂.CH₅.

Formed by reducing (B. 2)-phenyl-quinoline methylo-chloride with tin and HClAq (La Coste a. Sorger, A. 230, 24). Amorphous powder, insol. cold water.—B'HCl.—B'HI.—B'C₂H₂N₂O₃.

[147°].—B'MeI aq. [195°]. Yellow plates.

($\dot{P}y$. 3)-Phenyl-(\dot{B} . 4)-methyl-qui_oline CH:CH .C.CH:CH [50°]. Formed by distilling its (Py. 1)-carboxylic acid with soda-lime (Doebner a. Giesecke, A. 242, 299). Plates (from alcohol).—B'₂H₂PtCl₄. Brick-red needles. (Py. 5)-Phenyl-(B. 2)-methyl-quinoline CM:CH.C.CH:CH (68°). (above 360°). Got by distilling its carboxylic acid with soda-lime (D. a. G.). Small yellow needles (from dilute alcohol).—B'₂H₂PtCl₆: golden needles.

References .- NITRO - and OXY-PHENYL-METHYL-

QUINOLINE.

(Py. 3)-PHENYL-(B. 4)-METHYL-QUINOL-INE (Py. 1)-CARBOXYLIC ACID C₁,H₁₃NO₂. [245°]. Formed by heating equi-molecular proportions of o-toluidine, benzoic aldehyde, and pyruvic acid in alcoholic solution on the waterbath (Doebner a. Giesecke, A. 242, 298). Small yellow crystals (from alcohol), v. sl. sol. hot water.—CuA'₂ aq.—AgA' aq: needles (from Aq).

(Py. 3)-Phenyl-(B. 2)-methyl-quinoline (Py. 1)-carboxylic acid. [228°]. Formed in like manner from p-toluidine (D. a. G.). Yellow needles (from alcohol).—CuA'₂.—PbA'₂.—AgA'.
—H₂A'₂H₂PtCl₅: golden needles, v. sl. sol. hot Aq.
Reference.—Oxy-PHENYL-METHYL-QUINOLINE

CARBOXYLIC ACID.

PHENYL (Py. 3)-METHYL-(B. 2)-QUIN-OLYL KETONE C₆H₃-CO.C₆H₃-CH.CH. (B8°]. (above 300°). Formed from amido-benzophenone, paraldehyde, and HClAq at 100° (Hinz, A. 242, 323). Needles (from water) or plates (from alcohol).—B'₂H₂PtCl₆ 2aq.—B'₂H₂Cr₂O₇.—B'MeI. [220°]. Small dark-green needles; v. sol. water.

Phenyl (Py.3)-methyl-(B.4)-quinolyl ketone. [108°]. Formed by boiling o-amido-benzophenone with dilute H₂SO₄ and paraldehyde (Geigy a. Königs, B. 18, 2406). Matted needles (from

dilute alcohol).

PHENYL - DI - METHYL - QUINOLYL - THIO-UREA NHPh.CS.NH.C₁₁H₁₀N. [159°]. Formed by heating phenyl-thiocarbimide with amido-dimethyl-quinoline (Marckwald, B. 23, 1025). Prisms, v. sol. hot alcohol.—B'₂H₂PtCl₆: pp.

PHENYL-METHYL-QUINOXALINE

CMe:CH.C.N:CPh
CH:CH.C.N:CH. [79°]. Formed by reducing
C₆H₃Me(NO₂).NH.CH₂Bz with SnCl₂ and HCl
(Lellmann a. Donner, B. 23, 171). Needles, v.
sol. alcohol.—B'HgCl₂. [223°]. White needles.
Phenyl-methyl-quinoxaline

CMe:CH.C.N:CH CH:CH.C.N:CPh. [135°]. Formed, together with the preceding isomeride, by the action of chloroacetophenone on tolylene-o-diamine (Hinsberg, A. 237, 370). Formed also by warming phenylglyoxylic aldehyde C.H. CO.CHO with tolyleneo-diamine sulphate and NaOAc (Pechmann, B. 20, 2905). Needles, v. sol. hot alcohol and ether.

Phonyl-di-methyl-quinoxaline

C_sH_sMe</h>
N:CMe
N:CPh. [48°]. Formed by boiling phenyl methyl diketone with tolylene-o-diamine in ether (Müller a. Pechmann, B. 22, 2130). V. sol. most solvents.

Di-phenyl-methyl-quinoxaline C₂₁H₁₂N₂ i.e.

C₆H₄Me

N:CPh. [111°]. Formed by heating benzil with tolylene-o-diamine in alcoholic solution (Hinsberg, B. 17, 322). Silvery plates. May be distilled. Weak base. Nearly insol. Aq. Di-phenyl->-methyl-quinoxaline dihydride

C₆H₄

NMe.CHPh. [138°]. Formed by heating

phenylene-methyl-o-diamine with benzoin at 165° (Fischer a. Busch, B. 24, 2682). Yellow needles, m. sol. alcohol. Its solutions fluoresce greenish-yellow.

Reference. — OXY - PHENYL - METHYL - QUIN-

OXALINE.

PHENYL-METHYL-ROSINDULINE

 $C_{10}H_s(NPh) \gtrless_{NPh}^N > C_eH_sMe \begin{bmatrix} 1\\2 \end{bmatrix} 4$. [232°]. Formed from benzene-azo-p-tolyl-(a)-naphthylamine (Fischer a. Hepp, A. 256, 243). Reddishbrown bronzed plates. Split up by conc. HClAq at 200° into aniline and methyl-rosindone.

PHENYL-METHYL-SUCCINIC ACID v

PHENYL-PYROTARTARIC ACID.

Phenyl-tri-methyl-succinic acid C₁₃H₁₆O₁ i.e. CH₂Ph.CH(CO₂H).CMe₂·CO₂H. [140°]. Formed from bromo-isobutyric ether and sodium benzylmalonic ether, the resulting ether (200°–220°) being saponified (Bischoff, B. 24, 1060). Short needles (from water), v. sol. alcohol and ether.

Di-phenyl-di-methyl-succinic acid. Nitrile CPhMe(CN).CPhMe.CN. [227°]. Formed from CHPhMe.CN by treatment with NaOEt and I (Chalanay a. Knoevenagel, B. 25, 289). Insol.

benzene, sl. sol. alcohol.

PHENYL-METHYL-SULPHAMIC ACID

NPhMe.SO₃H. Formed from ClSO₃H and methyl-aniline in chloroform (Traube, B. 24, 362).—NH₄A'. Turns violet in air. Decomposed by boiling HClAq into methyl-aniline and H₂SO₄. The potassium salt is a crystalline powder, sl. sol. alcohol, v. sol. water.

PHENYL-DI-METHYL SULPHAMIDE SO₂(NMe₂)(NHPh). [85°]. Formed from NMe₂SO₂Cl and aniline (Behrend, A. 222, 128). Needles (from ether). Yields SO₂(NMe₂)(NPhNa) crystallising in needles, v. sol. water and alcohol.

PHENYL METHYL SULPHIDE PhSMe. (188°). Formed from Pb(SPh)₂ and MeI at 100°

(Obermeyer, B. 20, 2926). Liquid. PHENYL METHYL SULPHONE C, H_aSO₂ i.e. C₆H_aSO₂.CH₃. [89°]. Formed from benzene sulphinic acid, NaOEt, and MeI at 100°, and also by boiling O₆H₃.SO₂.CH₂.CO₂H with alcoholic potash (Michael a. Palmer, Am. 6, 254; 7, 65; Otto a. Rössing, B. 18, 156; 21, 652; 22, 1449, 1452; 23, 755). Plates (from hot water), v. e. sol. alcohol and ether.

PHENYL METHYL SULPHONE ω -CARB-OXYLIC ACID v. PHENYL-SULPHONO-ACETIC ACID.

PHENYL-METHYL-THIAZOLE C₁₀H₂NS i.e. N≪CMe: CH (279° cor.). Formed from thiobenzamide and chloro-acetone in alcohol (Hubacher, A. 259, 236). Oil.

Phenyl-methyl-thiazole N CH: CH. CMe.S . [68·5°]. (284° cor.). Formed from thioacetamide and ∞-bromo-acetophenone (Hantzsch, B. 21, 913; A. 250, 269). Weak base. Not attacked in boiling alcoholic solution by sodium (Schatzmann, A. 261, 7).—*B'HCl: sl. sol. Aq.

Phenyl-methyl-thiasole. Dihydride

NCH₂.CHMe. Formed by boiling propylene
bromide with thiobenzamide (Gabriel a. Heymann, B. 24, 785). Yellowish liquid.—Pierate
B'C₂H₂N₃O₇: crystalline.

Di-phenyl-methyl-thiasole N CPh: CPh

[52°]. Formed from thioacetamide and bromodeoxybenzoin in alcohol (Hubacher, A. 259, 244). Stellate groups of needles, insol. water. -Hydrochloride: [97°]; white crystals (from dilute HClAq).

PHENYL-METHYL-THIAZOLE CARBOXY-LIC ACID N CMe:C.CO₂H. [203°]. Formed

by saponifying its ether. Needles, sl. sol. ether. Ethyl ether EtA'. [43°]. Formed from thiobenzamide and chloro-acetic ether (Hubacher,

A. 259, 237).

PHENYL (a)-METHYL-THIENYL KETONE C_sH_s.CO.C₄H₂MeS. [124°]. Formed by heating O₄H₂Me(HgCl)S with BzCl at 100° (Volhard, A. 267, 181; *of.* Ernst, B. 19, 3280). Needles (from ligroïn).

TRI - PHENYL - DI - METHYL - DI-THIO-**BIURET** $C_{22}H_{21}N_{3}S_{2}$ *i.e.* $NPh(CS.NPhMe)_{2}$ or CS(NPhMe).S.C(NPh)(NPhMe). [202°]. Formed from Cl.CS.NPhMe and aniline (Billeter a. Strohl, B. 21, 108). Needles, v. sl. sol. cold

alcohol

PHENYL - METHYL - DI-THIO-CARBAMIC ACID NPhMe.C3.SMe. [88°].

Formed by heating NPhMe.C(NMe).SMe or NPhMe.C(NH).SMe with CS₂ at 160° (Bertram,

B. 25, 54). Plates.

PHENYL - METHYL - THIOCARBAMIC CHLORIDE NPhMe.CS.Cl. [35°]. Formed from methyl-aniline and CSCl₂ (Billeter, B. 20, 1631). Yellowish crystals. Converted by alcohol into (NPhMe.CS), O [116.5°]. PHENYL - METHYL-THIO-SEMI-CARBAZ-

IDE NHPh.NH.CS.NHMe. [89°]. Formed from methyl-thiocarbimide and phenyl-hydrazine (Dixon, C. J. 57, 262). Trimetric prisms, v. sl. sol. cold water. H2SO4 forms an azure-blue solution. CuSO, gives a deep-blue colour, and FeCl₃ a greenish-blue.

Phenyl-methyl-thio-semi-carbazide

methyl-hydrazine and phen (Brüning. A 252 Formed from phenyl-thiocarbimide (Brüning, A. 253, 11). Prisms (from alcohol).

Di-phenyl-methyl-thio-semi-carbazide NPhMe.NH.CS.NHPh. [154°]. Formed from phenyl-methyl-hydrazine and phenyl-thiocarbimide (Fischer, A. 190, 166). V. sol. hot alcohol.

PHENYL-METHYL-THIOCARBIZINE

 $cs<_{NMe}^{NPh}$. [123°]. Formed from phenyl-thiocarbizine and MeI at 100° (Fischer, A. 212, 330). Tables (from water). Not attacked by nitrous acid.

PHENYL-METHYL-THIOHYDANTOIC ACID NHPh.CS.NH.CHMe.CO.H. Formed alanine and phenyl-thiocarbimide in alcohol (Marckwald, B. 24, 3280; cf. Aschan, B. 17, 421).-KA': minute needles

PHENYL-METHYL-THIOHYDANTOÏN

CS<NPh.CO NH -CHMe. [184°]. Formed by treating phenyl-methyl-thiohydantoïc acid (v. supra) with HCl. Small prisms (from alcohol). Split up by HClAq at 150° into aniline, alanine, CO2, and CS.

Phenyl-di-methyl-thiohydautoin

CS NPh.CO [67°]. Formed from phenyland a-amido-isobutyric acid thiocarbimide (Marckwald, B. 24, 3282). Crystals, sl. sol. Aq. Vol. IV.

PHENYL-METHYL-THIOPHENE C,H,S i.e. S CMe:CH. [51°]. (271° uncor.). Got by heating acetophenone-acetone with P2S, at 125°, the yield being 70 p.c. of the theoretical amount (Paal, B. 18, 367). Needles, volatile with steam. Gives the indophenine reaction.

s<CMe:CH .CPh. Phenyl-methyl-thiophene

[73°]. Formed by heating CH, Ac. CHPh. CO, Na with P₂S₃ or P₂S₅ (Paal a. Püschel, B. 20, 2558).

Plates, v. sol. hot alcohol. Gives the indophenine reaction on warming. Br forms C₁₁H₅Br₄S [137°]

s-PHENYL-METHYL-THIO-UREA C.H.10N2S i.e. CS(NHPh)(NHMe). [113°]. Formed from methyl-thiocarbimide and aniline (Gebhardt, B. 17, 3038). Six-sided tables. Yields NH, Me and

CS(NHPh), on boiling with aniline.

u-Phenyl-methyl-thio-urea CS(NH₂)(NPhMe). [107°]. Formed from methylaniline hydrochloride and potassium sulpho-cyanide (Gebhardt, B. 17, 2094). Tables (from alcohol) or prisms (from hot water).

Phenyl-methyl-\u00fc-thio-urea

MeS.C(NH)(NHIPh). [71°]. Formed by warming phenyl-thio-urea with MeI and alcohol (Bertram, B. 25, 49). Colourless crystals, sl. sol. hot water. Converted by dilute H₂SO₄ at 160° into MeS.CO.NHPh.—B'HI. [147°].— B'H,8O,. [171°].—B',H,8O,. [1717].—B'HNO,. [113°].—B'HOAc. [115°].—B',H,PtCl₆. [184°]. —B'C₆H₂(NO₂)₃OH. [175°]. Yellow plates. Phenyl-di-methyl-thio-urea

CS(NHMe)(NPhMe). [114°]. Formed from methyl-thiocarbimide and methyl-aniline (G.).

Phenyl-di-methyl-ψ-thio-urea

MeS.C(NH)(NPhMe). Formed from phenylmethyl- ψ -thio-urea and methyl iodide (B.). Oil. Converted by dilute H.SO, at 160° into [184°]. -MeS.CO.NPhMe [54°]. — B'HI. B'C.H.N.O. Small prisms.

Phenyl-tri-methyl-\psi-thio-urea C10H14N2S i.e. MeS.C(NMe)(NPhMe). Methyl-imido-methylphenyl-carbamic thiic methyl ether. Formed by methylation of phenyl-di-methyl-+thio-urea (B.).—B'HI. [c. 184°].—B'₂H₂PtCl₅. [c. 174°]. —B'C₆H₃N₃O₇. [126°]. Lemon-yellow crystals.

Di-phenyl-methyl-thio-urea

CS(NHPh)(NPhMe). [87°]. (205°). Formed from phenyl-thiocarbimide and methyl-aniline (G.). Prisms, v. sol. hot alcohol. Decomposed into the parent substances by distillation with steam. Boiling aniline yields CS(NHPh)2 and methyl-aniline.

Di-phenyl-methyl-\psi-thio-urea

MeS.C(NPh)(NHPh). [110°]. Formed from CS(NHPh), and MeI. Conc. HClAq at 150° yields aniline and methyl-mercaptan.—B'HI.

Di-phenyl-di-methyl-thio-urea CS(NPhMe)2. [72.5°]. Formed from NPhMe.CS.Cl and methylaniline (Billeter, B. 20, 1631). Prisms (from ligroin).

Di-phenyl-di-methyl-\psi-thio-urea MeS.C(NPh)(NPhMe). (above 300°). Formed from CS(NHPh)(NPhMe) and MeI (Bertram, B. 25, 57). CS₂ at 160° forms 11. B'HI. Crystalline meal, sl. sol. hot water. CS, at 160° forms NPhMe.CS,Me.-B'HI.

Reference. -- OXY - DI - PHENYL - METHYL- -- THIO-

UREA.

PHENYL-METHYL-UREA C,H, N,O CO(NH₂)(NPhMe). [82°]. Formed from methylaniline hydrochloride and potassium cyanate (Gebhardt, B. 17, 2095). Thin crystals.

Phenyl-di-methyl-urea CO(NMe2)(NPhMe). Formed from NMe₂.COCl and aniline (Michler a. Escherich, B. 12, 1163). Crystals (from alcohol).

Di-phenyl-methyl-urea CO(NPhMe)(NHPh). [104°]. (204°). Formed from phenyl cyanate and methyl-aniline (G.). Small needles.

Di-phenyl-di-methyl-urea CO(NPhMe),. [121°]. (c. 350°). Formed from NPhMe.COCl and methyl-aniline (Michler a. Zimmermann, B. 12, 1165). Tables (from alcohol).

PHENYL-(B)-NAPHTHACRIDINE

 $C_{10}H_{0} < N > C_{10}H_{0}$. [297°]. S. (alcohol) ·04 at 15°; S. (benzene) 29 at 18°. Formed by heating a mixture of di-(3)-naphthylamine with BzCl or with benzoic acid and P2Os or ZnCl2 (Claus a. Richter, B. 17, 1595; Ris, B. 17, 2029; Klopsch, B. 18, 1586). Needles, v. sol. hot benzene.—B'HCl.—B'₂H₂PtCl₂: yellow needles.

Dihydride C10He CHPh C10He. from benzoic aldehyde (1 mol.) and (β)-naphthylamine (2 mols.) in presence of condensing

agents (Claisen, A. 237, 273). Oxidised to

phenyl-naphthacridine.

enyl-naphthacridine. (β)-PHENYL-NAPHTHALENE $C_{1e}H_{12}$ i.e. VD. 7·12 (calc. 7·05). C, H, C, H, [102°]. V.D. 7.12 (calc. Formed by passing a mixture of naphthalene and bromo-benzene through a red-hot tube (Watson Smith, B. 12, 2049; C. J. 39, 546; C. J. Proc. 5, 70). Formed also by the action of H₂SO₄ (80 g.) diluted with water (44 g.) on CHPh(OH).CH2OH (Zincke a. Breuer, A. 226, 23; 240, 187). Fluorescent plates. May be sublimed. Gives rise on oxidation to a quinone C16H10O2 [110°].

A hydrocarbon C16H12 [104°] got by distilling chrysoquinone with soda-lime is perhaps identical with the above (Graebe, B. 6, 66; 7, 792;

Schmidt, J. pr. [2] 9, 285).

(a)-Phenyl-naphthalene dihydride v. ATRON-

ENE.

Reference.—OXY-AMIDO-PHENYL-NAPHTHALENE. DI-PHENYL-NAPHTHAMIDINE C₂₈H₁₈N₂ Formed i.e. C₁₀H₇.C(NPh).NHPh. [183.5°]. Formed from (a)-naphthoic acid, aniline, and PCl₃ (Bössneck, B. 16, 642). Needles (from alcohol). PHENYL-(β)-NAPHTHINDOLE C₁₈H₁₈N i.e.

 $C_{10}H_{\bullet}$ CPh. [130°]. Formed by heating the (β) -naphthyl-hydrazide of acetophenone with ZnCl₂ at 170° (Ince, A. 253, 42). Crystals, v. sol. alcohol and ether. Colours pinewood green.—B'C,H,N,O,. [166°]. Brown needles.

Phenyl-(β)-naphthindole $C_{10}H_6 < \stackrel{CPh}{NH} > CH$. [211°]. Formed by the action of alcoholic HCl on the (3)-naphthyl-hydrazide of phenyl-acetic aldehyde (I.). Needles, suddenly decomposed on fusion. Colours pinewood green. ZnCl. at 170° converts it into the preceding isomeride. -B'C,H,N,O,. [120°]. Reddish-brown needles. PHENYL-WAPHTHOTRIAZINE. Dihy-

dride C1,H1,N, i.e. C10He N.CH2 [184°].

amine with formic paraldehyde at 140° (Goldschmidt a. Poltzer, B. 24, 1002). Plates (containing $\frac{2}{8}$ aq).—B'HCl. [254°].—B'₂H₂PtCl₂.

Di-phenyl- $(\alpha\beta)$ -naphthotriazine $C_{22}H_{17}N_{2}$ i.e. C₁₀H_s N.CHPh? [194°]. Got by heating a solution of CHPh:N.C₁₀H₆·N₂Ph in HOAc (Meldola, C. J. 57, 330). White flattened needles. PHENYL-NAPHTHOPHENANTHRAZINE

C.H., CH.N. C.H. The phenylo-nitrate B'PhNO₃, formed by boiling phenanthraquinone with phenyl-naphthylene-diamine in HOAc and then adding HNO₃ crystallises in needles or prisms (Witt, B. 20, 1185).

(Py. 3)-PHENYL-(a)-NAPHTHOQUINOLINE

C₁₉H₁₈N i.e. C₁₀H₈ CH:CH N=CPh. [68°]. Formed by heating its (Py. 1)-carboxylic acid with soda-lime (Döbner, A. 249, 114). Yellow needles (from alcohol-ether). Yields a syrupy tetrahydride. Salts. $B_2^*H_2$ PtCl_a2aq. $-B^*H_2$ Cr₂O₂, $-B^*C_8H_8$ N₃O₂, [167°]. Thin needles (from alcohol). (Py. 3)-Phenyl-(β)-naphthoquinoline. [188°].

Formed by heating its carboxylic acid with sodalime (D.). Needles or plates, insol. water, v. sol. alcohol.—B'₂H₂PtCl₅ aq.—B'C₅H₂N₅O₇. [250°]. Yellow plates, sl. sol. ether.—B'2H2Cr2O7.

Ethylo-iodide. [233°]. Plates, insol.

(Py.3)-PHENYL-(a)-NAPHTHOQUINOLINE (Py. 1)-CARBOXYLIC ACID C20H18NO. i.e. $C_{10}H_{\bullet} < N = CPh$ [300°]. Formed heating (a)-naphthylamine with pyruvic acid and benzoic aldehyde in alcoholic solution (D.). Yellow needles (from alcohol-acetone). May be reduced to a tetrahydride. Produces, on oxidation, di-phenyl-pyridine tri-carboxylic acid and $CO_2H.C_5NHPh < \stackrel{C_6H}{CO} - NaA' \frac{1}{2}aq. - CaA'_24aq.$

-ZnA'₂-PbA'₂.-AgA': insoluble powder. Ethyl ether EtA'. [103°]. Needles.

(Py,3)-Phenyl- (β) -naphthoquinoline (Py,1)carboxylic acid C20H12NO2. [296°]. Formed in like manner from (8)-naphthylamine (D.).

Needles, v. sl. sol. alcohol.—KA' 5aq.—NaA' 5aq.

—CaA'₂ 6aq.—ZnA'₂ 2aq.—Cu₂A'₂ aq (?).—AgA'.

PHENYL-NAPHTHOQUINOXALINE

 $C_{10}H_{e} < \stackrel{N:CH}{\sim} N:CPh$. The phenylo-bromide B'PhBr. formed from phenyl-naphthylene-diamine and bromo-acetophenone, crystallises in yellow plates, and is converted by NaOH into B'PhOH [148°], which crystallises in prisms, and yields B'PhNO, (Fischer a. Busch, B. 24, 1873).

Di-phenyl-naphthoquinoxaline

 $C_{10}H_6 < N:CPh \\ N:CPh$ [147°]. Formed by heating naphthylene-(1,2)-diamine hydrochloride with benzil in alcohol (Lawson, B. 18, 2426). Lightbrown plates, v. sol. alcohol and ether.

Phenylo-hydroxide C₁₀H₆<NPh(OH):CPh. [167°]. Formed by oxidation of C₁₀H_s<N=CPh NPh,CHPh [164°], which is got by heating phenyl-naphthylene-o-diamine with benzoin at 200° (Fischer, B. 24, 722, 1871). Formed by heating benzene-azo-(β)-naphthyl- and benzil (Fischer, B. 24, 2679). Yellow prisms, insol. water, v. sol. ether. -B'HNO₂.-B'HCl: dark-vellow needles, v. sol. alcohol.

Di-phenyl-naphthoquinoxaline dihydride $C_{10}H_{6} < NPh.CH_{2} N = CPh$ [165°]. Formed, together with a compound melting at 195°, from naphthylene-phenyl-diamine and benzoyl-carbinol at 155° (Fischer a. Busch, B. 24, 2680). Orange needles, v. sol. benzene, v. sl. sol. alcohol.

PHENYL-NAPHTHOROSINDULINE

 $C_{e}H_{4} < C(NPh).CH:C.NPh.\dot{C}:CH > C_{e}H_{4}. \quad [256^{\circ}].$ Formed, together with naphthyl-rosinduline, from benzene-azo-di-(a)-naphthylamine, hydrochloride, aniline, and alcohol at 165° (Fischer a. Hepp, A. 256, 247). Deep-red bronzed plates. Conc. HClAq forms aniline and naphthorosindone

DI - PHENYL - (a)-NAPHTHYL-ACETAMID-INE C₁₀H₂.CH₂.C(NPh).NHPh. [130°]. Formed from (a)-naphthyl-acetic acid (3 mols.), aniline (6 mols.), and PCl_s (Boessneck, B. 16, 642).

Needles, sol. ether and benzene.

PHENYL-(a)-NAPHTHYL-AMINE C16H18N i.e. C_eH_s.NH.O₁₀H₇. [60°]. Forms by heating (a)-naphthylamine hydrochloride with aniline at 250° (Girard a. Vogt, Bl. [2] 18, 67; Streiff, B. 13, 1852). Prepared by heating a mixture of (a)-naphthol (15 pts.), aniline (19.4 pts.) and CaCl₂ (11.6 pts.) under pressure for 9 hours at 280°; the yield being 26 p.c. of the theoretical (Friedländer, B. 16, 2077). White plates or prisms, v. sol. alcohol. Its solutions show blue fluorescence. Yields a tri-bromo- derivative [137°] and a di-nitro- derivative [77°]. H₂SO, at 100° forms a tetra-sulphonic acid.—B'HCl: prisms, decomposed by water.-B'C,H,N,O,.

Acetyl derivative C₁₈H₁₈NO. [115°].

Benzoyl derivative. [152°].

Nitrosamine. C₁₈H₁₈NPh.NO. [92°].

Reddish-yellow crystals (Fischer a. Hepp, B.20,

Phenyl-(β)-naphthylamine $C_{10}H_{18}N$. [108°] (395°). Formed by heating (B)-naphthol with aniline and ZnCl, at 190° (Merz a. Weith, B.13, 1300), or with aniline hydrochloride (Merz a. Weith, B. 13, 1850).

Preparation.—A mixture of (β) -naphthol (15) pts.), aniline (19.4 pts.), and CaCl, (11.6 pts.) is heated under pressure at 280° for 9 hours; the yield is 98 p.c. of the theoretical (Friedländer, B. 16, 2075).

Properties.—White needles, sol. Yields a tetra-bromo- derivative [198°] and a nitro- derivative [87°]. With nitroso-di-methylaniline hydrochloride and HOAc it yields lustrous black crystals, which form a violet solution in H₂SO₄ (Witt, B. 21, 723). Sulphur at 240° forms SC₀H₅NH [178°] (Kym, B. 23, 2466).

H2SO4 at 100° forms a trisulphonic acid

Salts .- B'HCl: unstable crystalline powder. -B'C₆H₃N₃O₇. Brownish needles, v. sol. CHCl₂. Acetyl derivative. [93°]. Crystals.

Bensoyl derivative. [136°] (Streiff, A. 209, 151); [148°] (Claus a. Richter, B. 17, 1591).
Nitrosamine C., H., NPh.NO. [98°].

Di-phenyl-naphthylamine C₁₀H, NPh₂. [142°]. (335°-340° at 85 mm.). Formed by adding abromo-naphthalene (20 g.) to a boiling solution of potassium (3.3 g.) in diphenylamine (20 g.) mixed with aniline (15 c.c.) (Herz, B. 23, 2541). Silky needles (from dilute alcohol), insol. water. References .- NITRO-AMIDO-, DI-NITRO-, and NITROSO- PHENYL-NAPHTHYL-AMINE.

PHENYL-(a)-NAPHTHYLAMINE BLUE v. TRI - NAPHTHYL - TRI - AMIDO-TRI-PHENYL-CARBINYL CHLORIDE.

PHENYL - (β) - NAPHTHYL - CARBAMIC ACID.

EthylC₁₀H₇NPh.CO₂Et. etherFormed by the action of NaOEt on the chloride (Paschkowezky, B. 24, 2919). Satiny needles.

Phenyl ether C10H,NPh.CO2Ph. [149°]. S. (96 p.c. alcohol) ·36 at 17°; S. (benzene) 2·5 at 17°. Formed from the chloride and NaOPh. Needles, v. sl. sol. cold alcohol.

Chloride C₁₀H₇NPh.COCl. [102°]. Formed from phenyl-(B)-naphylamine and COCl₂ in toluene (Kym, B. 23, 425). White plates.

PHENYL-(a)-NAPHTHYL CARBINOL Ph.CH(OH).C₁₀H₇. [86.5°]. (above 360°). Formed from the ketone, zinc-dust, and KOH (Elbs, J. pr. [2] 35, 504; cf. Lehine, B. 13, 359; Beckmann, B. 22, 915). Crystals, v. sol. alcohol. Gives a violet colour with H,SO.

Phenyl-di-(a)-naphthyl carbinol C27H200 i.e. Ph.C($C_{10}H_{7}$)₂OH. [160°-170°]. Formed, together with benzoic aldehyde, by boiling phenyl-(α)-naphthyl-(β)-pinacolin ($C_{10}H_{7}$)₂CPhBz (got by reducing phenyl (a)-naphthyl ketone with zinc and alcoholic HCl) with alcoholic KOH (Elbs, J. pr. [2] 35, 507). Grey crystalline crusts (from ether-alcohol), m. sol. alcohol, v. sol. ether.

PHENYL - (8) - NAPHTHYL - CARBINYL - THIO-UREA. Tetrahydride C₁₈H₂₀N₂S i.e. NHPh.CS.NH.CH₂C₁₀H₁₁. [140°]. Formed from C₁₀H₁₁.CH₂NH₂ and phenyl thiocarbimide (Bamberger a. Helwig, B. 22, 1913). Vitreous rosettes.

PHENYL-NAPHTHYL-CARBINYL-UREA. NHPh.CO.NH.CH2C10H11. Tetrahydride Formed from naphthyl-carbinylamine phenyl cyanate (Bamberger, B. 22, 1913). The (a)-compound melts at 126.5° , the (β)-isomeride at 141°. Both crystallise in needles.

PHENYL-(a)-NAPHTHYLENE-DIAMINE $C_{16}H_{14}N_2$ i.e. $C_{10}H_6(NH_2)(NHPh)$ [1:4]. [148°] Formed by reducing nitroso-phenyl-(a)-naphthylamine (Wacker, A. 243, 305). Needles (from

alcohol) or plates (from benzene).

Phenyl-o-naphthylene-diamine C₁₀H_e(NH₂)(NHC_eH_e) [1:2]. Amido-(β)-naph-thyl-phenyl-amine. [140°]. Formed, together with aniline, by reduction of benzene-azo-(β) naphthyl-phenyl-amine with SnCl₂ (Zincke a. Lawson, B. 20, 1170; Witt, B. 20, 1184). Broad needles or plates. Nitrous acid passed into HOAc (11 pts.) containing the hydrochloride in suspension forms $C_{1e}H_{s}N_{\star}O_{2}$ or $C_{1e}H_{1e}N_{\star}O_{2}$ [208°], whence SnCl₂ forms $C_{1e}H_{12}N_{\star}$ [194°], which yields B'HCl, $C_{1e}H_{1,\star}AcN_{\star}$ [261°] and $C_{1e}H_{1e}Ac_{2}N_{\star}$ [177°], and gives with benzoic aldehyde a compound C25H16N4 [189°] (Zincke a. Campbell, A. 255, 849).

Salts.—B'HCl: long colourless glistening needles, v. sl. sol. water, more readily in alcohol.

—×B'H₂SO₄: needles, similar solubility.

Phenyl-dinaphthylene-amine CaeH1, N i.s. C₁₀H NPh. [c. 144°]. Formed by heating (\$\$)-di-oxy-dinaphthyl with aniline-zinc-ohloride at 300° (Walder, B. 15, 2175). needles, sol.alcohol.— $B'2C_6H_3N_4O_7$. Prisms or [169°].

Di-phenyl-naphthylene-diamine

[2:2'] $\mathring{C}_{10}H_e(NH\mathring{P}h)_2$. [168°]. Formed, together with $C_{10}H_e(OH)(NH\mathring{P}h)$ [163°], by heating dioxynaphthalene with aniline and CaCl2 (Annaheim, B. 20, 1372; Clausius, B. 23, 528). Silvery plates (from hot benzene).

Acetyl derivative C28H22N2O2. [198°].

 ${f Di-phenyl-naphthylene-p-diamine}$ [1:4] $C_{10}H_6(NHPh)_2$. [144°]. Formed by reducing the di-anilide of (a)-naphthoquinone with zinc, HOAc, and alcohol (Fischer a. Hepp, A. 256, 255). Colourless prisms.

DI - PHENYL - NAPHTHYLENE - DI-THIO-UREA C₁₀H₆(NH.CS.NHPh)₂. [355°-360°]. Formed from naphthylene (1,2)-diamine and phenyl-thio-carbimide (Bamberger a. Schieffelin,

B. 22, 1376). Needles.

DÍ-PHÉNYL-NAPHTHYLENE-UREA

C₁₀H₆(NH.CO.NHPh)₂. [335°]. Formed from naphthylene (1,2)-diamine and phenyl cyanate (Bamberger a. Schieffelin, B. 22, 1376). Crystalline granules, v. sl. sol. benzene.

PHENYL-NAPHTHYL-ETHANE C₁₈H₁₆ i.e.

C₁₀H₇.CH₂.CH₂Ph. Formed by heating naphthyl benzyl ketone with HI and P at 155° (Graebe a. Bungener, B. 12, 1078). Yields chrysene on passing through a red-hot tube.

s-PHENYL-(B) NAPHTHYL-ETHYLENE

C₆H₃.C₂H₂.C₁₀H₇. [145°]. Formed by distilling (β)-naphthyl cinnamate, CO₂ being split off (Anschütz, B. 18, 1946). Silvery plates.

Di-bromide. [192°]. PHENYL-NAPHTHYL-ETHYLENE-\(\psi\)-THIO-UREA NPh:C< $^{N(C_{10}H_7)}_{S.CH_2}>$ CH₂. [185°]. Formed from phenyl-(a)-naphthyl-urea and ethylene bromide (Foerster, B. 21, 1870).

DI-PHENYL-NAPHTHYL-GUANIDINE $C_{23}H_{19}N_3CN_3H_2Ph_2(C_{10}H_7)$. [155°]. Formed by heating di-phenyl-thio-urea with (a)-naphthyl-Formed by amine, alcohol, and PbO (Tiemann, B. 3, 6; Z.

[2] 6, 309). Crystalline, sol. alcohol. PHENYL - (β) - NAPHTHYL - GUANIDINE GARBOXYLIC ACID

C10H2NH.C(NH).NH.C6H4.CO2H. Formed by heating cyan-carbimidamido-benzoic acid with (B)naphthylamine (Griess, B. 16, 338). Crystals

apinitylamine (Griess, B. 10, 536). Crystais sl. sol. hot alcohol.—B'HCl. Six-sided plates. PHENYL (a)-NAPHTHYL-KETONE C₁₇H₁₂O i.e. C_eH₅·CO.C₁₆H₇. Mol. w. 232. [75·5°]. (385°) (Schweizer, A. 264, 196). V.D. 119 (calc. 116). S. (alcohol) 2·5 at 12°.

Formation.— 1. Together with the (β)-

isomeride, by heating naphthalene with HOBz and P₂O₅ at 210° (Merz, B. 6, 541, 966, 1238).— 2. By heating (a)-naphthoic acid with benzene and P_2O_5 (M.).—3. Together with a smaller quantity of the (β) - isomeride, by the action of AlCl, on a mixture of naphthalene, CS₂, and BzCl. The two ketones are separated by crystallisation from alcohol-ether (Elbs, J. pr. [2] 35, 503; B. 19, 1965; Rospendowski, C. R. 102, 872).—4. By heating naphthalene with BzCl and ZnCl2, or Zn (Roux, A. Ch. [6] 12, 338; Kegel, A. 247, 178).

Properties.—Trimetric prisms, very slightly

volatile with steam.

Reactions.—1. Br forms C₁₇H₁₁BrO [98°] (E.); [100.5°] (Rospendowski), whence HNO₄

gives C₁₇H₀Br(NO₂)₂O decomposing at 90°.—2. Conc. H2SO, with a few drops of water forms at 100°-150° benzoic acid and naphthalene (B)sulphonic acid.—3. Soda-lime at 350° forms naphthalene and NaOBz. - 4. Chromic acid forms C₈H₅.CO.C₆H₃(CO₂H)₂[1:2:3]. mixture [155°].—5. Sodium acting on its ethereal soluforms a greenish-yellow compound 2.CPh)₂O₂Na₂, whence water produces tion (C₁₀H₁.CPh)₂O₂Na₂, whence water produces CPh(C₁₀H₁)₂.CO.C₆H₅ and other bodies, while CO₂ acting on the Na compound forms a yellow powder, split up by water into ketone and C₆H₅.C(C₁₀H₅)(OH).CO₂H [148°] (Beckmann a. Paul, A. 266, 10).

 $Oxim C_6H_5.C(NOH).C_{10}H_7.$ [142°]. Groups of white needles (from dilute alcohol) (Kegel, A. Oxidised by CrO₃ in HOAc to **247**, 181). C_8H_8 .CO. C_8H_8
CO.CH [152°], which yields an

anilide $C_0H_2Bz < CO.C.NHPh [200°]$ and a ptoluide [197°] both converted by boiling NaOHAq into $C_0H_3Bz < \stackrel{CO.C.OH}{CO.CH}$ [222°].

Phenyl (β)-naphthyl ketone [82°]. S. 2 at 12°. Formed, at the same time as the (a)- isomeride, from naphthalene (v. supra). Formed also by heating (β) -naphthoic acid with benzene and P₂O₅ (Merz). Needles, v. sol. hot alcohol. Oxim. [176°]. Needles. Oxidised by CrO₃

in HOAc to yellow C, H, Bz CO.CH, [132°], which is converted by heating with aniline into dark-red plates of $C_6H_3Bz < CO.C.NHPh$ [210°].

Reference. — DI - OXY - PHENYL - NAPHTHYL -KETONE

PHENYL NAPHTHYL KETONE o-CARB-OXYLIC ACID C₁₀H₇.CO.C₆H₄.CO₂H. The chloride is formed from naphthalene, phthalic anhydride, and AlCl_s (Ador a. Crafts, Bl. [2] 34, 531). Prisms (from dilute alcohol). PHENYL - NAPHTHYL - METHANE

BENZYL-NAPHTHALENE.

Phenyl-di-(a)-naphthyl-methane C_2 : i.e. C_0H_5 CH($C_{10}H_7$). [c. 180°]. Formed from phenyl-(a) naphthyl-(b)-pinacolin by distilling with zinc-dust. Got also by heating phenyl-di-(a)-naphthyl-carbinol with zinc-dust (Elbs, J. pr. [2] 35, 508). Grey powder; cakes together at 100°

Di-phenyl-naphthyl-methane C10H7.CHPh7 [134° and 149°]. Formed by heating di-phenylcarbinol with naphthalene and P.O. at 140° (Lehne, B. 13, 358; Hemilian, Bl. [2] 34, 326). Needles, sol. benzene and ether, sl. sol. alcohol.

PHENYL - (a) - NAPHTHYL - METHYL-PYR-ROLE CH:CPh NC₁₀H₇. [74°]. (above 360°). Formed by heating the carboxylic acid [244°] (Lederer a. Paal, B. 18, 2598). Plates. V. e. sol. alcohol, benzene, and ligroin.

Phenyl-(\$\beta\$)-naphthyl-methyl-pyrrole

CH:CPh NC₁₀H₇. [52°]. Formed by heating the carboxylic acid [249°] (L. a. P.). white concentric needles. V. sol. alcohol Small

PRENYL-(a)- NAPHTHYL-METHYL-PYR-BOLE CARBOXYLIC ACID

CO₂H.C=CMe>NC₁₀H₇. [244°]. Formed by

heating acetophenoneacetoacetic ether with (a)naphthylamine at 130°, and saponification of the product (Lederer a. Paal, B. 18, 2598). Needles. V. sol. alcohol, benzene, and acetic acid.

Phenyl- β -naphthyl-methyl-pyrrole oxylic acid. [249°]. Formed in like manner, using (β) -naphthylamine (L. a. P.). Small white needles, sol. alcohol and HOAc.

Ethyl ether EtA'. [115°]. Plates.

PHENYL - (B)-NAPHTHYL-METHYL-THIO-UREA C₁₀H₇NH.CS.NPhMe. [127°]. Got from (β)-naphthyl-thiocarbimide and methyl-aniline (Gebhardt, B. 17, 2091). Long yellow needles.

Phenyl-(α)-naphthyl-methyl-ψ-thio-urea $C_{10}H_*N:C(SMe).NHPh.$ [96°]. Formed from phenyl (a)-naphthyl-thio-urea and MeI (Foerster, 1870). Small white needles. (a)-naphthyl-thiocarbimide on heating with CS

PHENYL - (α) - NAPHTHYL-(β)-PINACOLIN $C_6H_5.C(C_{10}H_7)_2.CO.C_6H_5.$ [c. 130°]. From phenyl-(a)-naphthyl ketone by boiling with zinc and HCl (Elbs, J. pr. [2] 35, 505). Pale greenishyellow crystalline crusts, v. sol. ether, sl. sol. alcohol, insol. water. Cakes together at 100°-110°. Boiling alcoholic KOH forms benzoic aldehyde and phenyl-di-naphthyl-carbinol.

PHENYL - (a) - NAPHTHYL - PINACONE C₁₀H₂CPh(OH).CPh(OH).C₁₀H₂. [61°]. A product of the action of sodium-amalgam on an alcoholic solution of phenyl-(a)-naphthyl ketone (Lehne, B. 13, 1360). Needles (from ether)

 $DI - PHENYL - (\alpha) - NAPHTHYL - PYRROLE$ CH:CPh NC₁₀H, [149°]. Formed by heating its carboxylic acid with lime (Paal a. Braikoff, B. 22, 3092). Needles, v. sol. hot alcohol.

Di - phenyl - (β) - naphthyl - pyrrole. [208°]. Formed in like manner (P. a. B.). Needles.

DI - PHENYL - (a) - NAPHTHYL - PYRROLE CARBOXYLIC ACID CO2H.C=CPh NC10H, [272°]. Got by saponifying its ether, which is obtained from phenacyl-benzoyl-acetic ether and (a)-naphthylamine (Paal a. Braikoff, B. 22, 3091). Small white plates, sl. sol. hot alcohol.—KA'. Sl. sol. hot water.

Ethyl ether EtA'. [182°]. Needles.

Di-phenyl-(β)-naphthyl-pyrrole carboxylic acid. [above 350°]. Got in like manner, using (\$\beta\$)-naphthylamine. Plates.—KA'. Sl. sol. water.

Ethyl ether EtA'. [182°]. Needles. PHENYL-(α)-NAPHTHYL SULPHIDE $C_{10}H_7.SPh.$ [42°] (K. a. B.); [49°]. (c. 215° at 15 mm.). Formed from Pb(SPh)₂ and $C_{10}H_7Br$ (Krafft a. Bourgeois, B. 23, 3047). Formed also from (a)-diazonaphthalene chloride and NaSPh (Ziegler, B. 23, 2471). Prisms (from dil. alcohol).

Phenyl (β)-naphthyl sulphide. [52°]. (c. 224°

at 14 mm.). Formed in like manner. Needles.
PHENYL (a) - NAPHTHYL SULPHONE SULPHONE C.H.SO.C.OH. [100°]. Formed by oxidising phenyl (a)-naphthyl sulphide with CrO, and HOAc (Krafft a. Bourgeois, B. 23, 3047). Formed also, together with the (β) - isomeride, by heating benzene sulphonic acid with naphthalene and P₂O₅ at 175° (Michael a. Adair, B. 10, 585). Crystals (from alcohol).

Phenyl (β) - naphthyl sulphone. [116]. Formed in like manner, and also by heating naphthalena (β)-sulphonic acid with benzene and P.O. (M. s. A.), and by the action of zinc-

dust or AlCl, on a mixture of naphthalene and benzene sulphonic chloride (Crustschoff, B. 7, 1167; Otto a. Beckurts, B. 11, 2069). Needles.

PHENYL-(a)-NAPHTHYL-THIO-SEMICARB. AZIDE NHPh.CS.NH.NHC, H,. [185°]. Formed from (a)-naphthyl-hydrazine and phenyl-thiocarbimide (Preund, B. 24, 4191). Needles.

Phenyl - (β) - naphthyl - thio - semicarbaside. Formed in like manner from (B)-

naphthyl-hydrazine (P.). White plates.
PHENYL - (a) - NAPHTHYL - THIO - UREA NHPh.CS.NHC₁₀H, [163°]. Formed from (α)-naphthylamine and phenyl-thiocarbimide (Hofmann, Pr. 9, 274), and also from aniline and (a)-naphthyl-thiocarbimide (Mainzer, B. 15, 1414). Plates, v. sl. sol. alcohol. With ethylene bromide it gives two bases [185°] and [180°].

NHPh.CS.NHC, H, Tetrahydride[153°]. Formed from phenyl-thiocarbimide and (a)-naphthylamine tetrahydride (Bamberger, B. 21, 1794). Prisms, v. sol. benzene-alcohol.

Phenyl-(B)-naphthyl-thio-urea NHPh.CS.NHC₁₀H₇. [165°]. Formed from (β)-naphthylamine and phenyl thiocarbimide
 (M.; Freund a. Wolf, B. 25, 1468). Plates. Split up by HClAq at 150° into aniline, (3)naphthylamine, and phenyl and (β)-naphthyl COCl₂ in thiocarbimides. toluene forms NPh:C<S-U-107 CO [117°].

Tetrahydride NHPh.CS.NHC10H11. [161°]. Formed from phenyl thiocarbimide and (B)naphthylamine tetrahydride (Bamberger Müller, B. 21, 858). Prisms (from alcohol). PHENYL-(a)-NAPHTHYL-UREA. Tetr

Tetrahydride NHPh.CO.NHC₁₀H₁₁. [193°]. Formed from (a)-naphthylamine tetrahydride and phenyl cyanate (Bamberger, B. 21, 1794). Needles.

u-Phenyl- (β) -naphthyl-urea NH₂.CO.NPh.C₁₀H₇. [190°]. Formed by heating NPh(C₁₀H₁).COCl and alcoholic NH, at 130 (Kym, B. 23, 426). Needles, sl. sol. cold alcohol.

Phenyl- (β) -naphthyl-urea NHPh.CO.NHC, H,. [221°]. Formed from phenyl cyanate and (β)-naphthylamine (Goldschmidt, B. 21, 2567). Prisms (from alcohol).

Tetrahydride NHPh.CO.NHC, H₁₁, [166°]. Formed from phenyl cyanate and the tetrahydride of (B)-naphthylamine (Bamberger a. Müller, B. 21, 859). Needles, v. e. sol. alcohol.

Phenyl-di-(B)-naphthyl-urea C27H20N2O i.e. NHPh.CO.N(C₁₀H₁)_r [182°]. Formed from phenyl cyanate and di-(\$\beta\$)-naphthylamine (Gebhardt, B. 17, 3039). Formed also by heating N(C₁₀H₂)₂.COCl with aniline in CHCl₂ at 130° (Kym, B. 23, 429; Kühn a. Landau, B. 23, 811). Needles (from alcohol). By heating with aniline it is converted into CO(NHPh), and di-(8)naphthylamine.

Phenyl-tri-(\beta)-naphthyl-urea CO(NPhC₁₀H₁).N(C₁₀H₁)₂. [168°]. S. (alcohol) 9 at 16°; S. (benzene) 4.54 at 16°. Formed from (C₁₀H₂).N.COCl and phenyl-(\$\theta\$)-naphthylamine at 260° (Paschkowezky, B. 24, 2924). Granular crystals, sl. sol. cold alcohol.

Di-phenyl-(B)-naphthyl-urea

NHPh.CO.NPhC, H, [183°]. Formed from NPh(C₁₀H₂).COCl and aniline (Kym, B. 23, 426). White plates, sl. sol. cold alcohol.

u-Di-phenyl-di- (β) -naphthyl-urea NPh2.CO.N(C10H7)2 [104°]. Formed by heating NPh₂.COCl with (C₁₀H₇)₂NH at 220° or by heating (C₁₀H₇)₂N.COCl with NPh₂H at 260° (Paschkowezky, B. 24, 2923). Crystalline powder, m. sol. cold alcohol.

s-Di-phenyl-di- (β) -naphthyl-urea

CO(NPh.C₁₀H_r)₂, [186°]. S. (alcohol) 1 at 18·5°. S. (benzene) 59 at 18·5°. Formed by heating C₁₀H_rNPh.COCl with C₁₀H_rNPh.COCl at 250° (P.). Polyhedral granules. Converted by HClAq at 250° into aniline, (B)-naphthol, and CO2.

Tri-phenyl- (β) -naphthyl-urea NPh₂.CO.NPh.(C₁₀H₂). [128°]. Formed from C₁₀H,NPh.COCl and NPh₂H at 240° (P.). Crys-Formed from

talline powder, v. sol. alcohol.

PHÉNYL O-NITRO-BENZYL KETONE C₆H₈.CO.CH₂.C₆H₄NO₂. Nitro-deoxybenzoin. Formed, together with the p-isomeride, by nitrating phenyl benzyl ketone (Ney, B. 21, 2448). Sol. ether (difference from p-isomeride). p- isomeride yields a crystalline oxim [107°].

PHENYL p-NITRO-BENZYL OXIDE C₀H₅.O.CH₂.C₆H₄NO₂. [91°]. Formed C.H.O.CH.C.H.NO. [91°]. Formed from p-nitro-benzyl chloride and KOPh (Kumpf, B. 17, 1076). Plates (from alcohol).

Reference.—Nitro - Phenyl - Nitro - Benzyl

OXIDE.

PHENYL NITRODIPHENYLETHYL KE-TONE C₆H₅.CO.CHPh.CH₂.C₉H₄NO₂. Formed from nitro-benzyl chloride, deoxybenzoïn, and NaOEt (Buddeberg, B. 23, 2071). The o-compound melts at 102° and yields diphenylquinoline [96°] (420°) on reduction. The p- isomeride melts at 112° and yields an amido-compound [141°] on reduction.

DI-PHENYL-DINITROSACYL. So-called. C_{1e}H_{1e}N₂O₄. [87°]. The chief product of the action of HNO₃ (S. G. 1·4) on acetophenone (Hollemann, B. 21, 860, 2835). It is accompanied by an isomeride [179°]. Crystals (from ether). Converted by acids and alkalis into benzoic and oxalic acids, NH, and hydroxyl-[149°]. Ac₂O yields C₁₅H₁₀Ac₂N₂O₅ amine. Aniline and benzanilide form compounds melting at 205° and 160° respectively.

DI-PHENYL-NITROSAMINE v. Nitrosamine of DI-PHENYL-AMINE.

PHENYL-NITRO-TOLYL-THIO-UREA

NITRO-PHENYL-TOLYL-THIO-UREA. DI-PHENYL-NITRO-p-TOLYL-UREA

 $NPh_2.CO.NHC_0H_3Me(NO_2)$ [1:4:3]. [139·5°]. Formed from nitro-p-toluidine and NPh2.COCl at 125° (Lellmann a. Bonhöffer, B. 20, 2121). Yellow needles, v. sol. chloroform and benzene.

 $\begin{array}{cccc} \textbf{PHENYL-OCTINOIC} & \textbf{ACID} & \textbf{C}_{14}\textbf{H}_{16}\textbf{O}_{2} & \textit{i.e.} \\ \textbf{PhC(C}_{2}\textbf{H}_{2})(\textbf{C}_{2}\textbf{H}_{2}\textbf{Et}).\textbf{CO}_{2}\textbf{H} & (310^{\circ}\text{-}320^{\circ}). & \textbf{This} \end{array}$ appears to be one of the products of the action of CO at 170° on a mixture of NaOEt and PhCH₂.CO₂Na (M. Schroeder, A. 221, 46).

PHENYL-OCTOIC ACID. Nitrile.

C.H. CHPh.CN. (287°). Formed from phenylacetonitrile, hexyl iodide, and NaOH (Rossolymo, B. 22, 1237). Oil.

DI-PHFNYL-n-OCTYL TRICYANIDE $C_tH_{1r}.C_sN_sPh_2$. [43°]. (285° at 15 mm.). Formed from $C_sH_{1r}.COCl$, benzonitrile, and AlCl₃ (Krafft a. Koenig, B. 23, 2384).

PHENYL-OCTYL-THIO-UREA C15H24N2S i.e. C₈H₁₇NH.CS.NHPh. [53°]. Formed from aniline and sec-octyl-thiocarbimide (Jahn, B. 8, 804).

y-PHENYL-OSOTRIAZOLE C.H.N. CH:N>NPh. [224°]. Formed by warming the osotetrazone of glyoxal with FeCl, and HClAq (Pechmann, A. 262, 291). Sl. sol. water, v. sol. alcohol. HNO, yields $\stackrel{C(NO_2):N}{CH=N}$ NPh [184°].

Tri-phenyl-osotriazole CPh:N NPh. [122]. Formed by heating the diphenyl-dihydrazide of benzil with alcohol at 210° (Auwers a. V. Meyer,

B. 21, 2806). White plates.

PHENYL-OSOTRIAZOLE CARBOXYLIC ACID CO₂H.Ç=N CH:N>NPh. [192°]. Formed by oxidising phenyl-methyl-osotriazole with KMnO, (Von Pechmann, B. 21, 2760). Formed also by boiling the phenyl-hydrazide of di-nitroso-acetone with NaOHAq (Pechmann, A. 262, 283). White needles, sol. alcohol, not volatile with steam. Yields a nitro-compound [236°] which may be reduced to an amido-compound [252°].

Salts.—KA' aq.—BaA', 4aq.—CdA', 4aq.

Methyl ether Med. [90°]. (286°). Ethyl ether EtA'. [59°]. (306°). Amide C.N.HPh.CONH₂. [148.5°]. Formed by boiling the phenyl-hydrazide of acetyl-dinitroso-acetone with Na2CO3 Aq (P.). Needles.

Nitrile C2HN3Ph.CN. [94.56]. 60 mm.). Formed by the action of NaOH on the phenyl-hydrazide of di-nitroso-acetone (Pechmann, A. 262, 297). Plates, v. sol. ether. H₂S and alcoholic NH, convert it into the thio-amide C₂HN₂Ph.CS.NH₂ [132°]

Phenyl-osotriazole dicarboxylic acid

C(CO₂H):N NPh. [256°]. Formed by oxidising phenyl-di-methyl-osotriazole with KMnO4 (Pechmann, A. 262, 311). Crystalline powder. After fusion it melts at 184°.—CaA": needles.—Ag₂A".

PHENYL-OSOTRIAZOLE CARBOXYLIC ALDEHYDE CH NNPh. [70°]. Formed by boiling its oxim with dilute H2SO4 (Pechmann, A. 262, 294). Insol. cold water.

Oxim C₂HPhN₃.CH:NOH. [115°]. A product of the action of alkalis on the compound

CH(NOH).C(N2HPh).CH(NOAc).

Phenyl-hydraside C2HPhN2.CH:N2HPh. [118°-140°]. Plates, v. sol. alcohol.

PHENYL-OSOTRIAZYL ALCOHOL

CH₂(OH).Q=N NPh. [67°]. Formed, together with the carboxylic acid, by warming the aldehyde with conc. NaOHAq (Pechmann, A. 262, 296). Prisms, v. e. sol. alcohol, sl. sol. cold Aq.

PHENYL-OSOTRIAZYL-AMINE C.H. i.e. $CH_2(NH_2).C=N > NPh.$ (233° at 100 mm.).

Formed from C2HPhN3.CS.NH2 by treatment in alcoholic solution with Zn and HCl (Pechmann, A. 262, 300). — B'HCl. [229°]. Tables. -B'.H.PtCl. Pri [123°]. Tables. Prisms. - Di-thio-carbamate.

PHENYL-OXALACETIC ETHER C,4H,6O, i.e. CO₂Et.CO.CHPh.CO₂Et. Formed by the action of Na on a mixture of oxalic ether and phenylacetic ether (Wislicenus, B. 20, 591; A. 246, 339). Oil, decomposed by distillation, v. sol. alkalis. FeCl, colours its alcoholic solution red. Boiling dilute H2SO4 gives CO2 and phenyl-pyruvic acid [155°].

Phenyl-hydrazide

CO₂Et.C(N₂HPh).CHPh.CO₂Et. [70°]. Needles

(from dilute alcohol), v. sol. ether.

Ethyl ether of the semi-nitrile
CN.CHPh.CO.CO₂Et. [130°]. Formed Formed from phenylacetonitrile, oxalic ether, and Na or NaOEt (Erlenmeyer, jun., B. 22, 1483). Plates. FeCl, colours its alcoholic solution green.

PHENYL-OXAMIC ACID v. OXALIC ACID. PHENYL-OXAMIDE v. OXALIC ACID. PHENYL-OXAMIDE CARBOXYLIC ACID v.

OXALIC ACID

PHENYL-OXANTHRANOL $C_{20}H_{14}O_{2}$ $C_6H_4 < CO^{CPh(OH)} > C_6H_4.$ [208°]. Formed by oxidation of phenyl-anthranol with K2Cr2O, and HOAc (Baeyer, A. 202, 58). Colourless tables, insol. water, sol. alcohol. Conc. H₂SO₄ forms a purple solution. Reduced by zinc-dust and HOAc to phenyl-anthranol. Benzene and H₂SO, form crystalline C₂₈H₁₈O.

Acetyl derivative C20H18AcO2. [196°]. Reference. - DI-CHLORO- and OXY-PHENYL-OXANTHRANOL.

PHENYL-OXAZOLE C.H.NO i.e.

 $CPh \stackrel{CH.O}{\leqslant}_{N} CH$. [6°]. (221°). Formed by heating bromo-acetophenone with formamide at 140° (Lewy, B. 20, 2578; Blümlein, B. 17, 2580).-B'HČl. [80°].—B'2H2PtCl6 2aq: yellow needles.

Di-phenyl-oxazole CPh CH.O N :CPh. (339°). Formed by heating ω-bromo-aceto-phenone with benzamide at 145° (B.; L.). Plates (from alcohol).—B'HCl. Needles.

CH≪CPh.O. Phenyl-iso-oxazole Formed from the oxim of benzoyl-acetic aldehyde and AcCl (Claisen a. Stock, \vec{B} . 24, 134). PHENYL-OXAZOLE DIHYDRIDE $C_{\nu}H_{\nu}NO$

i.e. $CH_2 < \stackrel{CH_2.O}{N} = CPh$. (243°). Formed by warming bromo-ethyl-benzamide with alkalis (Gabriel a. Heymann, B. 23, 2495). Liquid, smelling like phenyl-thiazole dihydride. Miscible with alcohol and ether, sl. sol. water. HBrAq forms bromoethyl - benzamide and CH2(OBz).CH2NH2. B'₂H₂PtCl₈.—B'₂H₂Cr₂O₇.—B'C₈H₃N₃O₇. [177°]. **DI-PHENYL OXIDE** C₁₂H₁₀O i.e. Ph₂O. Mol.

w. 170. [28°]. (253°). $\mu_A = 1.5675$ at 25°.

Formation.—1. By distilling cupric benzoate (List a. Limpricht, A. 90, 190).—2. By warming diazobenzene sulphate or chloride with phenol (Hofmeister, B. 3, 747; A. 159, 204; Hirsch, B. 23, 3709).—3. By heating phenol with ZnCl₂ at 350°, the yield being 6 p.c. (Merz a. Weith, B. 14, 187).—4. By heating phenol with AlCl. (M. s. W.).—5. Together with diphenylene ketone oxide and a body melting at 111°, by distilling sodium salicylate with phenyl phosphate (R. Richter, J. pr. [2] 28, 273).

Preparation.—By distilling Al(OPh)_s (Gladstone a. Tribe, C. J. 41, 6).

Properties. - Prisms (from cold alcohol), smelling like geraniums, almost insol. water and KOHAq. Conc. H₂SO₄ forms O(C₆H₄.SO₂H)₂, which forms Na₂A" xaq, BaA", and Ag₂A".

References .- DI-AMIDO-, DI-BROMO-, and DI-

MITRO- and DI-OXY- DIPHENYL OXIDE.

PHENYL-OXY-ACETAMIDINE v. MANDEL-AMIDINE.

PHENYL-OXY-ACETIC ACID v. MANDELIC ACID.

DI-PHENYL-OXY-ANGELIC ACID v. Di. hydro-cornicularic acid.

PHENYL p-0XY-BENZYL KETONE
C.H., CO.CH., C.H., OH. [129°]. Formed from
CH.Bz.C., H., NH. by the diazo-reaction (Ney, B.
21, 2449). White spangles (from water).

Acetyl derivative. [87°]. Plates.

Isomeride v. Benzoïn.

PHENYL-OXY-BENZYL-UREA C14H14N2O2 i.e.[1:2]C₆H₄(OH).CH₂.NH.CO.NHPh. Formed by warming CeH4(OH).CH2NH2 with phenyl cyanate and benzene. Needles, sol. alcohol and ether.

Methyl derivative [1:2] C₆H₄(OM₆).CH₂.NH.CO.NHPh. [145°]. Formed from C₆H₄(OMe).CH₂NH₂ and phenyl cyanate (Goldschmidt a. Ernst, B. 23, 2743)

PHENYL-OXY-BROMO-PROPIONIC ACID v. BROMO-OXY-PHENYL-PROPIONIC ACID.

PHENYL OXYBUTYL KETONE C₁₁H₁,O₂ i.e. C₄H₃.CO.CH₂.CH₂.CH₂.CH₂OH. [41°]. Formed by boiling the anhydride of its carboxylic acid with water (Perkin, jun., C. J. 51, 733; 57, 310). Iridescent plates, v. sol. alcohol. If left to stand over H,SO, it is converted into the anhydride.

Oxim C, H, C(NOH). C, H, OH. [57°]. Minute

plates, sl. sol. ligroin.

 $\label{eq:chi} Anhydride\ C_{11}\Pi_{12}O\ \textit{i.e.}\ CH_2 < \begin{array}{c} CH\ :CPh\\ CH_2.CH_2 \end{array} > 0.$ (250° at 720 mm.). Formed by heating its carb. oxylic acid at 200° (Perkin, B. 16, 1792; 19, 2559; C. J. 51, 730). Oil. Converted by HBrAq into phenyl bromo-butyl ketone.

PHENYL OXY-BUTYL KETONE CARB. OXYLIC ACID. Anhydride C12H12O2 i.e. $CH_{2} < \begin{array}{c} C(CO_{2}H): CPh \\ CH_{2} \xrightarrow{\quad CH_{2} \\ \quad CH_{2} \end{array} > O. \quad \textit{Benzoyl-tetramethyl-}$ ene carboxylic acid. Phenyl dehydrohexone carboxylic acid. [144°]. Formed by saponification of its ether, which is made by the action of trimethylene bromide on sodium benzoyl-acetic ether (Perkin, jun., C. J. 51, 726; B. 19, 2557). a:b:c = 2.638:1:3.398; Monoclinic crystals, a:b:c=2.638:1:3.398; $\beta=74.44'$. V. sol. alcohol and ether. Converted by cone. HBrAq in the cold into C,H,CO.C,H,Br.

CaA".—PbA": white pp.—AgA': white pp.
Ethyl ether EtA'. [60°]. Monoclir Monoclini**c** prisms; a:b:c=1.002:1:.591; $\beta=68^{\circ}34'$.

Reference.—NITBO-PHENYL OXY-BUTYL KETONE ANHYDRIDE CARBOXYLIC ACID.

PHENYL - OXY - BUTYRIC ACID v. Oxy-PHENYL-BUTYRIC ACID.

PHENYL OXY- ψ -CUMYL KETONE

C_eH_s.CO.C_eHMe_sOH. [187°]. Formed by the action of nitrous acid on phenyl amido-\(\psi\-\)-cumyl ketone (Fröhlich, B. 17, 1806). Plates, v. sol. alcohol, sol. alkalis.

PHENYL OXYETHYL SULPHONE

C₆H₃·SO₂·C₂H₄OH. Formed by heating glycolic chlorhydrin with sodium-benzene sulphinate and a little water at 120° (Otto, J. pr. [2] 30, 186). Prepared by boiling $C_2H_4(SO_2C_4H_4)_4$ with KOHAq. Liquid, sl. sol. water, miscible with alcohol and Reactions. — 1. Conc. H₂SO₄ forms ether. $C_aH_a.SO_2.C_2H_4.O.SO_3H$, which yields $3\frac{1}{2}BaA'_2aq$, crystallising from water in needles.—2. Sodiumamalgam reduces it to alcohol and Ph.SO.H or PhSH.—3. PCl, forms PhSO₂.C₂H₄Cl, crystallis.

ing from benzene in six-sided tablets [56°]. 4. PCl, forms the sparingly soluble anhydride (Ph.SO₂.C₂H₄)₂O [70°].—5. Aqueous NH₃ at 120° forms (Ph.SO₂.C₂H₄)₂NH. - 6. Chromic acid oxidises it to Ph.SO2.CH2.CO2H

Acetyl derivative C₁₀H₁₂SO₄. Oil. Benzoyl derivative Ph.SO₂.C₂H₄.OBz

[125°

PHENYL OXY - METHYL KETONE v.

BENZOYL-CARBINOL.

PHENYL - OXY - NAPHTHYL KETONE. Ethyl derivative C₁₀H₆(OEt).CO.C₆H₅. [75°]. Formed from C₁₀H₇.OEt, BzCl, and AlCl₂ (Gattermann, B. 23, 1209). Needles.

Phenyl-di-oxy-naphthyl ketone

 C_6H_3 .CO. C_6H_3 <C(OH):CH. [191°]. Formed by boiling (a)-benzoyl-naphthoquinone with SnCl₂ and alcohol (Kegel, A. 247, 183). White plates. Acetyl derivative. [155°]. Plates.

Isomeride v. DI-OXY-PHENYL-NAPHTHYL-KE-

PHENYL-OXYPHENYLBIAZYL-HYDRAZ -

INE NPh.N C.NH.NHPh. [181°]. Formed by heating di-phenyl-carbazide with COCl₂ at 100° (Freund a. Kuh, B. 23, 2831). White crystals, insol. benzene. Oxidised by FeCl₃ to C₂N₂PhO₂N:NPh [200°]. PHENYL - OXY - PHENYLTHIOBIAZYL -

HYDRAZINE NPh.N C.NH.NHPh. [124°].

Formed by the action of COCl₂ on di-phenylthiocarbazide, the resulting C₁N₂PhSO.N:NPh [140°] being reduced by alcoholic ammonium sulphide (Freund a. Kuh, B. 23, 2827). Needles.

DI-PHENYL-DIOXYPHENYLENE DI-KETONE v. DIOXYPHENYLENE DIPHENYL DI-KETONE

PHENYL-OXY-PIVALIC ACID v. β-ΟΧΥ-

PHENYL-VALERIC ACID. PHENYL-OXY-PROPIONIC ACID v. Oxy-

PHENYL-PROPIONIC ACID

DI-PHENYL OXYPROPYLENE DISUL. PHONE CH(OH)(CH2.SO2.C6H5)2. Formed by oxidation of the product of the action of NaSPh on dichlorhydrin (Otto a. Rossing, B. 23, 758). Oil, v. sol. alcohol.

Benzoyl derivative. [150°]. Needles. DI-PHENYL OXY-PROPYL SULPHONE. Anhydride (C₆H₅.SO₂.CH₂.CH₂.CH₂)₂O. [85°]. Formed by heating di-phenyl trimethylene disulphone with alcoholic potash at 120° (Otto, B. 24, 1833). Hexagonal tables, insol. water.

PHENYL OXYTOLYL KETONE. C_eH₅.CO.C_eH₈Me.OMe. [80°]. derivative Formed from C.H.Me(OMe), BzCl, and AlCl, (Koenigs a. Carl, B. 24, 3897). Crystals.

PHENYL-OXY-VALERIC ACID v. OXY-

PHENYL-VALERIC ACID.

Di-phenyl-oxy-valeric acid v. Tetrahydro-CORNICULARIC ACID.

PHENYL-PARABAMIC ACID v. PARABAMIC

PHENYL-PARACONIC ACID v. Anhydride of Oxy-BENZYL-SUCCINIC ACID and NITRO-PHENYL-PARACONIC ACID

DI-PHENYL-PENTADECYL TRICYANIDE C₃N₃Ph₂(CH₂)₁₄CH₃. [64°]. (328° at 13 mm.). Formed by heating benzonitrile with palmityl chloride and AlCl, from 40° to 100° (Krafft a.

Globular groups of Hansen, B. 22, 809). needles (from isobutyl alcohol), v. sol. ether. PHENYL PENTADECYL KETONE

 $C_6H_3.CO.C_{15}H_{31}.$ [59°]. (251° at 15 mm.). Formed from palmityl chloride, benzene, and AlCl₂ (Krafit, B. 19, 2982; 21, 2266). Plates (from alcohol), sol. ether. Yields benzoic and pentadecoic acids on oxidation.

Reference.—OXY-PHENYL PENTADECYL KETONE. PHENYL-PENTANE v. AMYL-BENZENE.

PHENYL-PENTANE DICARBOXYLIC ACID CO₂H.C₆H₄.CEt₂.CO₂H. [148°]. Formed by the action of KOHAq upon its anhydride

 $C_6H_4 < CO \stackrel{CEt_2.CO}{O}$ [53°], which is got by the action of alcoholic potash and EtI on di-oxyethyl-isoquinoline (Pulvermacher, B. 20, 2494). Crystalline.—BaA".—Ag₂A"; yellow powder. Phenyl-pentane tricarboxylic ether

(336° cor.). CHEt(CO₂Et).C(CH₂Ph)(CO₂Et)₂. $\frac{20}{4}$ 1.0899. $\mu_{\rm D} = 1.4867$ at 20°. Formed from sodium butane tricarboxylic ether and benzyl chloride (Bischoff a. Mintz, B. 23, 654). Yields, on saponification, two benzyl-ethyl-succinic acids [157.5°] and [128.5°].

Phenyl-pentane tricarboxylic acid $\mathrm{CMe}_2(\mathrm{CO}_2\mathrm{H}).\mathrm{C}(\mathrm{CH}_2\mathrm{Ph})(\mathrm{CO}_2\mathrm{H})_2.$ [178°]. Ethyl ether $\mathrm{Et}_s\mathrm{A}'''$. (337° cor.). S.G. 29 1.0950. $\mu_\mathrm{D}=1.4834$ at 20°. Formed in like manner from isobutane tricarboxylic ether (B. a. M.). Yields, on saponification, the acid and also benzyl-di-methyl-succinic acid [155°]

Phenyl-pentane tri-carboxylic acid $CHMe(CO_2H).CH_2.C(CH_2Ph)(CO_2H)_2.$ [130]. One of the acids got by saponifying the product of the action of benzyl chloride on sodium isobutane tricarboxylic ether (Bischoff, B. 23, 1947). Plates (from water).

Di-phenyl-pentane tetra-carboxylic ether $CH_2(C(CO_2Et)_2.CH_2Ph)_2$. (c. 240° at 20 mm.). Formed from disodium propane tetra-carboxylic ether and benzyl chloride (Dressel, A. 256, 191). Thick oil.

PHENYL-PENTENOIC ACID $C_{11}H_{12}O_2$ i.c. $C_6H_5.C_4H_6.CO_2H.$ Hydrocinnamenylacrylic acid. [31°]. Formed by reduction of phenylpentinoic acid by sodium-amalgam (Baeyer a. Jackson, B. 13, 122). Plates. Yields di-oxyphenyl-valeric acid on oxidation by alkaline KMnO₄ (Fittig a. Mayer, A. 268, 51).

Dibromide. [109°]. Prisms. PHENYL-PENTINOIC ACID C₁₁H₁₀O₂ i.e. C₈H₂,CH:CH.CH:CH.CO₂H. Styryl-acrylic acid. [166°]. Formed from cinnamic aldehyde, Ac₂O, and NaOAc (Perkin, C. J. 31, 403; Fittig, A. 268, 50). Prisms (from water). On oxidation by alkaline KMnO, it yields benzoic aldehyde and racemic and oxalic acids (Doebner, B. 23, 2374).—SrA'₂ 2aq.—Ag₂A''.

Chloride C₁₂H₁₈OCl. [25°].

Amide. [186°]. Tables (from alcohol). Di-phenyl-pentinoic acid

C₆H₅CH:CH.CH:CPh.CN. [119°]. Formed from cinnamic aldehyde, phenyl-acetonitrile, and NaOEt (Freund a. Immerwahr, B. 23, 2856). Needles, v. sol. alcohol and ether.

Reference.—NITRO- and OXY-PHENYL PEN-TINOIC ACID.

PHENYL-PHENTRIAZINE. Dihydride. $C_{\bullet}H_{\bullet} <_{CH_{r}NPh}^{N=N}$ [128°]. Formed by adding NaNO₂ to a solution of o-amido-benzyl-phenylhydrazine or o-amido-benzyl-aniline in HClAq at 0° (Busch, B. 25, 448). Plates, v. sol. alcohol. Salts.-B'HCl.-B'2H2PtCle. [130°].-B'C₆H₃N₃O₇. [111°]. Orange-red needles.

PHENYL-PHENTRIAZOLE C12HpN2 $C_{\mathfrak{s}}H_{\mathfrak{s}} \overset{N}{\underset{N}{\otimes}} NPh.$ [109°]. Occurs as a byproduct in the manufacture of amido-azo-benzene (Gattermann, B. 21, 1633). Formed by the action of phenyl-hydrazine on bromo-di-nitrobenzene, the resulting nitro-phenyl-phentriazole [1:4:5]C₆H₃(NO₂):N₃Ph being reduced to amidophenyl-phentriazole [183°] and NH, then eliminated (Kehrmann a. Messinger, B. 25, 899).

Isomeride C.H. NPh N. Phenyl-phenyl-[90°]. Formed from o-amidoene-azimide. diphenylamine and nitrous acid (Schöpff, B. 23, 1843). Needles, sol. alcohol.

PHENYL-PHEN-(B)-NAPHTHACRIDINE

 $C_6H_4 < N_{CPh} > C_{10}H_6$. [198° uncor.]. Formed by heating a mixture of phenyl-(3)-naphthylamine and benzoic acid with P₂O₅ or ZnCl₂ (Claus a. Richter, B. 17, 1595). Sublimes in white needles.—B',H,PtCla: glittering yellow needles.

DI-PHENYL PHENYL-ACETYLENE DI-KETONE v. DEHYDRO-ACETOPHENONE-BENZIL, vol. i. p. 37.

Di-phenyl-diphenylacetylene diketone

OXYLEPIDEN, vol. iii. p. 137.

PHENYL-PHENŸL-AMIDO-ACETIC C₆H₄Ph.NH.CH₂.CO₂H. Formed from p-amidodiphenyl and chloro-acetic acid (Zimmermann, B. 13, 1966). Plates (from hot water).—EtA'. [95°]. Needles (from dilute alcohol).

PHENYL PHENYL-AMIDO-ETHYL KE-TONE C, H, CO.C, H, NHPh. [38°]. Formed from C.H. CO.C.H. Br and aniline (Pampel a. Schmidt, B. 19, 2896). Yellow crystals.

Acetyl derivative. [103°]. Needles. PHENYL PHENYL-AMYL KETONE

 $C_{18}H_{20}O$ i.e. C_6H_5 .CO.CHPh.CH $_2$ Pr. [78°]. (330° cor.). Formed from phenyl benzyl ketone, NaOEt, and isobutyl bromide (V. Meyer a. Oelkers, B. 21, 1295). Needles. Yields an oxim [118°] crystallising in prisms.

PHENYL-PHENYL-BENZAMIDINE

 $C_{19}H_{14}N_2$ i.e. $CPh \leqslant {}^{
m NH}_{
m N} > C_6H_3Ph$. [198°]. Formed from C₆H₃Ph(NO₂).NHBz, tin, and HOAc (Hübner, A. 209, 347). Plates (from alcohol).—B'HCl. $-\mathbf{B'_{o}H_{o}PtCl_{s}}$. -B'2H2SO4: needles, sl. sol. water. PHENYL-PHENYL BENZYL KETONE

 $C_{20}H_{16}O$ i.e. C_6H_4Ph . CO.CH₂Ph. [150°]. (above 360°). Formed from diphenyl, phenyl-acetic chloride, and AlCl. (Päpcke, B. 21, 1339). Plates (from alcohol). Converted by NaOEt and CSCl2 into C.H.Ph.CO.CPh:CS [above 320°]

PHENYL PHENYL-BUTYL KETONE

C_sH_s.CO.CHPhPr. [33°]. (330°). Formed from sodium deoxybenzoin and n-propyl bromide (Bischoff, B. 22, 346). Thread-like needles (from alcohol). Yields an oxim [100°].

Phenyl phenyl-isobutyl ketone C₆H₅.CO.CHPhPr. [48°]. (325°). Yields an oxim [70°] and a phenyl-hydrazide [72°].

DI-PHÉNYL-DÎ-PHÉNÝL-CARBÎNOL (C₄H₄Ph)₂CH.OH. [151°]. S. (alcohol) 1.25; (ether) 3; (ligroïn) .5 at 15°. Got by reducing

the corresponding ketone with sodium-amalgam [Adam, A. Ch. [6] 15, 260; cf. Weiler, B. 7, 1189). Plates.

PHENYL-DI-PHENYL-CARBINYLAMINE C_sH_s,C_sH_s,CHPh.NH_s. [77°]. Got by reducing the oxim C_sH_s,C_sH_s,C(NOH),C_sH_s by sodium-amalgam (Koller, M. 12, 508). Needles (from ether), insol. water. Conc. H_sSO₄ forms a deepviolet colour, becoming claret-red on warming. -B'HOAc. [161°]. White needles (from water).-White B'HCl. $[252^{\circ}].$ needles.—B'HNO... [211°].—B'₂H₂PtCl₆4aq. [191°]. Yellow needles. Phenyl-triphenylcarbinyl-amine

NHPh.CPh, [146°]. Formed from aniline and CPh,Br (Elbs, B. 17, 703; Hemilian a. Silberstein, B. 17, 746). Hexagonal prisms, v. sol. CS₂. Yields a nitrosamine [c. 156°] and a tetra-sulphonic acid C25H17(SO,H)4N which yields easily soluble Ba2A1 and Cu2A1.

PHENYL TRI-PHENYL-CARBINYL KE-

TONE v. (β)-Benzpinacolin.

DI-PHENYL - DIPHENYLENE - DI - UREA. NHPh.CO.NH.C.H., C.H., NH.CO.NHPh. [above Concentric needles (from aniline). Formed by adding phenyl cyanate (2 mols.) to an ethereal solution of benzidine (Kühn, B. 18, 1478).

PHENYL-ENNYL PHENYL KETONE $[61^{\circ}].$ C₆H₅.CO.CHPh.C₈H₁₇. (350°-355°). Formed from deoxybenzoin (E. Bischoff, B. 22, 348). Yields an oxim [101°] crystallising in long needles.

PHENYLPHENYL - ETHYL ALCOHOL [1:3]C₄H₄Ph.CHMe.OH. [86°]. Formed by reducing C₆H₄Ph.CO.CH₉ with sodium-amalgam (Adam, A. Ch. [6] 15, 257). Crystalline mass. Cannot be distilled.

DI-PHENYL DIPHENYLETHYLENE DI-KETONE CHPhBz. CHPhBz. Bidesyl. Hydrooxylepiden. [255°]. Formed as described in vol. iii. p. 139. Needles (from benzene), insol alcohol. On boiling for two hours with alcoho it changes to an isomeride [261°], sol. alcoho (Fehrlin, B. 22, 553).

Isomeride. [161°]. Isobidesyl. Accompanies bidesyl when prepared from deoxybenzoïn, NaOEt, and I, or from BzCHBrPh, NaOEt, and deoxybenzoin (Knövenagel, B. 21, 1356). Prisms (from alcohol). Reacts with hydroxylamine forming C₅₆H₁₇N₃O₄ [110°-120°]. Conc. H₂SO₄ forms with bidesyl and isobidesyl a green solution, turning brown. NH₂ forms tetraphenyl-pyrrole.

PHENYL-ETHYL PHENYL RETONE C_aH_b . CO.CHMe. C_aH_b . [53°]. (318° cor.). Formed from C_aH_b . CO.CHNa. C_aH_b and MeI (V. Meyer a. Oelkers, B. 21, 1297). Needles (from alcohol). Forms an oxim [120°].

Phenyl phenyl-ethyl ketone

C.H. CO.CH. CH. CH. [73°]. (above 360°). Formed by reducing phenyl styryl ketone with zinc-dust and HOAc (Schneidewind, B. 21, 1325). Plates (from alcohol). Yields an oxim Isoamyl nitrite and N: OEt form [87°]. C15H13NO2 [126°].

Phenyl diphenyl-ethyl ketone

C₆H₅.CO.CHPh.CH₂Ph. [120°]. Formed from phenyl benzyl ketone, NaOEt, and benzyl chloride (V. Meyer a. Oelkers, B. 21, 1300). Needles (from alcohol). Yields an oxim [208°].

PHENYL-ETHYL PHENYL KETONE | **CARBOXYLIC ACID** C₆H₃.CO.CHPh.CH₂.CO₂H. [152°] (J. a. M.); [161°] (M. a. O.). Formed by oxidising dehydro-acetone-benzil with CrO, and HOAc (Japp a. Miller, B. 18, 184). Needles .--

BaA', 2aq. AgA'. Ethyl ether EtA'. Formed Ph.CO.CHNaPh and ClCH2CO2Et (V. Meyer a. Oelkers, B. 21, 1305). When heated it yields an anhydride O CO .CH. [152°], whence am-

monia forms NH < CO .ÇH₂ [189°]. Phenylhydrazine acting on desylacetic acid forms $C_{2z}H_{18}N_2O$ [110°] and $(C_{zz}H_{1z}N_2O)_2$ [243°]. The former, NHPh.N<CO .CH $_2$? may be reduced

by Na and isoamyl alcohol to NH CO-CH2 [207°]. Aniline conver NPh $<_{\mathrm{CPh:CPh}}^{\mathrm{CO}}$ [190°] and converts the acid into

300°.

Phenyl phenyl-ethyl ketone carboxylic acid. Methylether C₈H₃·CO.CH(ČO₂Me).CH₂Ph. (c. 253° at 50 mm.). Formed from methyl benzoyl-acetate, NaOEt, and benzyl chloride (Perkin a. Calman,

C. J. 49, 155)

Phenyl di-phenyl-ethyl-ketone carboxylic acid $C_{22}H_{18}O_3$ i.e. C_6H_5 .CO.CH $_2$.CPh $_2$.CO $_2$ H. [183°]. Formed by heating the lactone of oxytri-phenyl-crotonic acid with alcoholic potash (Japp a. Klingemann, C. J. 57, 681). Plates. Reduced by boiling HIAq to tri-phenyl-butyrolactone. Phenyl-hydrazine in alcoholic solution yields $C_{28}H_{22}N_2O$, which is reduced by Na and isoamyl alcohol to oxy-tri-phenyl-pyrrole dihydride $C_{22}H_{18}NO.-NH_4\Delta'.-Ag\Delta'$. On heating with alcohol and hydroxylamine hydrochloride it forms C22H1,NO2 [152°] and C2H1,NO [143°].

Methylamide C₂₃H₂₁NO₂. [156°]. Formed from tri-phenyl-crotolactone and methylamine.

Ethylamide. [130°]. Needles.
Phenyl tri-phenyl-ethyl ketone carboxylic acid. Methylamide

C₆H₅.CO.CHPh.CPh₂.CONHMe. [260°]. Formed by heating the lactone of oxy-tetra-phenyl-crotonic acid with alcoholic NH₂Me at 100, to 200° (Klingemann a. Laycock, B. 24, 514). Plates, sl. sol. alcohol. Yields, on distillation,

sl. sol. alcono..

NMe CPh:CPh
CO .CPh
PHENYL-HEPTYL
[59°]. (345) KETONE $C_6H_9.CO.CHPh.C_6H_{13}.$ [59°]. (345°). Needles or plates (from alcohol). Yields an oxim [89°] (Bischoff, B. 22, 347)

PHENYL-DI-PHENYL-HYDRAZINE

NHPh.NHC, H, Ph. [127°]. Formed by reducing C,H₅.N₂.C₈H₄Ph with ammonium sulphide (Locher, B. 21, 911). Needles or plates (from dilute alcohol). Yields a di-acetyl derivative [203°

PHENYL-DI-PHENYL KETONE v. PHENYL-

PHENYL-PHENYL MERCAPTAN C12H10S i.e. C.H.Ph.SH. [111°]. Formed by educing C.H.Ph.SO.cl with tin and HCl (Gabriel a. Deutsch, B. 13, 386).—Pb(S.C.H.Ph)2. Reddish-

brown pp.—C₁₂H₀SMe. [108°]. Needles (Obermayer, \hat{B} . 20, 2927).

DI-PHENYL-DI-PHENYL-METHANE

(C_eH₄Ph)₂CH₂. [161°]. (360°). Formed from diphenyl, methylal, HQAc, and H₂SO₄ (Weiler, B. 7, 1188). Formed also from diphenyl, CH2Cl2, and AlCl3; the yield being bad (Adam,

A. Ch. [6] 15, 254). DI-PHENYL DI-PHENYL-METHYLENE **DISULPHIDE** CPh₂(SPh)₂. [139°]. Formed by passing dry HCl into a hot mixture of benzophenone, phenyl mercaptan, and ZnCl₂ (Baumann, B. 18, 888). Prisms (from ether).

PHENYL-PHENYL DI-PHENYL-ETHYL C₆H₄Ph.CO.CHPh.CH₂Ph. [158°]. Formed from C,H,Ph.CO.CH,Ph, NaOEt, and benzyl chloride (Päpcke, B. 21, 1339). Needles

(from alcohol). Yields an oxim [1756].
PHENYL PHENYL - PROPINYL KETONE CARBOXYLIC ACID CPh: C.CHBz.CO₂H. [135°]. (c. 275° at 40 mm.). Formed from di-benzoylpropionic ether and alcoholic potash (Kapf a. Paal, B. 21, 1488). Yellow needles or plates, insol. water. Phenyl-hydrazine forms C₂₀H₂₁N₄O [100°]. Boiling with alcohol and HClAq forms di-phenyl-furfurane carboxylic acid [217°]. Alcoholic NH, yields di-phenyl-pyrrole carboxylic ether.—KA'2aq: yellow needles (from alcohol).

PHENYL a. PHENYL - PROPYL KETONE C₀H₅.CO.CHEtPh. [58°]. (324° cor.). Formed from phenyl benzyl ketone, NaOEt, and EtI (V. Meyer a. Oelkers, B. 21, 1299). Needles.

Yields an oxim [130°].

Phenyl phenyl-propyl ketone C₆H₅.CO.CH₂.CHPh.CH₃ or C₆H₅.CO.CMe₂Ph [70°]. (340°-345° i.V.). Formed by heating acetophenone with HI and P at 140° (Graebe, B. 7, 1625). Plates (from alcohol), v. e. sol. ether.

PHENYL DI PHENYL PROPYL THIO-UREA NHPh.CS.NH.CH2.CHPh.CH2Ph. [129°]. Formed from di-phenyl-propyl-amine and phenyl thiocarbimide (Freund a. Remse, B. 23, 2862).

PHENYL PHENYL - PYRAZYL KETONE C₃H₂PhN₂.CO.C₆H₅. [123°]. Formed by heating phenyl-pyrazole with BzCl at 245° (Balbiano, G. 19, 139). Needles (from dilute alcohol). Yields an oxim [154°] and a phenyl-hydrazide [140°].

DI-PHENYL-DI-PHENYL SULPHIDE C₂₄H₁₆S *i.e.* (C₆H₁Ph)₂S. [172°]. Formed by distilling Pb(S.C₆H₁Ph)₂ (Gabriel a. Deutsch, B. 13, 386). Plates, sol. alcohol and ether.

Di-phenyl di-phenyl disulphide (C,H,Ph)2S2 [150°]. Formed by atmospheric oxidation of C₆H₄Ph.SH (G. a. D.). Needles, sol. alcohol

and CS2.

PHENYL-PHENYL SULPHOCYANIDE

C_sH_sPh.SCN. [84°]. Formed from Pb(SC_sH_sPh). and ICy (Gabriel a. Deutsch, B. 13, 389). White crystals.

DI-PHENYL-DI-PHENYL SULPHONE

(C₆H₄Ph) SO₂. [216°]. Made by oxidising (C_eH₄Ph)₂S with KMnO₄ (Gabriel a. Deutsch, B. 13, 387). Plates, sl. sol. ether.

PHENYL-PHENYL-SULPHONO-BENZAM-IDINE PhC(NPhH):NSO₂Ph. [139°]. Formed by the action of aniline on the imido-chloride obtained from the anilide of benzene sulphonic acid by the action of PCl_s (Wallach a. Gossmann, A. 214, 214). Narrow plates (from alcohol). Appears not to combine with HCl. On dry distillation it gives di-phenylamine, benzonitrile,

SO₂, and phenyl sulphides. PHENYL-PHENYL-THIOCARBIMIDE

C.H.Ph.N:CS. [58°]. Formed by distilling (C,H,Ph.NH)2CS with P2O, (Zimmermann, B. 13, 1964). Needles.

PHENYL PHENYL-THIO-GLYCOLLIC ACID [170°]. C,H,Ph.S.CH₂.CO₂H. Formed from chloro-acetic acid and C.H.Ph.SNa (Gabriel a. Deutsch, B. 13, 389). Colourless crystals, sl. sol. water.

Phenyl-phenyl-di-thio-di-glycollic acid $C_{12}H_{a}(S.CH_{2}.CO.H)_{2}.$ [252°]. Form $C_{12}H_{a}S_{2}Pb$ and $ClCH_{2}.CO.H.$ Crystals. Formed from

DI-PHENYL-DI-PHENYL-THIO-UREA CS(NH.C_eH₄Ph)₂ [228°]. Formed from p-amidodiphenyl and CS₂ (Zimmermann, B. 13, 1963). Plates.

PHENYL PHENYL DI-TOLYL-CARBINYL

KETONE v. PHENYL-TOLYL-PINACOLIN.

PHENYL-PHENYL-p-TOLYL-CARBINYL-UREA NHPh.CO.NH.CHPh.C.H.Me. Formed from C₆H₄Me.CHPh.NH₂ and phenyl cyanate (Goldschmidt a. Stöker, B. 24, 2802).

PHENYL PHENYL-DI-XYLYL-CARBINYL KETONE v. PHENYL-XYLYL-PINACOLIN.

PHENYL PHOSPHATES.

Mono - phenyl di - hydrogen phosphate C₆H₆O.PO(OH)₂. Phenyl-phosphoric acid. [98°]. A product of the action of P₂O₅ on phenol (Rembold, Z. 1866, 652; Jacobsen, B. 8, 1519; Rapp, A. 224, 157). Deliquescent needles. Partially decomposed by boiling water into phenol and phosphoric acid. Yields phenol and HPO, on distillation.—CaA".—BaA": prisms.

Chloride C.H.O.POCl₂. (242°). A product of the action of POCl₃ on phenol. Formed also from C₈H₅O.PCl₄ and SO₂ (Anschütz a.

Emery, A. 253, 110).

Di-phenyl hydrogen phosphate

(C₆H₅O)₂PO.OH. [56°] (R.). A product of the action of P₂O₅ on phenol. Formed also by boiling (PhO) PO (1 mol.) with aqueous KOH (1 mol.) (Glutz, A. 143, 193), and by decomposing the chloride with water. - BaA'2. - AgA'. - NH3PhA'. [c. 160°] (Wallach, B. 8, 1235).

Chloride (C₆H₅O)₂POCl. (315° at 272 mm.). Formed from phenol and POCl,. Oil, slowly de-

composed by water.

Anilide (C,H,O),PO.NHPh. [129°]. Six-

sided tables, v. sl. sol. water (W.).

Tri-phenyl phosphate (C_eH₂O)₈PO. [45°] (J.). (245° at 11 mm.) (A. a. E.) Formed, together with chloro-benzene, by the action of PCl_s on phenol (Williamson a. Scrugham, C. J. 7, 240). Formed also from phenol and POCl_s (Jacobsen), or PCl_s (Glutz), followed by water. It is also a product of the action of water on CoH, O.PCl, (Anschütz a. Emery, A. 253, 110).

Preparation.—A mixture of phenol (280 pts.) and phosphorus oxychloride (150 pts.) is heated to boiling with an inverted condenser for 16 hours, the excess of phenol is removed by distillation or by shaking with dilute NaOH; the yield is 90 p.c. of the theoretical (Heim, B. 16,

1763).

Properties.-Needles, insol. water, sl. sol. alco-

hol, v. sol. ether.

Reactions. - By distillation with sodium acetate or benzoate it gives phenylacetate or benzoate. Heated with potassium sulphide it yields tri-phenyl-thio-phosphate PS(OC,H,), together with phenol and some diphenyl and di-phenyl oxide. On distilling with MgO, PbO, ZnO, or CaO the chief product is phenyl, together with some diphenylene-oxide (Kreysler, B. 18, 1716). Chlorine passed into its cold ethereal solution forms (C₈H₅O)₈PCl₂ (A. a. E.). Reference.—NITRO-PHENYL-PHOSPHATES.

PHENYL-PHOSPHINE $C_6H_5PH_2$. Mol. w. 110. (161°). S.G. 15 1·001. A product of the action of alcohol on C, H, PI, H and on C, H, PCl, (Michaelis, B. 7, 6; 10, 807; 12, 338; A. 181, 803). Pungent liquid.

Reactions. - 1. Absorbs oxygen, forming C₆H,PHO.OH [70°]. - 2. Sulphur at 100° forms liquid C₆H₅PH₂S and solid (C₆H₅P)₃S [138°].—3.

CS₂ at 150° forms C₁₄H₁₂P₂S₃.

Salts.—C₆H₅PH₄I. Needles.—B'₂H₂PtCl₆. Phenyl-di-chloro-phosphine C.H.PCl.. Phosphenyl chloride. Mol. w. 179. (225° cor.). S.G. $\frac{0}{4}$ 1.3428 (Thorpe, C. J. 37, 347). Formed by passing benzene and PCl, through a red-hot tube. Formed also by heating HgPh2 with PCl3 at 180°, and by the action of AlCl, on a mixture of benzene and PCl₃ (Michaelis, B. 12, 1009; A. 181, 280). Pungent fuming liquid, decomposed by water into HCl and C₆H₅PHO.OH [70°]. Chlorine forms C₆H₅PCl₄ [73°], which is split up at 180° into C₆H₅Cl and PCl₃, and is converted by water into C₆H₅PO(OH)₂. SbCl₅ forms CH PCl SbCl₇ and CH PCl SbCl₇ [100] C.H.PCl.SbCl, an unstable yellow crystalline Br yields C6H5PCl2Br2 [2080] and powder. C₆H₅PCl₂Br₄. HI forms C₆H₅PHI₃. H₂S yields oily (C₆H₅PS)₂ and crystalline C₂₄H₂₆N₂S, [193°). S forms liquid C₆H₅PSCl₂ (270°) (Köhler, B. 13, 464).

Phenyl-di-bromo-phosphine C₆H₅PBr₂. [257°]. Got from HgPh2 and PBr3 (Michaelis, B. 9, 519). Br yields CaH.PBr. [207°] and

C₆H₅PBr₆.

Di-phenyl-phosphine (C₆H₅)₂PH. (c. 280°). S.G. 12 1.07. Formed, together with the acid (CaHa) PO.OH by the action of dilute NaOHAq on $(C_6H_5)_2$ PCl (Michaelis a. Gleichmann, B. 15, 801; Dörken, B. 21, 1508). Oil, v. sol. alcohol. Weak base. Yields (C,H,)₂PO.OH on oxidation. CS₂ forms (C₆H₅)₁P₂H₄CS₂ [157°]. — B'HCl. Crystalline.—B'₂H₂PtCl₅.—B'HI.

Di-phenyl-chloro-phosphine $(C_6H_b)_2PCl.$ S.G. 15 1.229. Formed from HgPh, and C₆H₂PCl₂ at 225° (Michaelis, B. 10, 627; 18, 2109; A. 207, 208). Formed also by heating C_aH_aPCl₂ in a sealed tube for 120 hours at 300° (Dörken, B. 21, 1505). Thick liquid, decomposed by water or Na₂CO₃Aq into (C₆H₅)₂PH and (C₆H₅)₂PO.OH. Absorbs oxygen from the air forming (C,H,)2POCl.

Tri-phenyl-phosphine P(C₆H₅)₃. [79°]. (above 360°) (Michaelis, B. 15, 802, 1610; A. 229, 297). Formed from C₆H₅PCl₂, bromo-229, 297). benzene and Na. Prepared by adding sodium to a mixture of PCl_s (1 mol.) and C_oH_sBr (3 mols.) diluted with 4 volumes of dry ether and kept cool. The reaction is completed by boiling for 12 hours, and the ether is filtered off and evaporated to crystallisation.

Properties.—Large prisms or tables, sol. alcohol, ether, and benzene, insol. water. weak base. Does not combine with CS2

Salts.—B'HI. [215°]. Prisms, sol. alcohol, insol. water.—B'₂H₂PtCl_e.—B'HgCl₂.

Alkylo-halides .-B'Mel. [183°]. B'MeClaq. [213°]. Crystals, v. e. sol. water and alcohol, insol. ether.—B'2Me2PtCl2. [238°]. B'EtI. [165°]. — B'PrI. [201·5°]. — B'PrI 2aq. [191°]. — B'CH₂PrI. [177°]. — B'C₅H₁₁I. [174°]. — B'CH₂PhCl aq. [288°].—B'CH₂PhBr. [275°].— B'CH,PhI. $[253^{\circ}]$. — $B'_{2}CH_{2}I_{2}$. $[231^{\circ}]$. [above 300°] B'2C2H4Br2.

Benzylo-nitrate B'C,H,NO. [203°]. S.

·838 at 15°. Needles (from water).

Bensylo-picrate B'C,H,C,H2N,O, [148°]. Benzylo-bichromate $(B'C, H_7)_2Cr_2O_T$ Benzylo-sulphocyanide B'C,H,SCN. [189°]. Prisms, v. sol. alcohol, sl. sol. water.

(C,H,),PO. Tri-phenyl-phosphine oxide [153.5°]. (c. 360°). Formed by oxidising triphenyl-phosphine with Br and NaOHAq or with conc. HClAq and KClO₃. Prisms (containing aq), sl. sol. water, v. e. sol. alcohol. Yields two trinitro- compounds [68°] and [242°], the latter of which yields a tri-amido-compound [258°], whence bromine water forms OP(C₆H₂Br₂(NH₂))₃ [206°]. -B'HNO₃. [75°]. Yellow needles, decomposed by water.

Tri-phenyl-phosphine sulphide (C6H2), PS. [157.5°]. Formed from Ph,P and S in CS2. Colourless needles, v. sol. alcohol, insol. water.

Tri-phenyl-phosphine selenide (C₈H₅)₃PSe. [184°]. Formed from Ph,P and Se. Necdles.

Tetra-phenyl-diphosphine P₂(C₆H₅)₄, [67°]. (400°). Formed from (C₆H₅)₂PH and (C₆H₆)₂PH (Dorken, B. 21, 1509). With CS₂ at 138° it forms Ph₂P.CS.SPH₂Ph₂ [157°], insol. ether.

PHENYL-PHOSPHINIC ACID v. PHENYL.

PHOSPHITES.

Di-phenyl-phosphinic acid (C,H,)2PO.OH. [190°] (Michaelis, B. 8, 922; 11, 885; 15, 802; 18.2113). Formed by oxidation of (C, H₅)₂PCl or of di-phenyl-phosphine. Needles, sl. sol. cold alcohol.-- CaA', Baq.-- AgA': bulky pp.

Ethyl ether EtA. [165°]. Needles. Phenyl ether (C.H.).PO.OC.H. [136°]. Formed by oxidising (C.H.).POC.H. (265°-270°), which is got by warming (C, H,)2PCl with phenol. Small needles, sol. hot alcohol, insol. water.

PHENYL PHOSPHITES.

Mono-phenyl dihydrogen phosphite $C_4H_5O.P(OH)_2$. Formed by adding water to $C_6H_5O.PCl_2$ (216°) which is got by the action of PCl_s on phenol (Noack, A. 218, 90; Anschütz, A. 239, 310). Uncrystallisable syrup, decomposed by addition of more water into phenol and HaPOs

Isomeride C₈H,PO(OH)₂. Phenyl phosphinic acid. Phosphenylic acid. Benzene phosphonic acid. [158°]. S. 23.5 at 15°. Formed by the action of water on C₆H₅PCl₄ (Michaelis, A. 181, 321). Plates, sol. alcohol and ether. Yields, on nitration, C₈H₄(NO₂)PO(OH)₂ [132°], which may be reduced to crystalline C₈H₄(NH₂).PO(OH)₂.—

NaHA"xaq. — Na₂A" 12aq. — CaH₂A"₂. — CaA" 2aq. — SrH₂A"₂ aq. — CuA". — Fe₂A"₂ 2½aq. — Ethers. — Me₂A". (247°). — Et₂A". (267°). — EtHA". — EtAgA". — PhHA". [57°]. Formed from C₈H, POCl₂ by successive treatment with phenol and boiling water. Needles, sl. sol. water.-Ph.A". [63.5°]. Formed from phenol and CoHsPCl4.

Chloride C_sH₅POCl₂. (258°). Formed from C.H.PCl, and water, and also from C.H.PCl2 and O. Thick liquid.

Di-phenyl hydrogen phosphite (C,H,O),POH. Formed by adding a little water to the compound $(C_sH_sO)_2$ PCl (295° at 731 mm.) which is got from phenol and PCl₃ (N.). Uncrystallisable. Decomposed by more water into H.PO. and phenol.

Tri-phenyl phosphite (C₆H₅O)₅P. (220° at 11 mm.). S.G. _{Th} 1184. Formed by heating phenol (3 mols.) with PCl₅ (1 mol.) at 230° in a current of dry CO₂ (Noack, A. 218, 91; Anschütz alcohol. Decomposed by water into phenol and H_3PO_3 . Dry bromine forms $(C_6H_5O)_3PBr_2$, which is converted by water into $(C_6H_5O)_3PO$.

PHENYL-PHTHALIDE v. Anhydride of Oxx-

BENZYL-BENZOIC ACID.

Di-phenyl-phthalide i.e. $C_8H_4 < \frac{CPh_2}{CO} > 0$. Phthalophenone. Tri-phenyl-carbinol carboxylic anhydride. [115°]. Formed by the action of AlCl, on a mixture of benzene and phthalyl chloride (Friedel a. Crafts, A. Ch. [6] 1, 523,

Baeyer, B. 12, 642; A. 202, 50) or of benzene with [1:2]C₆H₁Bz.CO.OAc (Von Pechmann, B. 14, 1866). Got also by heating phthalyl chloride with HgPh₂ (Noelting, B. 17, 387). Plates (from alcohol). Insol. KOHAq, sol. boiling alcoholic potash, but reppd. by acids.

Reference.—DI-AMIDO-DI-PHENYL-PHTHALIDE. DI-PHENYL-PHTHALIDE CARBOXYLIC ACID v. Anhydride of Tri-Phenyl-Carbinol Di-CARBOXYLIC ACID.

PHENYL-PHTHALIMIDE v. Phenyl-imide of Phthalic acid.

PHENYL-PHTHALIMIDINE C, H, NO i.e. $C_6H_4 < \stackrel{CH_2}{<} NPh.$ [160°]. Got by heating plithalide with aniline at 210° (Hessert, B. 10, 1450; 11, 239; Racine, A. 239, 88). Formed also by reducing the phenylimide of phthalia acid with tin and HClAq (Graebe, A. 247, 305). Plates (from alcohol), v. sl. sol. hot water.

PHÈNYL-PIPERAZINE v. PHENYL-PYRAZINE HEXAHYDRIDE.

PHENYL-PIPERIDINE v. PHENYL-PYRIDINE HEXALIYDRIDE.

PHENYL-PROPANE v. CUMENE.

Di-phenyl-propane $\mathrm{CH_2}(\mathrm{CH_2Ph})_2$. Di-benzyl-methane. (290°-300°). Formed by heating dibenzyl ketone with HIAq and P at 180° (Graebe, B. 7, 1623). Got also from CH₂Cl.CHCl.CH₂Cl,

benzene, and AlCl, (Claus, B. 18, 2935). Liquid. Di-phenyl-propane CH, Ph. CHPh. CH,. (291°-295°). S.G. 9 996. Formed from benzene, AlCl₃, and propylene chloride or allyl chloride (Silva, C. R. 89, 606). Formed also by adding H₂SO, to a cooled mixture of toluene and styrene (Kraemer, Spilker, a. Ebenhardt, B. 23, 3274). Oil, sol. alcohol and ether.

Di-phenyl-propane CMe,Ph2. $(282^{\circ}).$ Formed by the action of benzene and AlCl. on CCl₂Me₂ or CH₂:CCl.CH₃ (Silva, Bl. [2] 34, 674; 85, 289).

 $C_8H_5(C_6H_5)_2$ Tri-phenyl-propane Formed by the action of AlCl, upon a mixture of benzene and glyceryl tri-bromide or trichloride (Claus a. Mercklin, B. 18, 2935). Oil.

TRICARBOXYLIC PHENYL - PROPANE Formed from benzyl-malonic ether, NaOEt, and chloro-acetic ether (Fittig, A. 256, 92).

Small prisms, m. sol. warm water. benzyl-succinic acid [161°] when boiled with water.—Ca₃A'''₂ 6½aq.—Ba₃A'''₂ 2½aq.—Ag₃A'''.

Ethyl ether Et₂A'''. (336° cor.). S.G.

 $\mu_D = 1.484$ at 20°. Formed from sodium ethane tricarboxylic ether and benzyl chloride (Bischoff a. Mintz, B. 23, 653)

Di-phenyl-propane tetra-carboxylic acid. Di-nitrile of the di-ethyl ether (CO₂Et)₂C(CH₂.C₆H₄.CN)₂. [86°]. Formed from sodium malonio ether and [1:2]C₆H₄(CN).CH₂Cl (Hausmann, B. 22, 2019). Prisms.

Tri-phenyl-propane dicarboxylic acid. Imide $C_6H_4 < \frac{C(C,H_7)_2}{CO.NH^2} > CO.$ [174°]. Formed

from C₆H₄<CH₂.ÇO .NH, benzyl chloride, and NaOEt (Pulvermacher, B. 20, 2496). Yellow plates (from alcohol).

Benzyl-imide CanH25NO2. [109°]. PHENYL PROPARGYL OXIDE CoH, O i.c. C₈H₅.O.C₈H₈. (210°). S.G. ⁶ 1.246. Formed

by the action of alcoholic potash on the bromoallyl derivative of phenol (Henry, C. R. 96, 1233). Liquid.

PHENYL-PROPARGYL-UREA C10H10N2O i.e. CH:C.CH₂.NH.CO.NHPh. [133°]. Formed from propargylamine oxalate, phenyl cyanate, and K₂CO₃Aq (Paal a. Hempel, B. 24, 3042). Needles.

PHENYL-PROPENYL-AMIDINE C, H12N2 [68°]. Formed from propionitrile, HCl, and aniline (Michael a. Wing, Am. 7, 72). Hexagonal plates, sol. alcohol and ether. -B'2H2PtCl

PHENYL PROPENYL METHYL KETONĚ CARBOXYLIC ACID C₁₂H₁₂O₅ i.e. CH₂.CO.C(CH₂.CO₂H):CHPh. [125°]. S. 3 at 100°; '5 at 15°. Formed by boiling benzoic aldehyde with acetyl-propionic acid and NaOAc (Erdmann, A. 254, 187).—BaA', 5aq.—CdA', 2aq.

-MgA'₂.—CaA'₂3 $\frac{1}{2}$ aq: groups of needles. Methyl ether MeA'. (c. 215° at 38 mm.).

PHENYL PROPENYL SULPHIDE C₅H₅.S.CMe:CH₂. (208°). Formed by heating phenyl-sulphydro-crotonic acid (Autenrieth, A.

254, 232). Yellowish oil, volatile with steam. TRI-PHENYL PROPENYL TRISULPHONE C₉H₃(SO₂Ph)₃. [226°]. Got from CHBr(CH₂Br)₂ and sodium benzene sulphinate (Stuffer, B. 23, 1413). Needles, m. sol. hot alcohol.

DI-PHENYL-PROPENYL-DI-UREA

CEt N.CO.NHPh. [170°]. Formed from propionamidine and phenyl cyanate (Pinner, B. 23, 2924). Needles, m. sol. alcohol. Yields diphenyl-urea [234°] when boiled for a long time with alcohol. Boiling dilute HOAc forms NHPh.CO.NH.CO.C.H. [137°]. PHENYL-PROPIOLIC ACID C,HO, i.e.

C₆H₅.C:C.CO₂H. Mol. w. 146. [137°].

Formation.—1. By the action of Na and CO₂ on ω -bromo-styrene (Glaser, Z. [2] 4, 328; A. 154, 140).-2. By boiling αβ-di-bromo-phenyl-propionic acid or its ether with alcoholic potash G.; Perkin, jun., C. J. 45, 173; Weger, A. 221, 70; Roser, A. 247, 138).—3. By boiling a-bromocinnamic acid with alcoholic potash (G.; Bairisch, J. pr. [2] 20, 181).

Properties .- Trimetric twin crystals. Reduced by zinc and HOAc to cinnamic acid (Aronstein a. Hollemann, B. 22, 1181) and by sodium-amalgam to phenyl-propionic acid. Dry HCl passed through its solution in HOAc forms two chloro-cinnamic acids [142°] and [132°] (Michael, J. pr. [2] 40, 63).

Salts.-KA'.-BaA'2 aq: square tables.-BaA'₂ 2aq: needles.—BaA'₂ 3aq.—CuA'₂ 4aq.— AgA': sparingly soluble pp.

Ethyl ether EtA'. (260°-270°).

References .- AMIDO- and NITRO- PHENYL-PROPIOLIC ACID.

a-PHENYL-PROPIONIC ACID CoH10O2 i.s. CH3.CHPh.CO.H. Hydratropic acid. (268°). Formed by reducing atropic acid (15 g.) with (5 p.c.) sodium-amalgam (90 g.) (Trinius, A. 227, 262; cf. Kraut, A. 148, 244; Fittig, A. 195, 165). Got also from acetophonone by treatment with HCy, the resulting CH3.CPh(OH).CN being saponified and the oxy-phenyl-propionic acid reduced by HI and P (Janssen, A. 250, 135). Oil, with characteristic odour. Yields atrolaction acid CH_s.CPh(OH).CO,H on oxidation by alkaline KMnO4.—BaA'22aq.—CaA'23aq.— CaA'22aq. AgA': plates (from hot water).

Methyl ether MeA'. (221°).

Ethyl ether EtA'. (230°). S.G.

 $\begin{array}{ccc} 1\text{-}0147 & R_{\infty} = 83\text{-}3 \text{ (Bruhl)}. \\ & \textit{Amide CHMePh.CONH}_{2}. & [92^{\circ}]. \\ & \textit{Nitrile CHMePh.CN}. & (232^{\circ}). \end{array}$ $[92^{\circ}].$ Formed by the action of PCl, on the amide, and from sodium phenyl-acetonitrile and MeI (Oliveri, G. 18, 572).

β-Phenyl-propionic acid CH, Ph.CH2.CO, H. Hydrocinnamic acid. Mol. w. 150. [48.7°]. (280° i.V.). H.C.v. 1,084,600. H.C.p. 1,085,500. H.F. 105,500 (Stohmann, J. pr. [2] 40, 135). S. ·6 at 20°. A product of the putrefaction of albumen, 20 g. being got from 5 kilos of ox-brain

(Stöckly, J. pr. [2] 24, 17).

Formation.—1. By reduction of cinnamic acid by sodium-amalgam or HI (Erlenmeyer, A. 121, 375; 137, 327; Popoff, Bl. [2] 4, 375; Giacosa, H. 8, 108).—2. By reducing aß-dibromo-phenyl-propionic acid (Schmitt, A. 127. 319).—3. From ethyl-benzene vid CH, Ph.CH, Cl and CH₂Ph.CH₂·CN (Fittig, Z. 1869, 166; A. 156, 249.—4. By oxidising phenyl-propyl alcohol (W. von Miller, B. 9, 274).—5. By heating benzyl-malonic acid (Conrad, A. 204, 176).

Properties.—Needles (from water), sol. alcohol and ether; volatile with steam. taken internally it is secreted as hippuric acid

(E. a. H. Salkowski, B. 12, 653).

Salts.—NH₄A'. Plates, v. e. sol. water.—
CaA'₂ 1½aq.—CaA'₂2aq.—BaA'₂2aq.—PbA'₂aq.
ZnA'₂.—AgA'. Phenyl-hydrazine salt:
[572] siller peddag (Liebergen) 2, 24 1137. [57°]; silky needles (Liebermann, B. 24, 1107).

Methyl ether MeA'. (239° cor.). S.G. 2 1.0455. S.V. 196.0. Liquid, with peculiar odour. Ethyl ether EtA'. (248° cor.). S.G. 2

1.0343. S.V. 221.5 (Lossen, A. 254, 63). n-Propyl ether Pra'. (262° cor.). S.G. 2 1.0152 (Weger, A. 221, 79). S.V. 246.

Isoamyl ether C₄H₁₁A'. (292°). S.G. 2

·981; 49 ·952 (E.).

Benzyl ether C,H,A'. (c. 300°). $\frac{21}{17.8}$ 1.074. Formed by warming benzyl acetate with sodium (Conrad a. Hodgkinson, B. 10, 254;

A. 193. 300). Amide C.H., NO. [105°]. Formed by heating the ammonium salt at 230° (Hofmann, B. 18, 2740). Small needles.

Nitrile C.H.N. (261° cor.). S.G. 1 1.0014.

Occurs in the oil of Nasturtium officinale (Hofmann, B. 7, 520).

Di-a-phenyl-propionic acid CH3.CPh2.CO2H. [173°]. (above 300°). Made by oxidation of the (β)-pinacolin of acetophenone CPh2Me.CO.CH2 [41°] with chromic acid mixture (Thörner a. Zincke, B. 11, 1993). Formed also by dissolving pyruvic acid and benzene in H2SO4 at 0° (Böttinger, B. 14, 1595). White leaflets, sol. ether, alcohol, and hot water.-KA'-CaA', 11aq.-BaA' 2aq.—AgA': white pp.

aß-Di-phenyl-propionic acid

[84°]. Formed by the CH,Ph.CHPh.CO,H. action of sodium-amalgam on a mixture of benzyl chloride and ClCO₂Et (Wurtz, C. R. 70, 850). Got also by reducing phenyl-cinnamic acid (Oglialoro, J. 1878, 821). Needles, sl. sol. boiling water.—CaA'2.—PbA'2. [146°].—AgA'.

Nitrile CH2Ph.CHPh.CN. [58°]. (c. 335°)

(Janssen, A. 250, 133).

Di-β-phenyl-propionic acid

CHPh₂.CH₂.CO₂H. [145°]. Formed from allocinnamic acid, benzene, and H2SO4 (Liebermann a. Hartmann, B. 25, 960). Needles.—CaA'2.

aaß-Tri-phenyl-propionic acid

CH2Ph.CPh2.CO2H. [162°]. Formed by saponitying the nitrile with HOAc and fuming HClAq at 220° (Neure, A. 250, 147). White needles, v.

sl. sol. water.—AgA': white insoluble powder. Nitrile CH₂Ph.CPh₂CN. [126°]. Form from CHPh₂CN, benzyl chloride, and NaOEt.

Tri-\$-phenyl propionic acid Ph.C.CH2.CO2H. Formed from Ph.C.CH(CO2Et)2 by boiling with alcoholic potash (Henderson, C. J. 51, 226; B. 20, 1014). Prisms (from alcohol), v. sl. sol. water.—KA'aq.—NaA'aq.—BaA'₂aq. -AgA'. Minute needles.

Ethul ether EtA'. [81°].

References .- AMIDO-, AMIDO-SULPHO-, BROMO-, BROMO-SULPHO-, BROMO-AMIDO-, BROMO-NITRO-, CHLOBO-, IODO-, IODO-OXY-, NITRO-AMIDO-, NITRO-OXY-, NITRO-, OXY-AMIDO- and OXY- PHENYL-PROPIONIC ACID.

PHENYL-PROPIONIC ALDEHYDE

C₆H₅.CH₂.CH₂.CHO. [208°]. Formed from n-cumene by successive treatment with CrO₂Cl₂ and water (Etard, A. Ch. [5] 22, 254). Oil. Forms a crystalline compound with NaHSO,

Oxim. Oil. Yields a solid hydrochloride. Ac₂O rapidly forms phenyl-propionitrile; hence

it is a syn- oxim (Dollfus, B. 25, 1918)

PHENYL-PROPYL ALCOHOL C.H.2O i.e. PhEt.OH. Mol. w. 136. (216°). S.G. ² CHPhEt.OH. 1.016. Formed by reducing phenyl ethyl ketone with sodium-amalgam (Errera, G. 16, 322; cf. Barry, B. 6, 1007. Got also from benzoic aldehyde by successive treatment with ZnEt₂ and water (Wagner, Bl. [2] 42, 330). Yields water (Wagner, Bl. [2] 42, 330). Yie CHPhEt.OAc (228° i.V.) and oily CHPhEtCl.

Phenyl-propyl alcohol CH₂Ph.CH₂.CH₂OH. (235° uncor.) (213° at 360 mm.). S.G. ²⁰ 1.0079. μ_s 1.5478. $\hat{R}_{\infty} = 68.79$ (Brühl, A. 200, 191). Occurs in storax as phenyl-propyl cinnamate (Miller, A. 188, 202). Prepared by reducing cinnamyl alcohol with sodium-amalgam (Rügheimer, B. 6, 214; A. 172, 123; Hatton a. Hodgkinson, C. J. 39, 819; Errera, G. 16, 810). Viscid oil. Heating with conc. HClAq yields CH_Ph.CH_CH_Cl (219°), whence NaOEt forms CH_Ph.CH_CH_CH_CE (220°).

Acetyl derivative C.H.Ph.OAc. (245°).

Phenyl-isopropyl alcohol

CH₂Ph.CH(OH).CH₃. Methyl-bensyl-carbinol. (215°). Got by reducing benzyl methyl ketone (E.). Liquid, with pleasant odour.

Di-phenyl-propyl alcohol

(301°). CH₂Ph.CHPh.CH₂OH. Formed by warming di-phenyl-propylamine nitrite (Freund a. Remse, B. 23, 2863). Oil, sol. alcohol.

PHENYL-PROPYL-AMINE $C_9H_{13}N$ CH₂Ph.CH₂.CH₂NH₂. (222° i.V.). Formed by reducing the phenyl-hydrazide (20 g.) of cinnamic aldehyde dissolved in alcohol (300 g.) by (2) p.c.) sodium-amalgam (700 g.) and HOAc (70 g.) at about 30°, the yield being 30 p.c. of the theoretical (Tafel, B. 19, 1930; 22, 1857). Liquid, m. sol. water, miscible with alcohol and ether. Absorbs CO2 from air, forming a crystalline mass. The aqueous solution is strongly alkaline.—B'HCl. [218°]. Plates.—B'H2SO4.-[156°]. Needles. — B'H₂C₂O₄.— B'2H2C2O4. B',H,PtCl,. Plates, sol. hot water.

Phenyl-isopropyl-amine CH2Ph.CHMe.NH2. (203°). Formed by the action of Br and KOHAq on the amide of phenyl-isobut ric acid (Edeleano, B. 20, 618). Liquid.—B'2H2PtCls. Needles.

Isomeride v. Cumidine.

Di-phenyl-propyl-amine CH_Ph.CHPh.CH₂NH₂. (316°). Formed by reducing the nitrile (30 g.) of phenyl-cinnamic acid dissolved in alcohol by treatment with Na (75 g.) (Freund a. Remse, B. 23, 2860). Liquid, sol. alcohol. Forms with HgCl2 a crystalline compound [107°].—B'HCl. [190°]. Needles.— Aurochloride [145°]. Di-acetyl derivative. [85°]. Needles.

PHENYL - n - PROPYL-TRIAZOLE CARB. $\stackrel{N.NPh}{CPr.N} > C.CO_2H.$ OXYLIC ACID [161°].

Formed by saponifying the nitrile which is made from phenyl-hydrazine dicyanide and butyric anhydride (Bladin, B. 25, 179). Cubes (from alcohol) or prisms (from benzene). - CuA' 2 2 aq.-AgA'. Bulky white pp.

Amide C,N,PhPr.CO.NH, [122.5°]. The thio-amide C₂N,PhPr.CS.NH, [130.5°] is got by the action of alcoholic ammonium sulphide

on the nitrile.

Phenyl-isopropyl-triazole carboxylic acid N.NPh CPr.N ≥C.CO₂H. [135°]. Formed in like manner (B.). Prisms. - HA'HCl. Needles. -CuA', 2 aq. — AgA : minute tables.

Methyl ether MeA'. [76°]. Amide. [128°]. Needles. The thio-amide C₂N₂PhPr.CS.NH₂ [148°], formed by the action of alcoholic NH₂ and H₂S on the crude nitrile,

crystallises in yellow needles.

DI-PHENYL-DI-ISOPROPYL-TETRAZONE NPhPr.N:N.NPhPr. [79°]. Formed by oxidising phenyl-isopropyl-hydrazine with HgO (Michaelis a. Philips, A. 252, 281). Octahedra.

DI-PHENYL-PROPYL TRICYANIDE

 $CPh \stackrel{N.CPr}{\leqslant_{N:CPh}} N$. [78.5°]. (239° at 15 mm.). Formed from benzonitrile, butyryl chloride, and AlCl, at 75° (Krafft a. Von Hansen, B. 22, 807). Decomposed by dilute H,SO, into NH, butyric acid, and HOBz.—B'.H.PtCl. Needles.

PHENYL-PROPYLENE v. ALLYL-BENZENE.

PHENYL PROPYLENE - ψ - THIO - UREA

C₁₀H₁₂N₂S i.e. $\stackrel{\text{CHMe.S}}{\text{CH}_2.N} > \text{C.NHPh.}$ [117°]. Formed from the isomeric phenyl-allyl-thio-rea [95°] and HClAq at 100° (Prager, B. 22, 2992). Oxidised by potassium chlorate and HClAq to CHMe.SO₃>C.NHPh [192°]. -- B'₂H₂PtCl₆. -- B'C₆H₃N₃O₇. [154°]. Yellow needles.

Ďi - Phényl - Isopropyl - Glyoxaline

TETRA - HYDRIDE CHPr NPh.CH₂ [95°]. Formed from di-phenyl-ethylene-diamine and isobutyric aldehyde (Moos, B. 20, 734). Crystalline groups (from alcohol), sl. sol. water.

PHENYL-PROPYL-HYDRAZINE $C_0H_{14}N_2$ i.e. $C_0H_{5}NH$.NHPr. [158°]. Formed by heating propionamide with phenyl-hydrazine (Freund, B. 21, 2461). Plates (from chloroform).

Phenyl-isopropyl-hydrazine C₆H₂NPr.NH₂ (233°). Formed from sodium phenyl-hydrazine and isopropyl bromide (Michaelis a. Philips, B. 20, 2485; A. 252, 278). — B'HCl. [135°]. Crystalline. Yields a tetrazone [85°]. Phenyl-thiocarbimide forms NPhPr.NH.CS.NHPh. [116°].

Acetyl derivative. [102°]. Needles. DI-PHENYL-PROPYLIDENE DISULPHIDE Me₂C(SPh)₂. [56°]. Formed from phenyl-mercaptan and acetone (E. Baumann, B. 19, 2804). Crystals, insol. water, v. sol. alcohol.

DI-PHENYL-PROPYLIDENE DISULPHONE Me₂C(SO₂Ph)₂. [97°] (B.); [182°] (F.). Formed by oxidising Me₂C(SPh)₂ (Baumann, B. 19, 2810), and also by treating CH₂(SPh)₂ with NaOH and MeI (Fromm, A. 253, 162). Crystals, almost insol. cold water.

PHENYL PROPYL KETONE C₁₀H₁₂O *i.e.* Ph.CO.Pr. Mol. w. 148. (221°). S.G. ¹⁵ ·992.

Formation.—1. By distilling calcium butyrate with calcium benzoate (Schmidt a. Fieberg, B. 6, 498).—2. By the action of AlCl₂ on a mixture of benzene and butyryl chloride (Burcker, Bl. [2] 37, 4).—3. By boiling benzoyl-acetic acid with dilute alcoholic potash (Baeyer a. Perkin, B. 16, 2131; C. J. 45, 181).

Properties. — Oil. Does not unite with NaHSO, Oxidised by CrO, and H₂SO, to benzoic and propionic acids (Popoff, B. 6, 560). CrO₂Cl₂ forms C₁₀H₁₂O2CrO₂Cl₂, a brown powder.

Phenyl isopropyl ketone Ph.CO.Pr. (209°–217°). Formed by distilling calcium isobutyrate with calcium benzoate (Popoff, B. 6, 1255). Liquid. Yields benzoic acid, CO₂, and HOAc on oxidation.

Oxim.—Ph.C(NOH)Pr. [58°]. Plates (from

ligroin) (Rattner, B. 20, 506).

PHENYL PROPYL KETONE CARBOXYLIC ACID v. Ethyl-benzoyl-acetic acid, vol. i. p. 482.

Phenyl isopropyl ketone o-carboxylio acta, Vol. 1. p. 482.

C₁₁H₁₂O₃ i.e. Pr.CO.C₆H₄.CO₂H. [121°]. Formed by boiling isopropylidene-phthalide with alcohol and KOHAo (Roser B. 17, 2777). Crystals.

and KOHAq (Roser, B. 17, 2777). Crystals. PHENYL PROPYL METHYL KETONE CARBOXYLIC ACID CH_Ph.CHAc.CH_CO_H. [99°]. (235° at 40 mm.). Formed by reducing benzylidene-β-acetyl-propionic acid with sodiumamalgam in presence of H₂SO₄ (Erdmann, A. 254, 203). Needles. It yields on oxidation CHMe CH(C,H₇) CH₂ [86°].— CaA'₂ Saq.—AgA'.

PHENYL METHYLENE PROPYL DI-KETONE v. BUTYRYL-ACETOPHENONE.

PHENYL-PROPYL METHYL KETONE

CH₂.CO.CHMe.CH₂Ph. (239°). Formed by distilling calcium acetate with calcium phenylisobutyrate (Von Miller, B. 23, 1884). Oil.

PHENYL - PROPYL METHYL KETONE CARBOXYLIC ACID v. BENZYL-METHYL-ACETOACETIC ACID, vol. i. p. 25.

Di-phenyl-isopropyl methyl ketone tricarboxylic acid. Ethyl ether of the dinitrile CO₂Et.C(CH₂.C₀H₄.CN)₂CO.CH₃. [120°]. Formed from sodium acctoacetic ether and the nitrile [1:2;CH₂Cl.C₆H₄.CN (Gabriel a. Hausmann, B. 22, 2018). Prisms (from alcohol).

Hexa-phenyl-isopropyl methyl ketone carboxylic acid C₄₂H₁₈O₃ i.e. (CPh₃)₂C(CO₂H).CO.CH₃. [160°]. Formed from sodium acetoacetic ether and BrCPh₃ (Allen a. Kölliker, A. 227, 111). Crystals (from ether).

PHENYL-PROPYL-PYRAZOLE $C_{12}H_{14}N_2$ i.e. NPh $< \frac{C(C_3H_7):CH}{CH}$. (280°). S.G. 15 1.0435. Formed from butyryl-acetic aldehyde and phenylhydrazine (Claisen a. Stylos, B. 21, 1148). Liquid.

PHENYL PROPYL SULPHONE Ph.SO₂.Pr. [44°]. Formed by the action of KOHAq at 100° on Ph.SO₂.CHEt.CO₂H [124°], which is made from a-bromo-butyric acid and sodium benzene sulphinate (Michael a. Palmer, Am. 7, 67; cf. Otto, B. 21, 998). Pearly plates, v. sol. alcohol.

TRI . PHENYL . DI . PROPYL . DI . THIO. BIURET C.S.N., Ph., Pr. [154°]. Formed by the action of aniline (1 mol.) on NPhPr.CSCl (2 mols.) (Billeter a. Strohl, B. 21, 109). Yellow needles.

PHENYL - PROPYL - THIO - CARBAMIC CHLORIDE NPhPr.CS.Cl. [36°]. Formed from propyl-aniline and CSCl₂ (Billeter a. Strohl, B 21, 102). Prisms (from ligroin).

DI-PHENYL-ISOPROPYL-THIO - SEMI-CARBAZIDE NHPh.CS.NH.NPhPr. [116°]. Formed from phenyl-thiocarbimide and phenyl-isopropyl-hydrazine (Michaelis a. Philips, A. 252, 280). Crystalline.

PHENYL-PROPYL-THIO-UREA C₁₀H₁₄N₂S i.e. NHPh.CS.NHC₃H₂. [63°]. Formed from aniline and propyl-thio-carbimide and also from propyl-amine and phenyl thiocarbimide (O. Heoht, B. 23, 286). Pearly plates (from dilute alcohol).

Di-phenyl-propyl-thio-urea CSN₂HPh,Pr. [104°]. Formed by heating propyl-aniline with phenyl-thio-carbimide on the water-bath (Billeter a. Strohl, B. 21, 109). Needles.

Di-phenyl-di-propyl-thio-urea CS(NPhPr),. [103°]. Formed from propyl-aniline and CSCl, (B. a. S.). Large plates.

DI-PHENYL-PROPYL-UREA C_{1a}H_{1a}N₂O i.e. NH_.CO.NH.CH₂.CHPh.CH₂Ph. [112°]. Formed from di-phenyl-propylamine hydrochloride and potassium cyanate solution (Freund a. Remse, B. 23, 2861). Needles, sol. alcohol a.ld ether. PHENYL-PYRAZINE. Hexahydride

PHENYL-PYRAZINE. Hexahydride NPh CH₂.CH₂ NH. Formed by heating bromo-benzene (3 pts.) with pyrazine hexahydride (8 pts.) at 270° (Schmidt a. Wichmana, B. 24, 3239). Thick liquid, v. sol. water. Di-phenyl-pyrazine C18H12N2, i.e.

N≪CH:CPh CPh.CH>N. Iso-indole. [196° cor.]. V.D. 7.95 (Treadwell a. Meyer, B. 16, 342). Formed by the action of cold alcoholic NH₃ on w-bromo-acetophenone or on amido-acetophenone (Staedel a. Kleinschmidt, B. 11, 1744; 13, 837; Möhlau, B. 18, 163; Wolff a. Röders, B. 20, 432; Goedeckemeyer, B. 21, 2687). Broad yellow needles (from alcohol). — B'2H2PtCla. Bronzed plates (from HClAq) quickly decomposed by water.

Hexahy dride C₁₆H₁₈N₂. [109°] and [123°]. Formed by reducing di-phenyl-pyrazine with Na and isoamyl alcohol. Occurs in two forms. The (a)- variety [123°] crystallises from ligroin in white needles, and yields B"2HCl [c. 310°], B"H2PtCl aga, and a nitrosamine C18H16N6O, [143°]. The (\(\beta\))-variety [109°] crystallises from alcohol in white needles and yields B" 2HCl

[c. 295°] and B"H2PtCl6 2aq.

295°] and D Har took 2011.

Di-phenyl-pyrazine N CPh.CPh N. [119°]. (c. 340°). Formed by distilling the dihydride (M.). Needles (from ligroin), ppd. by adding water to its solution in HClAq.—B'2H2PtCl6.

 $N \leqslant_{CH_2.CH_2}^{CPh.CPh} N.$ Dihydride [181°]. Formed by boiling benzil with ethylene-diamine and alcohol (Mason, B. 20, 268; C. J. 55, 97). Prisms (from alcohol), insol. water. Split up by HClAq into the parent substances.

Di-phenyl-pyrazine. Hexahydride v.

DI-PHENYL-DI-ETHYLENE-DIAMINE.

Tetra - phenyl - pyrazine $N \leqslant_{CPh,CPh}^{CPh:CPh} N$. Di - tolane - azotide. [241°] (P.); [247°] (G.). Formed by heating benzoin with ammonium acetate (vol. i. p. 477). Formed also by reducing the di-oxim of di-phenyl-glyoxal in alcoholic solution by sodium-amalgam (Polonowska, B. 21, 489), and by heating benzoic aldehyde with CPhH(NH₂).CPhH(NH₂) (Grossmann, B. 22, White needles (from HOAc), v. sl. sol. alcohol. Conc. H₂SO₄ gives a blood-red colour. PHENYL-PYRAZOLE C,H,N, i.e.

NPh.CH N=CH>CH. [11°]. (247° cor.). 1.113. Formed by boiling epichlorhydrin with phenyl-hydrazine and benzene (Balbiano, G. 17, 176; 18, 354). Got also by distilling its tricarboxylic acid (Knorr a. Laubmann, B. 22, 180). Oil, sol. alcohol and ether. - B'2H2PtCl62aq: needles, decomposing at 175°.

Ethylo-iodide B'EtI. [117°]. Prisms. Dihydride N-CH₂ CH₂. [52°]. (274°

at 754 mm.). Formed from phenyl-hydrazine (120 g.) dissolved in ether (600 g.) by addition of arcolein (50 g.) in ether (100 g.) (Fischer a. Knoevenagel, A. 239, 196). Sl. sol. hot water, v. sol. alcohol, ether, and benzene. Sl. sol. dilute acids, sol. conc. HCl. May be distilled with steam. K₂Cr₂O₇ colours its solution in dilute H21'O4 reddish-violet, or, in very dilute solutions, blue. Gives no indole derivative when fused with ZnCl2

Reference. - DI-BROMO-PHENYL-PYRAZOLE DI-

HYDRIDE.

NPh.CH₂ CH₂ (210° at Tetra-hydride

165 mm.). Formed from tri-methylene bromide, sodium phenyl-hydrazine, and benzene (Michaelis a. Lampe, B. 24, 8738). Liquid. — B'C_sH₃N₃O₇. Short yellow needles.

Di-phenyl-pyrazole C15H12N2 i.e.

CH : CH NPh or CH: CPh NPh. [56°]. (336°). Formed by distilling its carboxylic acids (Beyer a. Claisen, B. 20, 2187; Knorr a. Laubmann, B. 21, 1212; 22, 176). White crystals.

Dihydride C₁₅H₁₄N₂. [136°]. Formed by reducing the base. Small needles. An isomeric or identical body [138°], got by distilling the phenyl-hydrazide of cinnamic aldehyde; differs by fluorescing slightly in alcoholic solution.

Tri-phenyl-pyrazole $C_{21}H_{18}N_2$. [206°]. A product of the action of phenyl-hydrazine on anhydro - acetophenone-benzil (Japp a. Klingemann, C. J. 57, 709). Stellate groups of needles.

Tri-phenyl-pyrazole CPh:NNPh. [138°]. Got by warming CH2Bz2 with phenyl-hydrazine (Knorr a. Laubmann, B. 21, 1205). Plates (from ether). Weak base.

Methylo-iodide B'Mel. [176°].

CH, CHPh CPh—N NPh. [135°]. Got Dihydrideby reduction or by warming benzylidene-acetophene with phenyl-hydrazine and alcohol. Yields C₂₁H₁₂Br₃N₂ [179°]. PHENYL-PYRAZOLE CARBOXYLIC ACID

 $C_{10}H_8N_2O_2$. [220°]. Formed by distilling the tricarboxylic acid (Knorr a. Laubmann, B. 22, 180). Needles (from Aq), v. sol. alcohol, sl. sol. ether. May be sublimed. AgA': amorphous pp.

Phenyl-pyrazole dicarboxylic acid $CH:C(CO_2H):N$ NPh. [256°] (B.); [266°] (C. a. C(CO₂H):N R.). Formed by oxidising phenyl-di-methylpyrazole or phenyl-methyl-pyrazole carboxylic acid with alkaline $KMnO_4$ (Balbiano, B.23, 1449; Claisen a. Roosen, B.24, 1892). Small white plates in spherical groups (from alcohol).-

(NH₄)₂A". [212°].—PbA".—Ag₂A".

Methyl ether MeA". [128°]. Amide C, N2HPh(CO.NH2)2. [190°]. Phenyl-pyrazole tri-carboxylic acid

CO2H.Q = CO₂H.C:C(CO₂H)>NPh. [184°]. Formed by oxidising phenyl-methyl-pyrazole dicarboxylic acid with alkaline KMnO4 (K. a. L.). White needles (containing aq) (from ether), sol. water and alcohol.—BaHA" ½aq: plates.

Di-phenyl-pyrazole carboxylic acid

CPh = N > NPhC₁₆H₁₂N₂O₂ CO₂H.C—N CH:CPh>NPh. Got by saponification of

its ether, which is obtained by boiling benzoylpyruvic ether CH2Bz.CO.CO2Et (20 pts.) with HOAc (100 pts.) and phenyl-hydrazine (10 pts.) (Beyer a. Claisen, B. 20, 2185). Crystals (containing EtOH), which melt at 185°, v. sol. acetone.

Ethyl ether EtA'. [90°]. (c. 400°). Di-phenyl-pyrazole dicarboxylic acid

 $C_{17}H_{12}N_2O_4$ i.e. $CO_2H.C.C(CO_2H)$ NPh. [218°]. Got by oxidation of di-phenyl-methyl-pyrazole carboxylic acid (Knorr a. Laubmann, B. 22, 175). Satiny needles (containing aq) (from HOAc).— NH,HA". [c. 270°].—BaA" aq.—CaA" 2aq.

References .- Oxy- and Oxy-AMIDO- PHENYL-PYRAZOLE CARBOXYLIC ACID.

DI-PHENYL-PYRAZOLONE OXY-DI-

PHENYL-PYRAZOLE.

PHENYL-PYRAZYL METHYL KETONE C₂N₂H₂Ph.CO.CH₂. [122° cor.]. Formed by heating v-phenyl-pyrazole with AcCl at 150° (Balbiano, G. 19, 136). Small needles. Yields an oxim [181°] and a phenyl-hydrazide [144°]
TETRA-PHENYL-PYRIDAZINE DIHY

DIHY.

DRIDE C₂₈H₂₂N₂ i.e. CH₂·CPh:N CPh:CPh.NPh ? [149°].

Formed from desyl-acetophenone, phenyl-hydrazine, and HOAc (Smith, C. J. 57, 649). Yellow needles (from alcohol), v. sol. hot HOAc. According to Klingemann (A. 269, 106), this body is probably phenyl-amido-tri-phenyl-pyrrole, since on boiling with HOAc it yields tetraphenyl-pyrrole.

(a)-PHENYL-PYRIDINE $N \leqslant_{CH:CH}^{CPh.CH} >_{CH}$. (270°). A product of the distillation of its dicarboxylic acid with lime (Skraup a. Cobenzl, M. 4, 472). Heavy oil. Yields picolinic acid on oxidation.—B'2H2PtCl, 2aq.—Pi crate [c. 170°].

 $N \leqslant_{CH:CH}^{CH.CPh} > CH.$ (\$)-Phenyl-pyridine (270°). Formed by distilling its carboxylic acid with lime (Skraup a. Cobenzl, M. 4, 453). Got also by heating pyrrole with benzylidene chloride and NaOEt (Ciamician a. Silber, B. 20, 191). Heavy oil. Yields nicotinic acid on oxidation. — B'₂H₂PtCl₈ 3aq. — B'C₈H₃N₈O₇. [163°]. Groups of yellow needles.

N≪CH.CH >CPh. (γ) -Phenyl-pyridine [78°]. (275° uncor.). Formed by distilling its tetra-carboxylic acid with lime (Hantzsch, B. 17, 1518). Plates (from water). Yields isonicotinic acid [304°] on oxidation.—B'₂H₂PtCl₂. $-B'_2H_2Or_2O_7$. [155°]. Orange Picrate. [196°]. Yellow needles. Orange needles. -

Hexahydride C.H., PhN. [58°]. (256° at 727 mm.). Got by reducing the base (Bally, B. 20, 2590). Strong base, almost insol. water.-B'HCl. Needles.—B'2H2PtCl. [204°-207°].

v-Phenyl-pyridine hexahydride

NPh CH2.CH2 CH2 Phenyl-piperidine. (249°). Formed by heating piperidine with iodobenzene or bromo-benzene at 260° (Lellmann, B. 20, 680; 21, 2279. Alkaline oil, v. sol. alcohol and ether. -B', H, PtCl, 2aq: needles or plates.—B'H2Cr2O7: plates.

References .- AMIDO- and NITBO- PHENYL-

PIPERIDINE.

 $Di-(\alpha)$ -phenyl-pyridine $C_{17}H_{18}N$ i.e.

N CPh:CH OH. [82°] (P. a. S.); [73°] (D.). Formed by distilling its carboxylic acids with soda-lime (Paal a. Strasser, B. 20, 2764; Doebner, A. 249, 121). Yellow needles, v. sol. Yields hexahydride.alcohol. an oily B',H,PtCl,. [205°]. - B'HAuCl. [202°]. — B'4H4Cr4O11. Orange-red needles.

Methylo-iodide B'MeI. [203°]. Tri-phenyl-pyridine C₂₂H₁₁N. [135°]. Formed by passing NH, through acetophenone, and adding P.O. (Engler, B. 6, 639; 19, 40; A. 238, 37).—B',H,PtCl.

Reference. - OXY-PHENYL-PYRIDINE.

Vol. IV.

PHENYL-PYRIDINE o-CARBOXYLIC ACID $C_{12}H_{\bullet}NO_{2}$ i.e. $N \leqslant \begin{array}{c} CH:C(C_{\bullet}H_{\bullet}.CO_{2}H) \\ CH.CH = CH \end{array} >$. [185°]. Formed by heating the dicarboxylic acid at 185°

(Skraup a. Cobenzl, M. 4, 450). Needles, sl. sol. cold water. Yields nicotinic acid on oxidation .-CaA', 2aq: long needles, v. sol. cold water.

(α)-Phenyl-pyridine dicarboxylic acid

C₁₂H₂NO₄ i.e. N C(C₆H₄.CO₂H):C.CO₂H [280°-Formed by oxidation of (a)-naphthoquinoline (Skraup a. Cobenzl, M. 4, 463). Minute crystals, gives off indigo-blue vapours when heated. When distilled with CaO it yields (a)-phenyl-pyridine and the compound $C_{12}H$, NO [140°-142°] (315°), which crystallises from alcohol in yellow plates and yields B'2H2PtCl. Yields a di-bromo- derivative [205°].

Salts. — CaA" 2aq. — CuA" 4aq: violet stalline pp. — Ag_2A " $1\frac{1}{2}$ aq. — H_2A "HCl. crystalline pp. -

 $(H_2A'')_2H_2PtCl_6$ 3aq.

(β)-Phenyl-pyridine dicarboxylic acid $N \leq \stackrel{\stackrel{\leftarrow}{C}(CO_2H):\stackrel{\leftarrow}{C}:C_1H_4.CO_2H[1:2]}{CH:\stackrel{\leftarrow}{C}H:\stackrel{\leftarrow}{C}H}$. [207°]. Formed by oxidation of (β)-naphthoquinoline (Skraup a. Cobenzl, M. 4, 442). Prisms (containing aq), sl. sol. cold water. Salts. — HA'HCl. —

 $(HA')_2H_2PtCl_6 2\frac{1}{2}aq. - K_2A'' 3aq. - KHA'' 2aq. - CaA'' 3aq. - BaA'' 4\frac{1}{2}aq. - CuA'' 4aq. - Cu_2H_2A''_8.$ -AgH,A"2.

Sulphonic acid C18HeNSO, i.e. $N \leqslant C(CO_2H):C.O_6H_3(SO_3H)(CO_2H)[1:2:6]$ CH.CH:CH

Formed, as well as C₁₂H₆(SO₃H)NO aq, by oxidation of (β)-naphtho-quinoline sulphonic acid with alkaline KMnO₄ (Immerheiser, B. 22, 405). Crystals (from water). $-K_sA'''$. $-Ba_sH_sA''$ Pb₄ $A'''_2(OH)_2$. $-Ag_sA'''$ (dried at 100°).

The by-product C₁₂H₄(SO₂H)NO aq yields KA' aq, BaA'₂ 2aq, PbA'₂ 8aq, AgA' aq, an oxim [290°] and a phenyl-hydrazide [c. 295°].

Phenyl-pyridine dicarboxylic acid

N CH:C(CO₂H) CPh. [230°] (hydrated); [246°] (anhydrous). Formed by heating the tetra-carboxylic acid at 170° (Weber, A. 241, 13). Greenish-yellow plates (containing aq). -CuA" 2aq : blue pp.

 $\begin{array}{c} (\gamma)\text{-Phenyl-pyridine tetra-carboxylic acid} \\ N \leqslant \stackrel{C(CO_2H)\cdot C(CO_2H)}{\subset} \text{CPh.} \quad [207^\circ]. \quad \text{Formed} \end{array}$ by oxidation of the phenyl-di-methyl-pyridine dicarboxylic acid, which is got from acetoacetic ether, NH, and benzoic aldehyde (Hantzsch, B. 17, 1515). Crystals (containing 3aq), v. sol. water. KH, A'vaq: small prisms or pyramids, sl. sol. cold water.—Ba,(NH4),A'v, 6aq: minute tables. -Cu2A'v 7aq.

Di-phenyl-pyridine (γ) -carboxylic acid

N CPh:CH C.CO.H. [275°]. Formed by heating (CH_Bz)₂C(CO₂H)₂ with alcoholic NH₃ at 120° (Paal a. Strasser, B. 20, 2761). Needles (from alcohol).—AgA': pp., sl. sol. hot water.

Dihydride C₁₈H₁₈NO₂ i.e.

NH CPh:CH CH.CO.H. Crystalline flakes.— NH,A'. [270°]. Formed from (CH,Bz),CH.CO,H and alcoholic NH, in the cold. Needles, v. sol. water. G

Hexahydride

NH CHPh.CH₂ CH.CO₂H. [339°]. Formed, in small quantity, when di-benzoyl-di-methylmalonic acid is heated with alcoholic NH, at 120° (P. a. S.). Crystals, v. sl. sol. water. Yields a nitrosamine [159°].

Di-phenyl-pyridine tricarboxylic acid

 $N < C(C_0H_0.CO_2H):C.CO_2H$ [250°]. Formed by oxidising phenyl-naphthoquinoline carboxylic acid with alkaline KMnO, below 50° (Doebner, A. 249, 120). Needles (from dilute alcohol).-Ag, A": white flocculent pp.

At 100° the chief product is phenylene phenylpyridylene ketone carboxylic acid C18H11NO2 [226°], which on distillation with soda-lime yields $CO < \stackrel{C_6H_4}{C_5H_5PhN}$ [68°], which forms an oxim [84°]

and B'2CrOs.

Reference.—OXY-PHENYL-PYRIDINE CARBOXYLIO ACID

PHENYL PYRIDYL KETONE C12HaNO i.e. C₃H₃.CO.C₃H₄N. (307°). Formed by heating its carboxylic acid (Bernthsen a. Mettegang, B. 20, 1209). Liquid. Yields a phenyl-hydrazide

[143·5°].—B',H,PtCl,.
PHENYL (β)-PŸRIDYL KETONE CARBOXYLIC ACID C,H,CO.C,H,N.CO.H. [147°]. Formed from quinolinic anhydride, benzene, and AlCl. (Bernthsen a. Mettegang, B. 20, 1209). Prisms, v. e. sol. hot water.

 $co<_{CH:CPh}^{CH:CPh}>0.$ DI-PHENYL-PYRONE

[139.5°]. Formed by heating its carboxylic acid (Feist, B. 23, 3734). Got also by heating dehydro-benzoyl-acetic acid with HClAq at 245°. Needles resembling asbestos (from benzene). Its solution in conc. H₂SO₄ exhibits violet fluorescence. Alcoholic NH₃ converts it, on heating, into oxy-di-phenyl-pyridine.

DI-PHENYL-PYRONE CARBOXYLIC ACID $co < C(CO_{\bullet}H):CPh > 0.$ [201°]. Formed by warming chloro-dehydro-benzoyl-acetic acid (1 pt.) in H₂SO₄ (3 pts.) at 140° (Feist, B. 23, 8730). Small crystals (from benzene), v. sol. chloroform.—NH₄A'. [135°].—BaA'₂6aq.—

Ag₂A'₂AgNO₈: curdy pp. Di-phenyl-pyrone dicarboxylic ether

CO C(CO,Et):CPh O. [140°]. Formed by allowing a mixture of cupric benzoyl-acetic ether (24 g.), COCl₂ (10 g.), and toluene (40 g.) to stand for six weeks (F.). Got also from acetone dicarboxylic ether, Et₂O, Na, and BzCl (Dünsch-mann a. Pechmann, A. 261, 189). Plates, v. sol, most solvents.

PHENYL-PYROTARTARIC ACID CeHs.CH2.CH(CO2H).CH2.CO2H. Benzyl-succinic acid. [161]. Got by heating phenyl-propane tricarboxylic acid and by reducing phenyl-itaconic acid (Bischoff, B. 23, 653; Fittig a. Roeders, A. 256, 87; cf. Baeyer a. Perkin, B. 17, 449). White plates, sl. sol. cold water. Yields an anhydride [102°].—BaA" 12aq.—CaA".—Ag.A".

Phonyl-pyrotartaric acid CO.H.CHPh.CHMe.CO.H. Two varieties [1719] and [193°] are formed by saponifying the product of the reaction of NaOEt on a mixture of s-cyano-propionie acid and a-bromo-phenylacetic ether (Zelinsky a. Buchstab, B. 24, 1877). At 320° it forms an anhydride which on treatment with water yields a mixture of the two isomeric phenyl-methyl-succinic acids.

Reference.— OXY-AMIDO-PHENYL-PYROTARTARIO

PHENYL-PYROXIMIDINE DIHYDRIDE

CH₂ CH₂ O CPh. Formed from γ-bromepropyl-benzamide by shaking with hot water (Gabriel, B. 24, 3214). Liquid, sl. sol. cold water.—B'₂H₂PtCl₈. [185°].—B'C₈H₂N₃O₇. [151°].

NH CPh:CH DI-PHENYL-PYRROLE Formed by the action of alcoholic potash on its carboxylic ether (Paal, B. 21, 3061). Got also by heating the dicarboxylic acid C₄NH₃(C₅H₄.CO₂H)₂ with lime (Baumann, B. 20, 1490). Plates (from HOAc or dilute alcohol). Forms a red solution in H2SO4. Colours pinewood, moistened with HClAq, red.

Tri-phenyl-pyrrole NPh CPh.CH [229°]. Formed by heating its carboxylic acid or its dicarboxylic acid C₄NH₂Ph(C₆H₄.CO₂H)₂ with lime (P.; B.). Needles, sl. sol. ether and alcohol.

Tri-phenyl-pyrrole NH<CPh:CPh . [141°]. Formed by heating desyl-acetophenone with alcoholic NH, at 150° (Smith, C. J. 57, 645). Slender needles with violet fluorescence.

Tetra-phenyl-pyrrole $C_{28}H_{21}N$ i.e.

NH<CPh:CPh [214.5°]. Formed from both bidesyls by heating with alcoholic NH, at 150° (Garrett, B. 21, 3107; Fehrlin, B. 22, 553; Magnanini a. Angeli, B. 22, 855). Got also by distilling its phenyl-v-amido- derivative (Klingemann, A. 269, 122). Needles or plates.

Acetyl derivative. [226]. Needles.

Tetra-phenyl-pyrrole NPh CPh:CPh

[197°]. Formed by boiling desyl-acetophenone with aniline and HOAc (Smith, C. J. 57, 646). White silky needles, v. sl. sol. cold alcohol. DI-PHENYL-PYRROLE CARBOXYLIC ACID

NHCPh:C.CO₂H. [261°]. Formed by saponifying the ether, which is got by the action of alcoholic NH, or of boiling NH,OAc and HOAc on di- β -benzoyl-propionic ether (Kapf a. Paal, B. 21, 1491, 3061). Orange needles, sl. sol. alcohol.

Ethyl ether EtA'. [159°]. Needles. [217°]. $A mide C_1, H_1, N_2O$.

 $\begin{array}{c} \textbf{Di-phenyl-pyrrole} \\ \textbf{NH} < \begin{matrix} \textbf{C(C_0H_1.CO_2H):CH} \\ \textbf{C(C_0H_1.CO_2H):CH} \end{matrix}. \end{array}$ di-o-carboxylic [232°]. Formed by heating $C_2H_4(CO.C_8H_4,CO.2H)_2$ with alcoholic NH_8 (Gabriel, B. 19, 840). Slender needles. Yields a nitrosamine $C_{18}H_{12}N_2O_8$ [o. 210°] crystallising from HOAc.

Tri-phenyl-pyrrole carboxylic acid

NPh CPh:C.CO₂H. [273°]. Formed by sapenifying the ether, which is got by boiling dibenzoyl-propionic ether with aniline and HOAc

(Paal, B. 21, 3061). Needles (from HOAc). Ethyl ether Eth'. [170°]. Needles. Tri-phenyl-pyrrole di-earboxylic acid NPh < 0(0, H, 00, H): CH [295°]. Formed by warming C,H4(CO.C,H4.CO,H)2 with aniline (Baumann, B. 20, 1487). Crystals, insol. water. Ag2A": bulky pp.

Ethyl ether Et, A". [122°]. Needles.

DI - a - PHENYL - B - PYRROYL - PROPIONIC [216° ACID C,H,N.CO.CH2.CPh2.CO2H. Formed by saponification of di-phenyl-pyrrylcrotolactone CH:C(C,H,N) O [184°] which is got by warming (a)-pyrryl methyl ketone with benzil and conc. KOHAq (Angeli, B. 23, 1356). Colourless crystals, almost insol. water.—KA'.-AgA': light-yellow pp.

PHENYL PYRRYL KETONE C₁₁H₂NO i.e., H₄.CO.C₆H₅. [78°]. Formed by heating NC,H,.CO.C,H,. [78°]. Formed by heating pyrrole with NaOBz and Bz,O (Ciamician a. Dennstedt, B. 17, 2955). Yields C,1H,AgNO.

PHENYL-PYRUVIC ACID v. PHENYL-GLY-

CH:N C₆H₄<N=CPh PHENYL-QUINAZOLINE

[101°]. Got by distilling o-amido-benzyl-benzamide (Gabriel a. Jansen, B. 23, 2810). Yellow needles. Weak base.

Phenyl-quinazoline. Dihydride.

 $C_{14}H_{12}N_2$ i.e. $C_6H_4 < \stackrel{CH_2.NPh}{N=CH}$. [95°]. Formed by reducing [1:2]C₆H₄(NO₂).CH₂.NPh.CHO with zine and HClAq (Paal a. Busch, B. 22, 2686; Donner, Ph. [3] 20, 43). Six-sided tables (from ether-ligroin), almost insol. water and alkalis. Split up into aniline and benzonitrile by heating with zinc-dust.—B'HCl 2aq. [80°] when hydrated; [221°] when anhydrous. Concentric needles, sol. water.—B'HSnCl₂. [130°-134°]. White crystals.—B'₂H₂PtCl₆. [208°]. Yellow crystals.—B'₂H₂SO₄ 2aq. [79°]. Melts at 140°-143° when anhydrous.

B'MeI. [170°] and Methylo-iodide [180°].—B'MeI₂. [157°]. Golden plates.

C.H. CH. NPh NH.CH. Tetrahydride Got by reduction. Needles. Yields a crystalline acetyl derivative and a brick-red nitrosamine.

Di-phenyl-quinazoline. Dihydride. C₂₀H₁₆N₂ i.s. C₆H₄ CH₂.NPh [115°]. Formed by reduction of the benzoyl derivative of o-nitrobenzyl-aniline in HOAc with tin and HCl (Lellmann a. Stickel, B. 18, 1608). Small needles,

v. sol. alcohol.—B'HCl: silky needles.

Reference.—Oxy-PHENYL-QUINAZOLINE. (B. 2)-PHENYL-QUINOLINE

OH: OH.O.N=CH. p-Phenyl-quinoline. [111]. (260° at 77 mm.). S.G. 20 1.1945. Prepared by heating p-amido-diphenyl with glycerin, nitrobenzene, and H₂SO₄ (La Coste, B. 15, 562; A. Trimetric plates (from ether). B',H,PtCl. Or B',H,Cr,O, [136 [153°]. Needles. Orange crystalline powder. — [136°]. — Tartrate B'C, H₆O₆ 3aq.

Methylo-iodide B'MeI 2aq. [194°]. Ethylo-iodide B'EtI 2aq. (169°7.

Tetrahydride C.H.PhN. Got by reduc-n. Unstable pp. Yields a nitrosamine tion. Unstable pp. Yields a nitrosamine C,H,Ph(NO)N [112°], an acetyl derivative [100°], and a benzoyl derivative [187°].—B'HCl 13aq. [204°].—B'C.H.N.O. [165°].

CH:CH .C.OH:CH CH:CPh.C.N—CH (B. 4)-Phenyl-quinoline o-Phenyl-quinoline. (273° at 80 mm.). Got in like manner, using o-amido-diphenyl (C.). Oil, with yellowish-green fluorescence. B'zHzPtClz -B'₂H₂Cr₂O₃. [126°]. Orange plates.

Methylo-iodide B'Mel. [163°]. Crystals,

v. sol. water.—B'2Me2PtCl, [193°].

(Py. 1)-Phenyl-quinoline $C_{s}H_{\bullet} < \stackrel{CPh:CH}{N} = CH$ [62°]. Got by heating its carboxylic acid (Königs a. Nef, B. 19, 2430). Needles (from ether or ligroin). Solutions of its sulphate and fluoresce hydrochloride blue. — B'H₂CrO₄: needles.—B'2H2PtCl6: yellow four-sided tables.

(Py. 2)-Phenyl-quinoline $C_{\mathfrak{g}}H_{\mathfrak{g}} < CH:CPh$ Formed by adding a little NaOH to a solution of o-amido-benzoic aldehyde and phenyl-acetic aldehyde in dilute alcohol (Friedländer a. Gohring, B. 16, 1836). Oil, sparingly volatile with steam. -B'HCl, [93°]. -B'2H2PtCl

(Py. 3)-Phenyl-quinoline $C_{e}H_{4} < \frac{CH:CH}{NCPh}$.

[86°]. (above 360°)

Formation.—1. By heating cinnamic aldehyde (30 pts.) with aniline (20 pts.) and conc. HClAq (20 pts.) for two hours at 210° (Grimaux.) C. R. 96, 584; Doebner a. Miller, B. 16, 1665; 19, 1194). — 2. By warming o - amido - benzoic aldehyde with acetophenone, dilute alcohol, and a little NaOH (Friedländer a. Gohring, B. 16, 1835).—8. By distilling m-oxy-phenyl-quinoline or di-oxy-phenyl-quinoline with zinc-dust (Miller a. Kinkelin, B. 18, 1908; Weidel, M. 9, 151).-4. By distilling its carboxylic acid with soda-lime (Doebner, B. 20, 280; A. 242, 294; Pfitzinger, J. pr. [2] 38, 583)

Properties .- Silky needles (from dilute alsol. ether, sl. sol. water. NHBz.C₆H₄.CO₂H [1:2] [182°] on oxidation by

KMnO, in acid solution.

Salts.—B'H_PtOl₂ 2aq.—B'HAuCl₄. [160°].
—(B'HCl)_AuCl₃. [204°]. Needles.—B'H₂Cr₂O₇;
stable golden plates.—B'C₂H₃N₂O₇. [188°]. Alkylo-iodides B'MeI. [197°].-B'EtI.

[195°].—B'₂Et₂PtCl₆.—B'EtCl 2aq.—B', Et₂PtCl₆.

Tetrahydride C₁₅H₁₆N. (342°). Got by reducing the base with tin and HClAq (Doebner, B.19,1198). Oil, yielding an oily nitrosamine. B'HCl. Needles (from alcohol), v. sl. sol. water.

(Py. 1,3)-Di-phonyl-quinoline

 $C_6H_4 < \begin{array}{c} CPh:CH \\ N = CPh \end{array}$. [112°]. Formed by heating the anilide of CH₂Bz₂ with H₂SO₄ (Beyer, B. 20, 1772). Mass of white crystals.—B'₂H₂PtCl₆ 2aq. $-\mathrm{B'H_2SO_4}$. Needles, v. sl. sol. dilute $\mathrm{H_2SO_4}$. $(Py.\ 2,3)$ -Di-phenyl-quinoline

 $C_eH < CH:CPh \\ N = CPh$. [96°], (c. 420°). Formed by reducing [1:2] C.H. (NO2).CH2.CHPh.CO.C.H. with iron and HOAc (Buddeberg, B. 23, 2075). Large crystals, sol. alcohol. - B'2H2PtCl

(Py. 2)-PHENYL-ISOQUINOLINE

O₁₅H₁₁N i.e. C₆H₄ CH:CPh. [105°]. Obtained by reduction of the (Py. 4)-chloro-derivative by heating it with HI and P at 170° for three hours; also by distilling phthalimide with zinc-dust (Gabriel, B. 18, 3477). Rhombic plates. Slightly volatile with steam .- B' H_Cl_PtCl : needles.

 $Tetrahydride C_{e}H_{4} < \substack{CH_{2}.CHPh.\\CH_{2}.NH}$ [45°-

48°]. Formed by reduction of an alcoholic solution of the (Py.)-mono- or di- chloro-derivatives by boiling with (3 p.c.) sodium-amalgam (Gabriel, B. 18, 3479). Crystals, v. sol. alcohol.

References. - AMIDO-, CHLORO-, CHLORO-NITRO-, NITRO-, OXY-AMIDO-, and OXY- PEENYL-QUINOLINE.

(Py. 1) - PHENYL - QUINOLINE (Py. 3)-CARBOXYLIC ACID C18H11NO2 i.e.

C.H. CPh:CH Got by oxidation of phenyl - methyl - quinoline - phthalone with chromic acid mixture (Kœnigs a. Nef, B. 19, 2428). Yellow needles (from dilute alcohol), sol. acids and alkalis.

(Py. 3)-Phenyl-quinoline (Py. 1)-carboxylic acid $C_0H_1 < N = CPh$. [207°]. Formed by warming pyruvic acid with benzoic aldehyde, aniline, and alcohol (Doebner, B. 20, 280; A. 242, 291). Formed also by heating isatin with acetophenone and alcoholic potash (Pfitzinger, J. pr. [2] 83, 538). Needles, m. sol. cold alcohol. Yields phenyl-quinoline [84°] on distillation with soda-lime. — (HA')₂H₂PtCl₈. — PbA'₂aq.— SuA', aq. - ZnA', aq. - AgA': white pp.

Phenyl - quinoline - o - di - carboxylic acid $C_9H_4(O_6H_5)N(CO_2H)_2 \quad \text{i.e.} \quad C_6H_4 < \begin{matrix} \mathrm{CPh:C.CO_2H} \\ \mathrm{N} = \mathrm{C.CO_2H} \end{matrix}$ Formed by oxidation of phenyl-acridine with KMnO.

Salts.—BaA"4aq: glistening needles (Claus **a.** Nicolaysen, B. 18, 2706).

(Py. 2.3) - Di - phenyl - quinoline carboxylic acid C_oH₄ C(CO₂H):CPh . [191°]. Formed by the action of isatin and alcoholic potash on deoxybenzoin (Pfitzinger, J. pr. [2] 38, 583). Needles, insol. water, v. sol. hot alcohol. (Py. 3)-PHENYL-QUINOLINE (B. 2)-SUL-

PHONIC ACID SO, H.C.CH.C.CH.CH HC:CH.C.N=CPh

together with the more soluble (B. 3)-sulphonic acid, by heating (Py. 3)-phenyl-quinoline with H₂SO₄ and SO₅ at 100° (Murmann, M. 13, 60).— Plates, sl. sol. water, almost insol. alcohol. Gives oxy-phenyl-quinoline [238°] when fused with potash at 250°.—BaA'2: plates.—NH,A': plates. -AgA': granules.

(Py. 3) Phenyl-quinoline (B. 3) sulphonic acid. Formed as above (M.). Granules. Yields oxy - phenyl - quinoline [156°] on fusion with potash. - KA'aq. -- BaA', 1 aq: needles.

AgA' $1\frac{1}{2}$ aq: needles.
(B. 2) - Phenyl - quinoline p-sulphonic acid C₁₅H₁₁NSO₂ i.e. [1:4]SO₂H.C₆H₄.C:CH.C.CH:CH HC:CH.C.N-CH

Formed, together with the following more soluble isomeride, by sulphonation of p-phenyl-quinoline (La Coste a. Sorger, A. 230, 30). Needles (containing 2aq). Turns brown, without fusion, at 800°. Alkaline KMnO, gives quinolinic acid and p-sulpl o benzoic acid.—NH, A'. [above 810°].— NaA'aq.—HgA',: white pp.

(B.2) - Phenyl-quinoline (β) - sulphonic acid. Formed as above. Plates (containing aq), not

melted a. 300°.—NH, A': small scales.
PHENYL-DIQUINOLYL-METHANE. Octohydride CHPh(C.H.,N). [153°]. Formed from

quinoline tetrahydride, benzois aldehyde, and ZnCl, in alcohol (Einhorn, B. 19, 1243). Its solution in HClAq is coloured green by FeCl.

Di-phenyl-(B. 1)-quinolyl-methane CHPh₂.C:CH.C.CH:CH

[104°]. Formed by HC:CH.C.N=CH Skraup's method from amido-tri-phenyl-methane (Fischer a. Fränkel, B. 19, 749; A. 241, 364). Prisms, v. sol. alcohol. Yields a nitro-derivative [213°].—B'₂H₂PtCl₆ (dried at 100°).

DI-PHENYL-(Py. 3)-QUINOLYL-UREA C.H.N.NPh.CO.NHPh. [150°]. Formed by heating carbostyril with phenyl isocyanate and benzene at 220° (Goldschmidt a. Meissler, B. 23, 276). Needles (from benzene), sol. HClAq and re-ppd. by NH,

DI-PHENYL-QUINONE C18H12O2 i.e.

CPh CO.CH CPh. [214°]. Formed by heating phenyl methyl diketone with NaOHAq and a little K_sFeCy_s at 100° (Müller a. Pechmann, B. 22, 2130). Orange-yellow plates, m. sol. benzene. Zinc-dust and HOAc reduce it to di-phenylhydroquinone [219°].

DI-PHENYL-QUINOXALINE C.H. N:CPh.

[126°]. Formed by heating benzoin with ophenylene-diamine in the air at 165° (Fischer, B. 24, 720). Needles.

Dihydride $C_6H_4 < NH.CHPh \\ N=CPh$ [149°]. Formed by heating benzoin with o-phenylenediamine at 165° with exclusion of air. Prisms. Its solutions in ether and benzene fluoresce yellowish-green.

Phenylo-hydroxide $C_6H_4 < N = CPh$ [135°].

Formed from benzil and o-phenylene-phenyldiamine (Kehrmann a. Messinger, B. 24, 1239). -C₂₆H₁₉N₂ClFeCl₃. Golden prisms, m. sol. cold water, insol. ether. $-(C_{28}H_{19}N_2Cl)_2PtCl_4$. $-C_{28}H_{19}N_2HgCl_4$. Yellow scales.

Tri-phenyl-quinoxaline. Dihydride

C_sH₄<NPh.CHPh.CPh. (117°]. Got by heating benzoïn with o-phenylene-phenyl-diamine in a sealed tube at 175° (Kehrmann a. Messinger, B. 24, 1875). Crystalline, v. sol. ether. Its solutions fluoresce bluish-green.

Tetra-phenyl-diquinoxaline, so-called. ÇPh:N.C.CH:C.N:CPh [289°]. Formed from CPh:N.C.CH:C.N:CPh tetra-amido-benzene and benzil (Nietzki a. Müller, B. 22, 446). V. sol. hot HOAc. Forms a blue solution in conc. H,SO.

DI-PHENYL-QUINOXALINE CARBOXYLIC **ACID** $C_{21}H_{14}N_2O_2$, i.e. $\begin{bmatrix} 4^2_1 \end{bmatrix}C_0H_4(CO_2H) < N:CPh$ [288°]. Formed by mixing benzil and di-amidobenzoic acid, both dissolved in hot HOAc (Zehra, B. 23, 3627). Yellow plates or needles. BaA'₂Saq: needles, sl. sol. hot water.

Ethyl ether EtA'. [151°]. Needles.

TETRA-PHENYL-DIQUINOXALYL Out H20 N ... i.e. $\stackrel{\mathrm{CPh:N}}{\mathrm{CPh:N}} > C_{e}H_{s}.C_{e}H_{s} < \stackrel{\mathrm{N:CPh}}{\mathrm{N:CPh}}.$ [above 270°]. Formed by mixing acetic acid solutions of benzil and tetra-amido-diphenyl {C,H,(NH,),(1:3:4)}, (Brunner a. Witt, B. 20, 1026). Crystals (from phenol-HOAc). Forms a bright-red solution in | conc. H.SO

PHENYL SELENIDE v. Organic compounds of SELENIUM.

TETRA - PHENYL - SILICANE v. Organic

compounds of SILICON.

PHENYL SILICATE Si(OPh), [48°]. (420° Prepared by heating phenol with SiCl.; the yield being 78 p.c. of the theoretical amount (Hertkorn, B. 18, 1679). Long colourless prisms, v. Decomposed by hot water into sol. alcohol. phenol and silicic acid. Boiling alcohol forms ethyl silicates and phenol.

TRI-PHENYL-STIBINE v. vol. i. p. 294.

PHENYL-STYRYL-HYDRAZINE C, H16N2i.e. CHPh:CH.CH₂.NPh.NH₂. [54°]. Formed from styryl bromide and sodium phenyl-hydrazide (Michaelis a. Claessen, B. 22, 2239). Crystals, v. sol. alcohol and ether.

PHENYL-STYRYL KETONE v. BENZYLIDENE-ACETOPHENONE. Its carboxylic ether is described as Benzylidene-benzoyl-acetic ether, vol. i. p. 481.

PHENYL - SUCCINIC ACID C10H10O4, i.e.

CO.H.CHPh.CH₂.CO.H. [167°].

Formation. - 1. By saponification of the nitrile which is got from ω-chloro-styrene and KCy (Rügheimer, B. 14, 428).—2. By saponification of the product of the action of a-bromophenyl-acetic ether on sodium acetoacetic ether (R.).—3. At 191° from CO₂H.CHPh.CH(CO₂H)₂, got from a-chloro-phenyl-acetic ether and sodium malonic ether (Spiegel, B. 14, 873, 1693; A. 219, 32; Alexander, A. 258, 74).-4. From hydrocornicularic acid by potash-fusion (S.).

Properties.—Needles, sol. hot water and al-cohol, sl. sol. chloroform. Br and PBr_s yield a bromo- derivative [119°] converted by hot water

into phenyl-maleic acid.

Salts.—CaA".—Ag₂A".

Anhydride C10H8O3. [c. 54°].

(a)-Di-phenyl-succinic acid C₁₆H₁₄O₄ i.e. CO₂H.CHPh.CHPh.CO₂H. [222°] (when dry); [183°] (when hydrated). H.C. 1,848,300 (Ossi-

poff, C. R. 109, 223). S. (alcohol) 44 at 21°.

Formation.—1. By heating α-bromo-phenylacetic acid with KCy (Franchimont, B. 5, 1048). 2. Together with the isomeric (β)-acid, by reduction of di-phenyl-maleïc anhydride with sodium-amalgam or Zn and HCl. Separated through the sparing solubility of the Ba salt (Reimer, B. 14, 1802; Anschütz, A. 259, 67).-8. By heating the (β)-isomeride with barytawater at 200° (R.).

Properties. - Prisms (containing aq) or needles (from dilute HOAc). Yields di-phenyl-ethane and di-phenyl-ethylene when distilled with lime.

HClAq at 200° converts it into the (β)-acid. Salts.—BaA" 2aq. S. 32 at 18°.—BaA" 4aq.

-Ag₂A": white pp.

Mono-ethylether HEtA".

wono-ethyl ether HEtA". [140°]. Di-ethyl ether Et,A". [84°]. Needles. [116°]. (240° at Anhydride C16H12O2. 11 mm.). Formed when either of the two s-diphenyl-succinic acids is heated. Got also by heating the (a)- acid with AcCl at 120° (Tillmanns, A. 258, 87). Trimetric crystals; a:b:c = .508:1:.546. V. sol. chloroform. Yields the (a)- acid on boiling with water, and a mixture of (a)- acid (86 p.c.) and (β)- acid (14 p.c.) when

treated with KOHAq. CN.CHPh.CHPh.CN. [160°]. Nitrile

Formed, together with the (3)- isomeride by heating phenyl-acetonitrile with mandelic nitrile and alcoholic KCy at 55° (Chalanay a. Knoevenagel, B. 25, 289). V. e. sol. alcohol. Converted into the (β) - isomeride by boiling with HOAc.

Phenylimide $C_{22}H_{1}$, NO₂. [231°]. Got by heating either the (a)- or the (β)- anhydride with

aniline.

Phenyl-amic acid C₂₂H₁₈NO, i.e. CO₂H.CHPh.CHPh.CO.NHPh. [220°]. Got by boiling the phenylimide with baryta (Anschütz a. Bendix, A. 259, 92).

β)-Di-phenyl-succinic acid

CO₂H.CHPh.CHPh.CO₂H. [230°]. H.C. 1,822,900.

S. (alcohol) 1.124 at 21°.

Formation.—1. By heating its nitrile with HClAq at 200°.—2. By reduction of di-phenylmaleïc anhydride.—3. By heating the a-isomeride with HCAq at 200°.

Properties.—Needles, sol. alcohol, sl. sol. benzene, insol. water. On distillation in vacuo at 250° it yields a mixture of the (a). anhydride (89 p.c.) and the (β) - anhydride (11 p.c.).

Reactions.—1. Yields s-di-phenyl-ethane and di-phenyl-ethylene on distillation with lime (Reimer, B. 14, 1802). - 2. Baryta-water at 200° converts it into the (a)- isomeride.

Salts.—BaA" 7aq. S.21 at c. 18° .—Ag₂A". Ethyl ether Et₂A". [141°]. Needles.

Anhydride CHPh.CO O. [c.

Formed by heating the (β) - acid with AcCl at 100°, and also by the action of AcCl on its salts. When boiled with water it yields the (a)- acid (75) p.c.) and some (β)- acid (25 p.c.). Cold KOHAq forms only the (β)- acid. On heating with aniline it gives the phenylimide [231°].

Diphensuccindone

Formed by heating the acid (5 g.) with conc. H₂SO₄ (8 g.) until dissolved, and then pouring into water (Roser, A. 247, 153). White crystals, insol. water, sol. alcohol. Yields a dioxim C₁₆H₁₀(NOH)₂ [254°] and a phenyl-hydrazide C₁₆H₁₆(N,HPh)₂ [c. 265°]. Reduced by HI and P to C₁₆H₁₁ [100°].

Nitrile CN.CHPh.CHPh.CN. [240°]. Accompanies the (a)- isomeride (v. supra). Colour-

less needles, sol. hot alcohol.

Tetra-phenyl-succinic acid CO₂H.CPh₂.CPh₂.CO₂H. [262°]. Formed by heating a-chloro-di-phenyl-acetic ether with finely-divided silver at 125° (Bickel, B. 22, 1538). Insol. water, v. sol. alcohol and ether.

Ethyl ether Et,A". [89°

Nitrile CN.CPh2.CPh2.CN. [c. 280°]. Formed by the action of NaOEt and I on diphenyl-acetonitrile (Auwers a. V. Meyer, B. 22, 1227; cf. Anschütz a. Romig, A. 233, 349). Formed also by the action of nitrous acid on diphenyl-acetonitrile (Neure, A. 250, 148). Flat needles (from HOAc), v. sl. sol. hot alcohol. Liquefied by heating for a long time at 180°.

Phenyl-isosuccinic acid v. Benzyl-Malonio

ACID. References. - NITBO- and OXY- PHENYL-SUCCINIC ACIDS.

DI-PHENYL-SUCCINIMIDINE C, H, N, i.e. C₂H₄ C(NH) NPh. Formed from ethylene cyanide and aniline hydrochloride at 154° (Blochmann, B. 20, 1856). Greyish-yellow crystals, insol. water. Yields the phenylimide of succinic acid when boiled with HClAq.— B'HCl.—B',H.PtCl.,—B'H.SO.,—B'HBr.—B'HI.

PHENYL-SULPHAMIC ACID C.H, NSO, i.e. C.H.NH.SO.H. Formed as a salt from CISO.H (1 mol.), aniline (3 mols.) and chloroform in the cold. Got also by the action of solid fuming sulphuric acid on aniline in CHCl, at a low temperature (Wagner, B. 19, 1157; Traube, B. 23, 1655; 24, 360).—NaA'.—KA': leaflets, sol. hot alcohol.—BaA'22aq: leaflets or needles, sol. hot water. Decomposed by acids into aniline and H₂SO₁.-NH₂PhHA'. [192°]. Plates.

PHENYL SULPHATE,

Hydrogen phenyl sulphate C, H, O.SO, OH. Phenyl-sulphuric acid. Occurs in urine of horses, men, and dogs (Baumann, B. 9, 55; 11, 1907; H. 2, 335), the quantity being greatly increased by administration of phenol. The K salt is formed by adding K.S.O. (125 pts.) to a solution of phenol (100 pts.) and KOH (60 pts.) in water (90 pts.) at 65°. The free acid quickly splits up, in aqueous or alcoholic solution, into phenol and H₂SO₄.—KA'. S. 14 at 15°. Plates (from alcohol). Not attacked by potash. The dry salt changes at 150°-160° into phenol psulphonic acid.—BaA', 3aq: needles.
DI-PHENYL SULPHAZIDE v.

Phenylhydrazide of Benzene sulphonic acid.

DI-PHENYL SULPHIDE C12H10S i.e. Ph2S.

Mol. w. 186. (293°). S.G. 1·119. Formation.—1. By the dry distillation of sodium benzene sulphonate (Stenhouse, Pr. 14, 351; A. 140, 288).—2. From S(C₀H₄.NH₂)₂ by elimination of NH₂ by the diazor reaction (Krafft, B. 7, 384, 1164).—3. By distilling Pb(SPh)₂ (Kekulé a. Szuch, C. R. 64, 752).—4. By the aution of Na on SO(C₆H₅)₂ (Colby a. McLoughlin, Am. 9, 67).—5. By the action of NaSPh on C_eH_b, N₂Cl in molecular proportions (Ziegler, B. 23, 2471).-6. By heating benzene with S and AlCl. at 80° (Friedel a. Crafts, A. Ch. [6] 14,

Preparation.—Ammonium sulphide is slowly added to an acid solution of diazobenzene, cooled with ice, the oil which separates is cohobated for 2 or 3 hours to decompose the disulphide, and finally distilled; the yield is good

(Graebe a. Mann, B. 15, 1683).

Properties .- Oil, with slightly alliaceous odour, v. sol. hot alcohol, miscible with ether. Its alcoholic solution is not ppd. by AgNO, or HgCl2. Yields di-phenyl-sulphone on oxidation.

Di-phenyl disulphide Ph₂S₂. Mol. w. 218.

[61^{\text{\text{0}}}. (310°).

Formation.—1. By oxidising phenyl meraptan with dilute HNO₃ (Vogt, A. 119, 142; Otto, A. 143, 213), with chromic acid mixture (Kekulé, Z. 1867, 194), and by exposure of its ammoniacal solution to air.—2. From NaSPh and CyCl (Clemm, J. pr. [2] 1, 147).—8. Together with me cury by distilling Hg(SPh), (Dreher a. Otto, A. 154, 178).—4. From NaSPh and I (Hübner a. Alsberg, A. 156, 330).-5. In small quantity by adding zinc-dust to a mixture of Scil and benzene (Schmidt, B. 11, 1173).—6. By reducing C₂H₂SO₂O with HI (Cleve, B. 21, 1100).—7. By heating C₂H₃SO₂H with HSPh (Otto, B. 9, 1589).—8. By passing SO₂ into a ling with alcohol into C₂₂H₂₂SO₂, sol. alcohol

solution of phenyl-hydrazine in benzene, and heating to boiling (Michaelis a. Ruhl, B. 23;

Properties.-Needles, with faint odour, insol.

water, v. sol. alcohol and ether.

Reactions. -1. Reduced by zinc and dilute H₂SO, to C₆H₅SH.—2. Oxidised by nitric acid to benzene sulphonic acid.—3. Split up by long boiling into Ph₂S and S (Graebe, A. 174, 189).— 4. Alcoholic potash forms KSPh and C₆H₅SO₂K. 5. Alcoholic K2S forms KSPh.-6. Bromine forms crystalline Ph.SBr.,.

Di-phenyl tetra-sulphide (C₆H₃)₂S₄. S.G. 145 1.297. Formed by the action of H₂S on a conc. alcoholic solution of benzene sulphinic acid (Otto, J. pr. [2] 37, 208). Formed also by the action of S_2Cl_2 on phenyl mercaptan. Thick yellow oil, m. sol. alcohol, v. sol. ether.

Di-phenyl hexasulphide $(\mathbf{C_6H_5})_2\mathbf{S_6}$. amorphous solid formed by heating benzene (20 g.) with S₂Cl₂ (10 g.) and iodine (1 g.) for 100 hours at 120° (Onufrovitch, B. 23, 3368).

References .- AMIDO-, AMIDO-IMIDO-, BROMO-, CHLORO-, CHLORO-NITRO-, IDDO-, NITRO-, and OXY-DI-PHENYL SULPHIDES. V. also DI-PHENYL-DI-PHENYL SULPHIDE

DI-PHENYL SULPHIDE O-CARBOXYLIC ACID C.H.S.C.H.CO.H. Formed by the action of alkalis on the product of the action of NaSPh on diazotised o-amido-benzoic acid (Ziegler, B.

23, 2471). Plates (from benzene), insol. water. PHENYL SULPHINIC ACID is BENZENE SULPHINIC ACID.

Diphenyl sulphinic acid C.H.Ph.SO.H. Got by reducing C12H9.SO2Cl in ether with sodium-

amalgam (Gabriel a. Deutsch, B. 13, 388). Crystalline powder, decomposing at about 70°. PHENYL SULPHITE (Schall, B. 25, 1490, 1875). The salt NaSO2.OPh is formed from NaOPh and SO2, and exhibits the following re-

actions :- 1. Alkyl iodides convert it into phenyl ethers of the sulphonic acids R.SO₂OH.— 2. Benzyl ether forms phenyl benzyl oxide, SO₂, and NaCl at 180° (yield 16 p.c.).—3. COCl, forms CO(OPh)2, SO2, and NaCl at 180°.-4. Iodoform (1 mol.) heated with NaSO, OPh (3 mols.) at 180° forms MeI, CH, SO, OPh, and rubbadin.

Rubbadin C₄₄H₃₂S₄O₈, mol. w. 830 (by Raoult's method, calc. 806). Rub. crystallises from dilute alcohol in minute brownish-red plates, S. (alcohol) 43 at 20°, and forms a claret-coloured solution in alkalis. Its solution in NH, Aq dyes silk and wool pale-claret colour, and cotton

yellowish-violet.

Reactions .- 1. Reduced by zinc-dust and NaOHAq to $C_{44}H_{34}S_3O_5$.—2. Conc. HNO₃, free from nitrous fumes, at 0° forms $C_{44}H_{24}(NO_2)_3S_3O_5$, a red crystalline powder, m. sol. hot alcohol, which may be reduced to $C_{44}H_{23}(NO_2)_4(NH_2)_2S_3O_6$, a glittering black crystalline powder (from etherphenol) which yields a black crystalline di-acetyl derivative. - 3. HClAq at 200° forms phenol (2 mols.), H_2S (2 mols.), and $C_{32}H_{20}S_2O_8$, a reddish-brown crystalline powder which yields the derivatives $C_{s_1}H_{1s}Ac_2S_2O_s$, $C_{s_2}H_{1s}Me_2S_2O_s$, and $C_{12}H_{1s}H_{1s}Ac_2S_2O_s$,—4. Fusion with NaOH forms SO(C_sH_sOH)₂ [96°] and salicylic acid.

Di-acetyl derivative $C_sH_{1s}Ac_2S_sO_s$.

and $C_{2i}H_{2i}S_2O_4$, insol. alcohol. Br in chloroform converts the di-acetyl derivative into

C44H24Br6Ac2S4O8

Di-methyl-rubbadin C44H30Me2S4O8. from rubbadin, KOHAq, and Mel. Dark reddish-

brown powder.

SULPHOCYANIDE PHENYL C.H.SCN. (231° cor.). S.G. 17.5 1.155. Formed from Pb(SPh)2 and CyCl. Prepared by adding cuprous sulphocyanide to a solution of diazobenzene sulphate and potassium sulphocyanide (Billeter, B. 7, 1753; Gattermann a. Haussknecht, B. 23, 738). Liquid. Yields phenyl mercaptan when heated with alcoholic potash.

Polymeride (Ph.SCy), [97°]. Got from PhSNa and cyanuric chloride (Klason, J. pr. [2]

33, 120). Prisms (from HOAc).

Reference.—NITRO-PHENYL SULPHOCYANIDE.
PHENYL SULPHOCYANO-ETHYL KETONE C₁₀H₀NSO i.e. C₆H₄.CO.C₂H₄.SCy. Got from C.H. CO.C.H. Br and KSCy in alcohol (Pampel a. Schmidt, B. 19, 2897). Liquid.

DI-PHENYL SULPHONE (C.H.), SO... Sulphobenzide. Mol. w. 218. [128°]. (376°) at

722 mm.

Formation. - 1. From benzene and SO. (Mitscherlich, P. 31, 628; Freund, A. 120, 76; Otto, A. 136, 160), fuming H₂SO₄ (Berthelot, B. 9, 349), or HSO₃Cl (Knapp, Z. [2] 5, 41).—2. By oxidation of Ph.S (Stenhouse, A. 140, 290; Kekulé a. Szuch, Bl. [2] 8, 204).—3. By distilling benzene sulphonic acid (F.). -4. From benzene, C₈H₈SO₂Cl, and AlCl₃; the yield being 80 p.c. of the theoretical amount (Beckurts a. Otto, B. 11, 2066).—5. By heating benzene (200 c.c.) with conc. H₂SO₄ (300 c.c.), the yield being 6 p.c. of the benzene employed (Istrati, Bl. [3] 1, 492).— 6. By heating C.H. SO.Cl with HgPh. (Otto, B. 18, 248).-7. By oxidation of di-phenyl sulphoxide with KMnO, and HOAc (Colby a. McLough. lin, Am. 9, 67).

Properties. - Monoclinic prisms (from benzene) or plates (from alcohol), v. sol. ether. PCl, at 160° forms C.H. SO.Cl and chloro-benzene. Converted by potash-fusion into phenol, diphenyl, Ph₂S, PhSH, and other bodies (Otto, B. 19, 2425). Hot fuming H2SO4 forms benzene sul-

phonic acid.

References. - AMIDO-, Bromo-, Chloro-, NITRO-, OXY-AMIDO-, and OXY- DI-PHENYL SUL-PHONES.

PHENYL SULPHONE ACETIC ACID v.

PHENYL-SULPHONO-ACETIC ACID.

DI-PHENYL SULPHONE o-CARBOXYLIC ACID C13H10SO4 i.e. C6H5.SO2.C6H4.CO2H. [152°]. Formed by oxidising PhS.C.H..CO2H with HNO. (Graebe a. Schultess, A. 263, 7). Needles (containing aq). Melts at 99° when hydrated.

Di-phenyl sulphone p-carboxylic acid. [above 300°]. Got by oxidation of phenyl tolyl sulphone with KMnO₄ (Michael a. Adair, B. 11, 119). Prisms (from alcohol), sl. sol. hot water. - AgA'.

Di-phenyl sulphone di-carboxylic acid SO₂(C₆H₄,CO₂H)₂. [above 300°]. Got by oxidising di-p-tolyl sulphone (M. a. A.). Small prisms, sol. nitrobenzene.

DI-PHENYL SULPHONE m-SULPHONIC ACID C.H.SO.C.H.SO.H. Got by heating diphenyl sulphone (1 mol.) with HO.SO₂Cl (1 mol.) at 150° (Otto, B. 11, 2075; 19, 2417). Fibrous crystalline mass, v. sol. water. — KA'aq.— NaA' 3aq.—CaA', 7aq.—BaA', 4\frac{1}{2}aq.—PbA', 3\frac{1}{2}aq. -CuA', 7 gaq. Chloride.

[99°]. White needles. Amide Ph.SO₂.C₆H₄.SO₂NH₂. [154°].
Anilide. [181°]. Hard nodules. Ethers.-EtA'. [89°].-PhA'. [106°].

Di-phenyl sulphone di-sulphonic acid SO₂(C₆H₄.SO₃H)₂. Formed from di-phenyl sulphone (1 mol.) and HO.SO₂Cl (2 mols.) at 155° (Otto, B. 19, 3124). Deliquescent mass.- $K_2A''aq.-Na_2A''3aq.-CaA''6\frac{1}{2}aq.-BaA''5aq.$ -PbA" Baq. - CuA" Baq.

Chloride C₁₂H₈S₂O₆Cl₂. [176°]. Plates.

Amide. [242°]. Nearly insol. ether.

Anilide. [212°]. Lustrous plates.

Ethers Et₂A". [82°].—Ph₂A". [198°].

DIPHENYL SULPHONIC ACID C₁₂H₁₆SO,

i.e. C. H. Ph. SO. H. Formed, together with the disulphonic acid, by heating diphenyl with H SO. (Engelhardt a. Latschinoff, B. 4, 561; 6, 193). -KA'aq. Crystals (from 40 p.c. alcohol).-BaA'2. - CuA'2 6aq.

Ethyl ether EtA'. [74°]. Needles. Chloride C12H9.SO2Ci. [115°].

 $Amide C_{12}H_9.SO_2NH_2$. [227°-230°]. Slender needles (Gabriel a. Deutsch, B. 13, 386)

Diphenyl di-o-sulphonic acid C12II1082O8 i.e. SO₃H.C₆H₄.C₆H₄.SO₈H. Formed by elimination of NH, from benzidine disulphonic acid (Limpricht, A. 261, 327). Viscid mass. Yields dioxy-diphenyl [99°] by potash-fusion. BaA"6 aq.

-PbA" 5aq: needles, v. e. sol. water.

Chloride C₁₂H₈(SO₂Ol)₂. [138°]. Prisms.

Amide C₁₂H₈(SO₂NH₂), 2aq. Prisms.

Anilide C₁₂H₈(SO₂NHPh)₂. [157°].

Diphenyl di-p-sulphonic soid C₁₂H₈(SO₂H)₂.

[72.5°]. Formed by sulphonating diphenyl (Fittig, A. 132, 209; Engehardt, Z. 1871, 260). Deliquescent prisms. - K,A" 2 aq. - CaA". -BaA"

Chloride. [203°] (G. a. D.). Prisms. Amide. [above 300°]. Needles. References .- NITRO-, OXY-AMIDO-, and OXY-DIPHENYL BULPHONIC ACID.

PHENYL-SULPHONO-ACETIC ACID C₆H₈SO₄ i.e. C₆H₈.SO₂.CH₂.CO₂H. [112°].

Formation .- 1. From sodium benzene sulphinate and chloro-acetic acid (Gabriel, B. 14, 834).—2. By saponifying its ether, which is got by boiling sodium benzene sulphinate with chloro-aceto-acetic ether (Otto a. Rössing, B. 23, 755).—3. By oxidising phenyl-thio-glycollic acid (Claesson; Blomstrand, B. 4, 712; 8, 120; Otto, B. 19, 3138).—4. By warming phenyl oxyethyl sulphone (1 vol.) with H₂SO₄ (1 vol.), diluted with water (2 vols.), and extracting with ether (Otto, J. pr. [2] 80, 840).

Properties. Monoclinic crystals, m. sol. water, but separated by addition of NaCl. Sol.

alcohol and ether.

Reactions .- 1. Reduced by sodium amalgam to benzene sulphinic and acetic acids. - 2. Gives off CO₂ above 160°, forming phenyl methyl sulphone.—3. Chlorine, passed into its yarm aqueous solution, forms CHCl₂SO₂C₄H₅ [59°].— 4. PCl₅ gives the chloride [58°], and at 110° forms C₆H₅SO₂CCl₂COCl, which on treatment with water gives C.H. SO2. CHCl2 (Otto, J. pr. [2] 40, 540).
Salts.—BaA', 2aq.—CuA', 2aq.—CaA', 2½aq.
—PbA', 2aq.—AgA'. Sparingly soluble needles.

Methyl ether MeA'. Oil (Otto, J. pr. [2]

Ethyl ether Eth'. [42°] (O.). Readily saponified by NaOHAq, while KOHAq gives phenyl methyl sulphone. Bromine at 90°, followed by NaOHAq, forms C.H.SO2.CHBr2 [76°]. Alcoholic potash forms Ph.SO.Me (Michael a. Milner, Am. 7, 65). NaOEt forms the compound C,H,SO,CHNa.CO,Et, which yields Ph,S, Ph,S, PhSEt, and Ph.SO₂Me on distillation (Otto a. Rössing, B. 22, 1453; 23, 1647).

Chloride Ph.SO2.CH2.COCl.

Amide Ph.SO₂.CH₂.CONH₂. [153°]. Needles (from water). Yields Hg(C₈H₈NSO₃)₂ [215°] on boiling with HgO.

PHENYL-SULPHONO-ACETONE v. PHENYL

METHYLENE METHYL SULPHONE-KETONE.

Di-phenyl-di-sulphono-acetone v. DI-PHENYL DI-METHYLENE DISULPHONE-KETONE.

PHENYL-SULPHONO-ANGELIC ETHER C₁₈H₁₆SO₄ i.e. C₆H₅.SO₂.CH(C₃H₅)CO₂Et. [64·5°] Got from PhSO₂.CHNa.CO₂Et and allyl iodide (Michael, Am. 7, 67). Prisms (from alcohol).

PHENYL-SULPHONO n-BUTYRIC ACID C.H., SO. CHEt. CO.H. [124°]. Formed from benzene sulphinic acid, a-bromo-butyric acid, and Na₂CO₃Aq (Otto, B. 21, 996). Needles, m. sol. water. Yields phenyl propyl sulphone on heating.

Di-phenyl-di-sulphono-butyric ether (C₆H₅.SO₂)₂CMe.CH₂.CO₂Et. [97°]. Got by oxidation of (PhS)₂CMe.CH₂.CO₂Et (Autenrieth, A. Small crystals, insol. water. Con-259, 367). verted by KOHAq into benzene sulphinic acid and β -phenyl-isocrotonic acid.

PHENYL-SULPHONO-CROTONIC ACID

C_cH_a·SO₂·CMe:CH_cCO₂H. [158°]. S. 15 38; 120 25. Formed by heating the Na salt of β-chloro-crotonic acid [94·5°] with sodium benzene sulphinate in aqueous solution at 170° (Autenrieth, A. 259, 343). Plates (from water), m. sol. ether. At 210° it slowly changes to the following isomeride [127°]. — KA' 1½aq. — BaA', aq. — MgA', 7aq. — ZnA', 6aq. — CuA', aq. — AgA'. [200°]. Matted needles (from water).

Phenyl-sulphono-isocrotonic acid

C.H., SO. CMe: CH.CO.H. [127]. S. 26 at 15°; 5 at 100°. Formed by heating \$\beta\$-chloro-isocrotonic acid with sodium benzene sulphinate in aqueous solution at 145°. Got also by saponifying di-phenyl-di-sulphono-isobutyric ether (Autenrieth, A. 259, 335). Feathery needles. Yields benzene sulphinic acid [81°] on boiling with potash.— KA' 3aq.—BaA', 21aq.— MgA', 6aq.-ZnA', 6aq. [120°] (hydrated).—AgA'. [c. 243°].

Ethyl ether EtA'. Oil.

DI-PHENYL-DI-SULPHONO-DI-ETHYL **OXIDE** C₁₆H₁₈S₂O₅ *i.e.* O(C₂H₁.SO₂Ph)₂. [70°]. Formed from C₆H₅.SO₂.C₂H₄Cl and dry Ag₂O (Otto, *J. pr.* [2] 30, 202). Needles, sol. benzene. A polymeride [88°] is got by heating the compound (C,H,SO,2)2C,H, with KOHAq.

DI-PHENYL-DI-SULPHONO-DI-ETHYL SULPHIDE S(C₂H₄,SO₂Ph)₂. [124']. Got from C₂H₃,SO₂C₂H₄Cl and alcoholic KSH (Otto, *J. pr.* [2] 30, 348). Needles (from dilute alcohol).

DI - PHENYL - DI - SULPHONO - HEXOIC ETHER (C.H., SO.) CMe. CHEt. CO. Et. [111°]. Got by oxidising (PhS) CMe. CHEt. CO. Et. (Autenrieth, A. 259, 872). Crystals, v. sol. alcohol.

DI-PHENYL-DI-SULPHONO-DI-PHENYL OXIDE (C.H.SO.C.H.).0. [70°]. (above 200°). Got by heating C.H.(SO.Ph). with alcoholic potash at 165° (Otto a. Rössing, B. 20, 187). Needles, v. e. sol. alcohol, insol. water.

PHENYL-a-SULPHONO-PROPIONIC ACID C₆H₅.SO₂.CHMe.CO₂H. [116°]. Formed by heating a-bromo-propionic ether with sodium benzene sulphinate and ether. Minute needles. -NaA'.—BaA', 2aq.—EtA'. [c. 17°].

Phenyl-\$-sulphono-propionic acid

 $C_6H_5.SO_2.CH_2.CH_2.CO_2H.$ [124°]. Got from β -bromo-propionic acid, benzene sulphinic acid, and Na₂CO, Aq (Otto, B. 21, 89). Plates, sl. sol. cold water.

DI-PHENYL SULPHOXIDE $(C_sH_s)_sSO.$ [71°]. Formed by the action of SO, or SOCl. on benzene in presence of AlCl, (Colby a. McLoughlin, Am. 9, 67; B. 20, 195). Triclinia crystals, v. sol. alcohol. Oxidised by KMnO, to Ph₂SO₂. Reduced by sodium to Ph₂S. NaNO₃ and H₂SO₄ give SO(C₆H₄·NO₂)₂ [116°] and SO₂(C₆H₄·NO₂)₂ [163°].

Di-phenyl-di-sulphoxide v. Phenyl ether of

BENZENE THIOBULPHONIC ACID.

PHENYL-SULPHURIC ACID v. PHENYL BULPHATE and PHENOL, Reaction 19.

PHENYL SULPHUROUS ACID v. BENZENE SULPHONIC ACID.

PHENYL SULPHYDRATE v. PHENYL MER-

PHENYL-SULPHYDRO-ACETIC ACID v. PHENYL-THIOGLYCOLLIC ACID.

PHENYL - SULPHYDRO - ACETOPHENONE C₆H₈.CO.CH₂.SPh. [53°]. Formed from ω-bromo-acetophenone and NaSPh (Delisle, B. 22, 306). Crystals, v. sol. ether and acetone.

DI - PHENYL - DI - SULPHYDRO - BUTYRIC ETHER C₁₈H₂₀S₂O₂ i.e. (PhS)₂CMe.CH₂.CO₂Et. [58°]. Got by passing HCl into a mixture of acetoacetic ether and phenyl mercaptan (Escales a. Baumann, B. 19, 1790). Pearly plates (from alcohol), v. sol. ether.

PHÉNYL-SULPHYDRO-CROTONIC CH₃.CH:C(SPh).CO₂H. [86°]. Formed by the action of Na and phenyl mercaptan on the Na salt of a-chloro-crotonic acid [97°] (Autenrieth, A. 254, 246). Needles or plates. Its K salt is hygroscopic, S. (alcohol) 20.8.

Isomeride CHMe:C(SPh).CO2H. [80°]. Got in like manner from the stereo-isomeric liquid chloro-crotonic acid. White plates. Its K salt

is hygroscopic, S. (alcohol) 10.4.

Phenyl-8-sulphydro-crotonic acid CH₂.C(SPh):CH.CO₂H or CH₂:C(SPh).CH₂.CO₂H. [177°]. S. (alcohol) 2 in the cold. Formed, together with small quantities of an isomeride [145°], by warming di-phenyl-di-sulphydro-butyric ether with alcoholic soda (Escales a. Baumann, B. 19, 1791). Plates (from alcohol), insol. water.—BaA', 2aq.—AgA': amorphous pp. Isomeride CH, C(SPh):CH.CO, H. [158°].

Formed from phenyl mercaptan and the Na salt of chloro-crotonic acid [95°] (Autenrieth, A. 254, 230). Both acids form a cherry-red solution in H,80.

DI-PHENYL SULPHYDRO-ETHY DISULPHONE. Phenyl derivative SULPHYDRO-ETHYLIDENE (C₆H₄.SO₂)₂CMe.SPh. [194°]. Got by heating (C₆H₄.SO₂)₂CH.SPh with alcoholic soda and

MeI at 100° (Laves, B. 23, 1416). Crystals (from

chloroform)

DI - PHENYL - DI - SULPHYDRO - HEXOIC ETHER CH. C(SPh) CHEt. CO Et. [71°]. Got from ethyl-acetoacetic ether, phenyl mercaptan, and HCl (Autenrieth, A. 259, 871). Shining crystals, v. sol. ether and alcohol.

DI-PHENYL SULPHYDRO-METHENYL DI-

SULPHONE. Phenyl derivative

(C_sH₅.SO₂)₂CH.SPh. [175°]. Got by the action of KMnO, and dilute H.SO, on CH(SPh), dissolved in benzene (Laves, B. 23, 1414). Silky needles, m. sol. chloroform, nearly insol. water and alcohol, sol. alcoholic NaOH. May be oxi-

dised to CH(SO₂Ph)_s [215°]. DI-PHENYL-DI-SULPHYDRO-PHENYL-ACETIC ACID C,H,C(SPh)2.CO,H. Formed from phenyl-glyoxylic acid, phenylmercaptan, and HCl (Escales a. Baumann, B. 19, 1789). Insol. water, v. sol. ether. Crystallises from benzene with $\frac{1}{2}C_{d}H_{e}$.—KA' $1\frac{1}{2}$ aq.

PHENYL - SULPHYDRO - PHENYL - THIO-

NPh.N CS—S > C.NH.NHPh. BIAZYL-HYDRAZINE [142°]. Got by reducing the corresponding azocompound with alcoholic ammonium sulphide (Freund a. Kuh, B. 23, 2830). Plates, sl. sol. alcohol.

DI-PHENYL - DI - SULPHYDRO - PROPIONIC **ACID** CH₃.C(SPh)₂.CO₂H. [117°]. Got from pyruvic acid, phenyl mercaptan, and HCl (Escales a. Baumann, B. 19, 1787). Needles (from ligroin).—NaA'.—BaA', 2aq: v. sol. hot Aq.

Amide. [93°]. Needles (from alcohol)
DI-PHENYL SULPHYDRO-PROPYLE SULPHYDRO-PROPYLENE DISULPHONE. Phenyl derivative C₆H₅.SO₂.CH₂.CMe(SPh).SO₂.C₈H₅. Formed by oxidising C_eH₅.SO₂.CH₂.CMe(SPh)₂

234). Crystalline granules, sl. sol. cold alcchol. DI - PHENYL - DI - SULPHYDRO - VALERIC ACID CH₂.C(SPh)₂.CH₂.CH₂.CO₂H. [69°]. Got from phenyl mercaptan, β-acetyl-propionic acid and HCl (Escales a. Baumann, B. 19, 1795).

with KMnO, in the cold (Otto a. Rössing, B. 24,

Prisms (from chloroform-ether).—BaA'2. PHENYL-TETRIC ACID $C_{11}H_{10}O_{8}$ CH₂Ph.CH.CO CO.CH₂>O. Formed by heating bromo-

benzylacetoacetic ether (Moscheles, B. 21, 2609). Not affected by HClAq at 170°. Yields a benzoyl derivative C11H2BzO2 [110°] crystallising in needles.

PHENYL-THIAZOLE C.H., NS i.e.

N CH .S [52°]. (273° cor.). Got by the diazo- reaction from the amido-phenyl-thiazole that is produced by the action of thio-urea on bromo-acetophenone (Popp, A. 250, 279; cf. Arapides, A. 249, 25). Sl. sol. cold water.—
"B'HCl. [80°]. Needles.—B'_2H_2PtCl_2aq. [196°].
—*B'HHgCl_. [163°].—Picrate. [165°].— *B'HAuCl. [175°]. Needles (from alcohol).

N≪CH :CH μ-Phenyl-thiazole (269°). Formed by heating thio-benzamide with dichloro-di-ethyl oxide at 100°. Oil.—B'HCl 2aq. [62°].—B'₂H₂PtCl₂2sq. [175°] (when anhydrous). —Picrate. [125°]. Yellow needles.

Dihydride N CPh.S CH, CH,

with ethylene bromide (Gabriel a. Heymann, B. 23, 158; 24, 784). Formed also by the action of PCl₅ on S₂(O₂H₄.NHB₂)₂ (Gabriel a. Coblentz, B. 24, 1124). Oil. Yields a crystalline dibromide. — B'₂8HCl. — B'C₅H₄N₂O₇. [172°]. Yellow needles. -B' H2PtCl.

v-Phenyl-thiazole. Tetra-hydride

 $NPh {<}_{\mathrm{CH}_2,\mathrm{CH}_2}^{\mathrm{CH}_2,\mathrm{S}}.$ Got by reducing ethylene phenyl-di-thio-carbamate [128°] with tin and HCl (Förster, B. 21, 1871). Oil. - B'₂H₂PtCl₆.

N CPh.S CPh:CH Di-phenyl-thiazole (above 360°). Got from thiobenzamide and bromo-acetophenone in alcohol (Hubacher, A. 259, 237). Plates, v. sol. alcohol and ether.

N≪CPh.S CPh:CPh Tri-phenyl-thiazole Got from thiobenzamide and bromo-deoxybenzoïn (Hubacher, A. 259, 245). Prisms (from ether), m. sol. alcohol, nearly insol. HClAq.

Reference.—Oxy-Phenyl-Thiazole. PHENYL-DITHIENYL C.H.S.C.H.PhS. [209°]. A product of the action of S on toluene at a red heat (Renard, C. R. 111, 48). Plates, v. sl. sol. alcohol and other. Gives a blue colour with isatin and H₂SO₄, and a green colour with phenanthraquinone and H_2SO_4 . Gives rise to $C_{14}H_1Br_8S_2$ [320°] and $C_{14}H_8(NO_2)_2S_2$ [273°].

PHENYL THIENYL KETONE v. THIENYL

PHENYL KETONE

(α)-PHENYL-(β)-THIO-ALLOPHANIC ACID Ethyl ether. C₁₀H₁₂N₂SO₂ i.e. NH₂.CS.NPh.CO₂Et. [127°]. Formed from NHAc.CS.NHPh and ClCO2Et (Seidel, J. pr. [2] 32, 275). Monoclinic tablets, converted by alcoholic NH, at 100° into phenyl-thio-urea.

(β)-Phenyl-thio-allophanic acid. ether NHPh.CS.NH.CO2Et. Got from phenylthio-urea and ClCO₂Et (Seidel, J. pr. [2] 32, 270). Oil. AcCl at 60° forms NHAc.CS.NHPh [170°].

Phenyl di-thio-allophanic acid. Isoamyl NHPh.CS.NH.CO.SC,H11. [102°]. ether Formed from phenyl-thio-urea and Cl.CO.SC, H11 (Schöne, J. pr. [2] 32, 256). Needles. unstable. Forms an acetyl derivative [240°].

Di-phenyl-thio-allophanic acid. Ethyl [95°]. ether NHPh.CS.NPh.CO2Et. Formed from di-phenyl-thio-urea and ClCO,Et (Seidel, J. pr. [2] 32, 262). Prisms (from alcohol). Alcoholic AgNO₃ ppts. C₁₆H₁₆N₂SO₂AgNO₃.

Reactions.—1. Caustic potash regenerates CS(NHPh)₂.—2. Aqueous ammonia at 100° forms NHPh.CS.NH2 and NHPh.CO.Et. - 3. Alcoholic aniline at 100° forms CS(NHPh), and NHPh.CO₂Et.—4. HgO in presence of alcoholic ammonia forms (NHPh.CO.NPh.CO,Et)2HgO [129°], a crystalline body that is reconverted by H.S into di-phenyl-thio-allophanic ether .- 5. By heating in a current of HCl it is converted into CS(NHPh)2, aniline, and phenyl-thiocarbimide.

Di-phenyl-di-thio-allophanic acid. Isoamyl ether NPhH.CS.NPh.CO.SC,H,,. Got from di-phenyl-thiures and Cl.CO.SC, H11 (Schöne, J. pr. [2] 32, 258). Needles (from alcohol). It is insoluble in cold water, but decomposed by hot water. Its alcoholic solution is (276). Got by heating thio-benzamide ppd. by AgNO,, HgCl, or PtCl,. HgO and NH, convert it (in alcoholic solution) into di-phenyl-guanidine, [147°]. Formed also by the action of alcoholic potash on phenyl-thiocarbimide (R. Schiff, B. 9, 1316).

TRI-PHENYL-THIO-AMMELINE

C₂₁H₁₇N₅Si.e. C(SH) N C(NPh) NH. [238°]. Formed from phenyl-cyanamide and phenyl-thiocarbimide, and got also by heating tri-phenyl-biguanide with CS₂ at 100°, and by heating diphenyl-guanidine with phenyl-thiocarbimide, alcohol, and mercury sulphocyanide (Rathke, B. 20, 1065; 23, 1673). Plates (from chloroform). Converted by alcoholic potash into tri-phenyl-ammeline. Alcohol and EtBr at 100° form C₂₁H₁₅EtN₅SHBr, crystallising from alcohol in plates.—B'HCl.—AgC, HPh₃N₃S: yellow pp. DI-PHENYL-THIO-BENZAMIDE

DI-PHENYL-THIO-BENZAMIDE Ph.CS.NPh₂. Formed by heating di-phenylbenzamidine with H_2S or CS_2 at 130° (Bernthsen, A. 192, 38). Yellow triclinic crystals (from benzene); a:b:c=:927:1: '766, a=93° 29', $\beta=$ 100° 55', $\gamma=$ 84° 14'. Insol. cold water, sl. sol. cold alcohol.

PHENYL-DI-THIO-BIURET C₈H₈N₃S₂ i.c. NHPh.CS.NH.CS.NH₂. [174°]. Formed by heating aniline with perthiocyanic acid (Glutz, A. 154, 44). Got also by boiling carbimidophenyl-thio-urea with ammonium sulphide (Wunderlich, B. 19, 452). Prepared by adding phenyl-thiocarbimide (36 g.) to an alcoholic solution of cyanamide (11 g.) and Na (6 g.), and decomposing the pp. by dissolving in a solution of NH₄Cl (14·5 g.) containing NH₂Aq (50 c.c.), passing in H₂S, and heating to boiling (Hecht, B. 25, 756). Pearly plates, sl. sol. hot water.—B'HCl.—B'HNO₃.

Ethyl derivative C.H.EtN.S. [109°]. Got by adding alcohol, NH.Aq, and EtI to the base (Tursini, B. 17, 584). Trimetric tables. PHENYL-THIO-CARRAMIC

PHENYL-THIO-CARBAMIC ACID
NHPh.CO.SH. Methyl ether NHPh.CO.SMe.
[84°]. Formed by heating di-phenyl-methyl-\$\psi\$thio-urea NHPh.C(NPh).SMe with dilute \$H_2\$O_4
(Will, B. 15, 339). Plates, sol. alcohol, ether,
and benzene. Decomposed by KOH into aniline,
MeSH, and CO_2. Boiling alcoholic NH₃ forms
MeSH and phenyl-urea. Aniline gives methyl
mercaptan and CO(NHPh)₃.

Ethyl ether NHPh.CO.SEt. [73°]. Needles. Got by heating NHPh.C(NPh).SEt with dilute H₂SO₄ and by the action of aniline on

Cl.CO.SEt (Will).

Isoamyl ether NHPh.CO.SO.H... [67°]. Got from aniline and Cl.CO.SC.H... (Schöne, J. pr. [2] 82, 249). Needles (from ether).

Ethylene ether CO SPh.CH₂. [79°]. Got by heating di-phenyl-ethylene-ψ-thio-urea with dilute HClAq at 200° (W.). Formed also by the action of chloro-acetic acid on the ethylene-ther of phenyl-di-thio-carbamic acid (Evers, B. 21, 976). Needles, sol. alcohol and ether, insol. acids and alkalis.

Phenyl ether NHPh.CO.SPh. [125°]. Formed by heating phenyl mercaptan with phenyl cyahate (Snape, C. J. 47, 778; B. 18, 2432). Needles, v. sol. alcohol and ether.

Phenyl-\psi-thio-carbamic acid NHPh.CS.OH. Ethyl ether C.H., NSO i.e. NHPh.CS.OEt or NPh:C(SH).OEt. [72°]. Formed by heating phenyl-thiocarbimide with alcohol (Hofmann, B. 2, 120; 8, 772; Bamberger, B. 15, 2164).

Formed also by the action of alcoholic potash on phenyl-thiocarbimide (R. Schiff, B. 9, 1316). Triclinic prisms, sol. alkalis and reppd. by CO_Resolved by distillation into alcohol and phenyl-thiocarbimide. Oxidised by alkaline K₂FeCy₄ to C₁₈H₂₆N₂O₂S₂ or S₂(C(NPh).OEt)₂ [102°], which is also got by the action of I on C₆H₁₆NOSAg (Liebermann a. Natanson, B. 18, 1575; Jacobsen, B. 19, 1076).—(C₆H₁₆NOS)₂Pb 2aq: small needles.—(C₈H₁₆NOSAg.—C₆H₁₆NOSHgClHCl: needles.—(C₆H₁₆NOS)₂HgAgNO₃. Needles.—C₉H₁₆NOSMe. (260°-265°). Oil.—C₆H₁₆NOSEt. [30°]. (275°). Prisms; decomposed by dilute H₂SO₄ at 200° into aniline and CO(OEt)(SEt) (Liebermann, B. 13, 682; A. 207, 148).—C₉H₁₆NOSC,H₁₆. [75°] (Mylius, B. 5, 977). Phenyl ether NHPh.CS.OPh. Got by

Phenyl ether NHPh.CS.OPh. Got by heating phenyl-thiocarbimide with phenol at 145° (Dixon, C. J. 57, 268). Sulphur-yellow

octahedra; decomposed by heat.

Phenyl di-thio-carbamic acid NHPh.CS.SH. Splits up into CS₂ and aniline when set free from its salts.

Salts.—The salt NHPh.CS.SK is formed by boiling potassium xanthate EtO.CS.SK (1 mol.) with an alcoholic solution of aniline (1 mol.) (Rathke, B.11, 958). Prepared from CS₂, aniline, ether, and alcoholic potash (Losanitsch, B. 24, 3024). Yellow monoclinic needles, v. e. sol. water and alcohol, insol. ether. Yields CS(NHPh)₂ when heated. Boiling water forms phenyl-thiocarbimide and 'CS(NHPh)₂—NHPh.CS.SNH₄. Yellow prisms. Converted by I into (NHPh.CS)₂S [138°]. BaA'₂: yellowish plates.—NiA₂.

Yellow prisms. Converted by I into (NHPh.CS.)₂S [138°]. BaA'₂: yellowish plates.—NiA'₂:

Methyl ether NHPh.CS.SMe. [93·5°].

Formed from the NH₄ salt and MeI (L.) and also by heating NHPh.C(NPh).SMe with CS₂ at 160° (Will, B. 15, 342). Decomposed by alkalis into

MeSH and aniline.

Ethylether NHPh.CS.SEt. [60°]. Formed from phenyl-thiocarbimide and mercaptan at 150° (Hofman, B. 2, 120; Will, B. 15, 1305), and also by the action of EtI on the ammonium salt (L.). Tables, v. sol. alcohol, sol. alkalis.

Isoamyl ether C, H, A'. [171°].

 $\begin{array}{cccc} Ethylene & derivative & \mathrm{CS} \leqslant \mathrm{NPh.CH_2} \\ \mathrm{[134^\circ].} & \text{Formed, together with phenyl-thiocarbimide, by heating C(NPh)} \leqslant \mathrm{NPh.CH_2} \\ \mathrm{with CS_2 at} \\ \mathrm{200^\circ (W.).} & \mathrm{Needles.} & \mathrm{Converted by MeI into C_3H_pNS_MeI [149^\circ].} \end{array}$

References .- Bromo-, Chloro-, and Nitro-

PHENYL-THIO-CARBAMIC ACID.

PHENYL-THIOCARBAMINE-CYAMIDE v. Carbimido-phenyl-thio-urea.

PHENYL - THIOCARBAMINE - ETHYL - CYAMIDE v. ETHYL-CARBIMIDO-PHENYL-THIO-

PHENYL-THIO-CARBAZIC ACID

NHPh.NH.CO.SH. The phenyl hydrazine salt NHPh.NH.CO.SN₂H₂Ph [84°] is formed from phenyl-hydrazine and COS (Heller, A. 263, 269).

Phenyl di-thio-carbasic acid NHPh.NH.CS.SH. Formed, as phenyl-hydrazine salt NHPh.NH.CS.SN₂H₂Ph [97°] by adding CS₂ to an ethereal solution of phenyl-hydrazine (Fischer, A. 190, 114). Plates, v. sol. alcohol and ether. Readily decomposed by heat into CS(N₂H₂Ph)₂, CS₂, H₂S, and NH₂.

PHENYL-THIO-SEMICARBAZIDE C.H.N.S i.e. NH2.CS.NH.NHPh. [201°] (F.; P.); [190°] (S. a. R.).

Formation.-1. By warming di-phenyl-thiocarbazide with alcoholic or aqueous alkalis (Fischer, A. 212, 324).—2. By heating phenylhydrazine sulphocyanide.—3. By heating phenylhydrazine with thio-urea or with phenyl-thiourea (Pellizzari, G. 16, 203; Skinner a. Ruhemann, C. J. 53, 550; B. 20, 3373).

Properties.—Needles, insol. ether, sl. sol. hot

water, v. sol. hot alcohol. Conc. HClAq at 120° forms phenyl-thiocarbizine C,H,N,S. COCL forms CO:N₂Ph.CS.NH₂ [c. 270°] crystallising from HOAc (Freund, A. 21, 2465).

Acetyl derivative NHPh.NH.CS.NHAc. [179°]. Formed by mixing solutions of acetylthiocarbimide and phenyl-hydrazine in dry benzene (Dixon, C. J. 55, 303). Small prisms. Benzoyl derivative C₁,H₁₈N₃SO. Silky needles. Does not melt at 220°.

CS(NH.NHPh)2. Di-phenyl-thiocarbazide [c. 150°]. Formed by heating the compound of phenyl-hydrazine and CS2 at 110°-140° (Fischer). Formed also from phenyl-hydrazine and CSCl₂ (Heller, A. 263, 278). Triangular prisms (from alcohol). Dissolves in hot aniline, forming a solution which is green in thin layers, but red in thick layers. Hot alcoholic potash also forms this dichroic substance, di-phenyl-thiocarbazone $C_{13}H_{12}N_4S$ or $CS(N_2Ph)NH.NHPh$, which rields $(C_{18}H_{12}N_4S)_2ZnO$, and may be reduced by zinc-dust and NaOHAq to phenyl-thio-semicarbazide and aniline. Di-phenyl-thio-carbazone may be oxidised by ppd. MnO₂ in presence of alcoholic potash to red needles of di-phenylthiocarbodiazone CS(N₂Ph)₂.

Di-phenyl-thio-semicarbazide NHPh.NH.CS.NHPh. [177°]. Formed from phenyl-hydrazine and phenyl-thiocarbimide in alcoholic solution (Fischer, A. 190, 122). Prisms from alcohol), sl. sol. ether. COCL, forms CO:N,Ph.CS.NHPh [188°], crystallising in

needles (Freund, B. 21, 2466).

Butyryl derivative C.H.O.NPh.NH.CS.NHPh. [156°]. Formed from sodium phenyl-hydrazine, phenyl thio-carbimide, and n-butyryl chloride (Michaelis a. Schmidt, A. 252, 309). V. sol. hot alcohol.

Bensoyl derivative

NPhBz.NH.CS.NHPh. [310°]. Obtained from NPhBz.NH, and phenyl-thiocarbimide (M. a. S.). Reference. - NITRO-DI-PHENYL-THIO-SEMICARB-AZIDE

DI - PHENYL - THIO - SEMICARBAZIDE CARBOXYLIC ACID C14H13N3O2S i.e.

NHPh.CS.NH.NHC, H4.CO, H. [204°]. Formed by warming m-hydrazido-benzoic acid with phenyl thiocarbimide and HOAc (Roder, A. 236, Colourless needles.

PHENYL-THIOCARBIMIDE C,H,NS i.e. C.H.N.CS. Mol. w. 185. (220°) at 750 mm. (Schiff, B. 19, 568). S.G. 2 1059 (Nasini a. Scala, G. 17, 66). S.V. 148.4 (Schiff; Losser.

A. 254, 73).

Formation.-1. By distilling phenyl-thiourea with P.O. (Hofmann, Pr. 9, 274, 487; C. J. 18, 309; B. 15, 980).—2. By heating di-phenylthio-urea with conc. HClAq at 165° (Merz a. Weith, Z. 1869, 589).—3. By the action of I on a benzene solution of tri-phenyl-guanidine (Hofmann, B. 2,453; Rudneff, J. R. 10, 184).-4. By heating phenyl cyanate or phenyl-carbamic ether with P₂S₅ at 160° (Michael a. Palmer, Am. 6, 258).—5. Formed, together with o- or p- tolylthiocarbimide, by heating phenyl-o- or p. tolyl-cyanamide (C_eH_sN:C:NC,H_r) with CS₂ at 180°-200° (Huhn, B. 19, 2409).

Properties. - Pungent oil, volatile with

steam.

Reactions. — 1. Alcohol at 100° forms NHPh.CS.OEt, while mercaptan gives rise to NHPh.CS.SEt. - 2. Alcoholic potash forms CS(NHPh)2 and CO(NHPh)2. - 3. Ammonia forms CS(NHPh)(NH2), aniline gives CS(NHPh)2, and other amines act in like manner.—4. PEt, acts violently, and forms $C_{13}H_{20}NPS$, while PMe, forms oily C10H14NPS, which yields crystalline B'HCl.-5. Chlorine passed into its solution in chloroform forms (PhNCS)₂Cl₂ [150°-160°] and PhNCCl₂ (212°). The chloride (PhNCS)₂Cl₂is converted by H₂S into CS(NHPh)₂, and by warm spirit into (PhNCS)2O, crystallising in yellow needles [118°] (Helmers, B. 20, 786). 6. Bromine in chloroform or HOAc forms red crystals of (PhNCSBr₂)₂, decomposed at 100° into pale-yellow plates (PhNCSBr)₂ [190°] (Helmers; cf. Proskauer a. Sell, B. 9, 1262).—7. SO, forms C,H₅NS₂O₃ [180°-183°], crystallising from benzene, and converted by water at 100° into amido-benzene p-sulphonic acid, CO_2 , and H_2S (Magatti, B. 11, 2267).—8. PCl₃ at 100° forms $C_eH_5NCCl_2$ and $C_eH_4{<}^N_S{>\!\!\!>}CCl$ (248°) (Hofmann, B. 12, 1126).—9. CISO₃H forms C₁₄H₁₀N₂S₃

[151°], insol. water, and SO, H.C, H, NOS aq, sol. water (Pawlewski, B. 22, 2200).-10. Reduced copper at 200° forms benzonitrile.—11. H₂S forms phenyl-thio-urea.—12. Boiling water forms CS(NHPh)₂, CO₂, and H₂S (Bamberger, B. 14, 2642). — 13. HOAc forms, on heating, CO(NHPh)2, Ac2O, H2S, and CO2; the di-phenylurea being further split up into aniline and CO₂, with formation of Ac₂O (Cain a. Cohen, C. J. 59, 327; cf. Gumpert, J. pr. [2] 32, 294).—14. Chloro-acetic acid and alcohol at 170° form PhN:C S.CH₂ [148°], which yields aniline, CO₂,

and thioglycollic acid on boiling with water (Liebermann a. Voeltzkoff, B. 13, 276).-15. Sodium malonic ether in presence of alcohol forms C14H18SNO4Na, whence acids liberate C₁,H₁,SNO₃, or (CO₂Et)₂CH.CS.NHPh [60°] (Michael, J. pr. [2] 35, 451).—16. Sodium cyanamide and alkyl iodides form cyano-phenylalkyl-thio-ureas (Hecht, B. 23, 1664). In this way the following compounds may be prepared: NHPh.CS.NMeCy [186°], NHPh.CS.NEtCy [144°], NHPh.CS.NCyC.H. [100°], crystallising in needles, NHPh.CS.NCyC.H. [108°], crystallising in needles, and NHPh.CS.NCy.CH.Ph [182°]. 17. Aldehyde-ammonia forms $C_{22}H_{21}N_5S_2O_2$ [148°] crystallising from alcohol in silvery needles (R. Schiff, B. 9, 567; G. 6, 244).—18. Benzamidine hydrochloride and dilute NaOHAq form CPh(NH).NH.CS.NHPh [125°] (Pinner, B. 22, 1609). Prisms.

References .- Bromo-, Chloro-, Iodo-, Nitho-, and Oxy- PHENYL-THIOCARBIMIDE

PHENYL-THIOCARBIZINE [129°]. Formed by heating phenylthio-semicarbazide (10 g.) with dilute (20 p.c.) HClAq (30 c.c.) for 12 hours at 130° (E. Fischer, A. 212, 326). Plates (from water), v. e. sol. alcohol. May be distilled. McI forms C,H₂McN₂S [123°]. Does not reduce Fehling's solution. HIAq at 200° forms aniline, H₂S, CO₂, and NH₃. Aqueous NaOCl forms a violet pp. which dissolves in H₂SO₄ with deep-red colour. Br in chloroform gives C,H₅BrN₂S [210°] crystallising in needles.

Salts. — B'HCl. [240°]. Needles. — B'₂H₂PtCl₃. Yellow prisms.—C₇H₃N₂SAg: floculent pp. got by adding ammoniacal AgNO₃.

Acetyl derivative C,H,AcN₂S. [187°]. Got by heating phenyl-thiocarbizine with Ac₂O (F.). An isomeride [74°] (275°), got by heating acetyl-phenyl-hydrazine with CSCl₂ and benzene (Freund, B. 21, 2468), is converted by boiling with alcohol and HgO into acetyl-phenyl-carbizine.

Benzoyl derivative C,H₅BzN₂S. [180°]. Got from the base and BzCl (Fischer). An isomeride [110°] is got by heating NHBz.NPhH with COCl₂ in benzene at 100° [F). PHENYL THIOCARBONATE CS(OPh)₂.

PHENYL THIOCARBONATE CS(OPh)₂. [97°]. Formed from NaOPh and CSCl₂ (Bergreen, B. 21, 346). White crystals, sol. alcohol. PHENYL-TRI-THIO-ORTHOFORMATE

CH(SPh)₃. [40°]. Got by boiling aqueous NaSPh with chloroform (Gabriel, B. 10, 185). Thick prisms, sol. ether. Split up by fuming HClAq at 100° into formic acid and PhSH.

PHENYL-THIOGLYCOLLIC ACID C₈H₈SO₂, i.e. CH₂(SPh).CO₂H. [62°]. Formed by saponifying its ether, which is got by adding ClCO₂Et to a solution of NaSPh in absolute alcohol (Claesson, Bl. [2] 23, 441; Gabriel, B. 12, 1639). Thin tables, sl. sol. cold water. Oxidised by KMnO₄ to phenyl methyl sulphone (Otto, B. 19, 3138). Salts.—KA'.—NaA'.—BaA'₂.—CaA'₂.—MgA'₂3aq.—CdA'₂aq.—ZnA'₂2aq.—PbA'₂. [60°].—MnA'₂5aq.—CuA'₂.—AgA' aq: crystalline pp. Ethyl ether EtA'. (277°). S.G. ½ 1·136;

Amide. [104°]. Tables (from alcohol).

References. — Amido- and Bromo- PhenylTHIOGLYCOLLIC ACIDS.

(a)-PHENYL-THIOHYDANTOÏC ACID

C_sH₁₀N₂SO₂, i.e. NH₂.C(NPh).S.CH₂CO₂H. [148°–152°]. Got by heating aniline with ammonium sulphocyanide, chloro-acetic acid, and alcohol (Claesson, B. 14, 732). Needles (from HOAC); m. sol. hot water and ether. Cold NaOHAq converts it into C₁₈H₁₇N₃S₂O₄ [112°–115°] (Liebermann, A. 207, 129).

(β)-Phenyl-thio-hydantoïc acid

NHPh.C(NH).S.CH₂.CO₂H. Got by evaporating an alcoholic solution of ammonium chloroacetate and phenyl-thio-urea at 100° (Meyer, B. 14, 1659). Decomposes on heating without melting. Insol. water, alcohol, and ether. Boiling

dilute HCl forms CO NPh.CO [148°].

Isomeride v. Phenyl-thio-hydantoïn. Di-phenyl-hydantoïc acid

NHPh.C(NPh).S.CH., CO.H. Got from potassium chloro-acetate and di-phenyl-thio-ures (Lange, B. 12, 597). Yellow octahedra.

PHENYL-THIO-ψ-HYDANTOÏN C.H.N.OS,

i.s. C(NH) NPh.CO [178°]. Got from chloro-

acetic anilide and thio-urea, and obtained also by boiling (B)-phenyl-hydantoic acid with HOAc or conc. NH₈Aq (Meyer, B. 14, 1661), and by the action of phenyl-cyanamide on thioglycollic acid (Andreasch, B. 15, 324). Needles, Bol. NaOHAq.

Phenyl-thio-hydantoïn CS NPh.CO_{NH.CH}. Got by melting glycocoll with phenyl-thiocarbimide (Aschan, B. 17, 424). Monoclinic plates, decomposing near 200°. Boiling alcoholic potash converts it into a salt of an oily phenyl-thio-hydan-

toïc acid NHPh.CS.NH.CH2.CO2H which, when

set free, rapidly changes again to phenyl-thio-hydantoin.

Di-phenyl-thio-hydanto'in $C_{1b}H_{12}N_2SO$, i.e. CS < NPh.CO? [176°]. Got from chloro-acetic acid and di-phenyl-thio-urea (Lange, B. 12, 595). Iridescent plates, insol. water, v. sol. hot alcohol. Boiling dilute HClAq forms the compound $C_bH_1NSO_2$ [148°].— $B'_2H_2PtCl_s$. Yellow needles.

Di-phenyl-thio-hydantoin CS NPh.CO NH. CHPh [233°]. Formed by the action of alkalis on NHPh.CS.NH.CHPh.CO₂Et (Kossel, B. 24, 4152).

Reference.—OXY-DI-PHENYL-THIOHYDANTOIN.

PHENYL-DI-THIO-DI-METHYL-KETURET NH CHPh.S CMe₂. [239°]. Formed from phenyl-di-thio-biuret and acetone (Fromm, B. 25, 1278). Sol. alkalis. Converted by benzyl chloride into C,H,N CHPh S CMe₂ [128°] which is insol. alkalis.

DI-PHENYL-THIO-OXAMIDE C₂S₂(NHPh), [133°]. Formed from di-phenyl-oxamide by successive treatment with PCl₃ and H₂S (Wallach, B. 13, 527). Golden plates.

(a)-PHENYL-THIOPHENE C₁₀H₃S i.e.

 C_4SH_3Ph . [41°]. Formed by heating either CH_Bz.CH(CO_H)₂ or β -benzoyl-propionic acid with P_2S_4 (Kues a. Paal, B. 19, 3141). Small tables, insol. water, v. e. sol. ether, v. sol. alcohol. Volatile with steam. With isatin and cold H_2SO_4 it gives a splendid bluish-violet colour, changing to dark blue. Bromine gives rise to

C_{1e}H₄Br₁S [56°] and C₄SBr₂.C₆H₄Br [1:4] [146°].
(β).Phenyl-thiophene C₄H₃PhS. [170°].
(330°). A product of the passage of the vapours of toluene and sulphur through a red-hot tube (Renard, C. R. 109, 699). White spangles, m. sol. cold alcohol. Gives a blue colour with isatin and H₂SO₄, and a green colour with phenanthraquinone and H₂SO₄. Yields C₄H₃BrS.C₆H₄Br[1:4] [195°], C₄H₂(NO₂)S.C₆H₄,NO₂ [1:4] [178°], a disulphonic acid, and a tetrasulphonic acid.

Di-(a)-phenyl-thiophene S CPh:CH. [153°] Formed by heating C₂H₄Bz₂ with P₂S₅ at 180° (Paal, B. 21, 3058). White plates (from alcohol).

Tri-phenyl-thiophene C₄Ph₂HS. [127°]. Got by heating desyl-acetophenone with P₂S₃ at 150° (A. Smith, C. J. 57, 647). Plates, v. sol. benzene. Gives a greenish-yellow colour with H₂SO₄ and isatin or phenanthraquinone.

Tetra-phenyl-thiophene C₂₅H₂₆S i.e. S

CPh:CPh. Thionessal. [182°].

Formation.—1. By the dry distillation of thiobenzoic aldehyde (Laurent, A. 52, 354).—2. By distilling benzyl sulphide or benzyl di-

sulphide (Märcker, A. 136, 94).—3. By distilling benzylidene sulphide (Fleischer, A. 140, 239; 144, 194).-4. By distilling barium phenyl-acetate with S (Forst, A. 178, 876).-5. By heating phenyl-acetic acid with S for 6 hours at 260° (Ziegler, B. 23, 2473).—6. By heating phenyl benzyl ketone with S.—7. By heating s-diphenyl-ethylene with S at 250° (Baumann a. Klett, B. 24, 3311). Minute needles, sl. sol. alcohol, v. sol. benzene and CS,. Yields a sulphonic acid, and also the following derivatives: $C_{2a}H_{16}Cl_2S$ [219°] (Dorn, A. 153, 350), $C_{2a}H_{1}$, Br_2S [265°–270°], $C_{2a}H_{16}(NO_2)_4S$ [above 250°].

DI - PHENYL - DI - THIO - DI - PHENYL - KE-

TATE CPh₂(SPh)₂. Phenyl-mercaptol of benzophenone. [139°]. Formed by passing HCl gas into a hot mixture of benzophenone, phenylmercaptan, and ZnCl₂ (Baumann, B. 18, 888). Short prisms, sl. sol. alcohol and ether.

THIOPHOSPHATE PS(OPh), PHENYL [49°]. (above 360°). Got by boiling PSCl, with phenol (Schwarze, J. pr. [2] 10, 222) and by heating phenyl phosphate with K_2S (Kreysler, B. 18, 1718), or with S at 190° (Anschütz, A. 253, 118). Needles (from alcohol), turning red in air and light. Not decomposed by water.

Phenyl tri-thiophosphate PO(SPh)₂. [72°]. Got by boiling PhSH with POCl₃. Prisms (from alcohol). Decomposed by warm water.

Phonyl PS(SPh)3. tetra-thio-phosphate [86°]. Got from PhSH and PSCls. Silky white needles, not affected by boiling water.

PHENYL THIOSULPHURIC ACID C.H.S.SO2.OH. Phenyl hydrogen thiosulphate. Got by dissolving Ph₂S in H₂SO₄ (Stenhouse, Pr. 17, 62; A. 149, 254).—BaA'₂ 2aq: crystalline.
PHENYL-THIO-URAMIDO-ACETIC ORTH-

ALDEHYDE. Di-ethyl derivative NHPh.CS.NH.CH₂.CH(OEt)₂. [96°]. Got by mixing phenyl-thiocarbimide with amido-acetal NH₂.CH₂.CH(OEt)₂ (Wohl a. Marckwald, B. 22, 569). Needles (from dilute alcohol), insol. water and ligroin, v. sol. ether. Cold conc. H2SO4 forms NHPh.CS.NH.CH₂.CH(OH)(OEt) [94°] which yields, by loss of H₂O, the salts $(C_{11}H_{14}N_2SO)_2H_2PtCl_e$ and $C_{11}H_{14}N_2SOC_eH_sN_3O_{7}$ [190°].

PHENYL-THIO-URAMIDO-BENZOIC ACID

v. m-Amido-Benzoic acid.

o-PHENYL - THIO - URAMIDO - CINNAMIC ACID NHPh.CS.NH.C,H,.CH:CH.CO,H. [237°] Formed by heating o-amido-cinnamic acid with phenyl-thiocarbimide on the water-bath (Rothschild, B. 23, 3343)

DI-PHENYL - DI-THIOURAMIDO - TETRA-OXY-DI-PHENYL. Tetra-methyl derivative C₁₂H₄(OMe)₄(NH.CS.NHPh)₂. [184°]. Formed from C₆H₂(OMe)₂(NH₂).C₆H₂(OMe)₂(NH₂) and phenyl thiocarbimide (Baessler, B. 17, 2128). White flakes, sol. hot alcohol and benzene.

PHENYL-THIOURAMIDO-PHENYL-ACETIC ETHER NHPh.CS.NH.CHPh.CO.Et. [162°]. Formed from phenyl-amido-acetic ether and phenyl-thiocarbimide (Kossel, B. 24, 4151). phenyl-thiocarbimide (Kossel, White crystals, sol. hot alcohol.

PHENYL-THIO-URAMIDO-THIO-FORMIC ACID v. PHENYL-DI-THIO-ALLOPHANIC ACID.

PHENYL - THIOURAMIDO - TOLYL - CARB-AMIC ETHER NHPh.CS.NH.C.H.Me.NH.CO.Et. [155°]. Formed, together with CS(NHPh)2, by the action of phenyl-thiocarbimide on amidotolyl-carbamic ether (Schiff, B. 23, 1818). Prisms (from alcohol), v. sl. sol. alcohol.

DI-PHENYL-DI-p-THIOURAMIDO -DI. TOLYL-o-SULPHIDE $C_{2a}H_{2a}N_iS_i$ i.e. $S(C_aH_aMe.NH.CS.NHC_aH_a)_2$. Thio-p-tolyl-diphenyl-thiourea. [184°]. Formed by the action of phenyl-thiocarbimide upon di-p-amido-di-tolylsulphide (Truhlar, B. 20, 670). White prismatic needles. V. sol. alcohol, ether, and benzene.

PHENYL-THIO-UREA C, H, N, S i.e. NHPh.CS.NH₂. Mol. w. 152. [154°]. S. 26 at 18°; 5.93 at 100°. S. (alcohol) 5.59 at 16°; 68 at 78°.

Formation.—1. By heating phenyl sulphocyanide with alcoholic NH₃ (Hofmann, Pr. 9, 276).—2. By boiling aniline with ammonium sulphocyanide (Schiff, A. 148, 338; Rathke, B. 18, 3104).—3. By heating aniline sulphocyanide at 190° (Salkowski, B. 24, 2728; cf. De Clermont, C. R. 82, 512).-4. By passing H₂S into a solution of phenyl-cyanamide in benzene (Weith, B. 9, 819).

Properties.—Needles, sol. alkalis and reppd. by acids. Tastes bitter.

Reactions .- 1. Heated in a scaled tube at 180° it yields CS(NHPh)2, ammonium sulphocyanide, NH₃, H₂S, and aniline.—2. Ammonia at 140° forms aniline and ammonium sulphocyanide.—3. Boiling aniline forms CS(NHPh) -4. HClAq at 120° forms aniline, H2S, NH2, and CO₂.-5. By boiling with alkaline lead acetate solution it gives phenyl cyanamide N:C.NHC,H, which polymerises on heating to tri-phenyl-isomelamine (PhN:C)_s(NH)_s.—6. When boiled for several hours with freshly-precipitated HgO it yields asymmetrical tri-phenyl-melamine PhN C(NH).NPh C.NHPh (Hofmann, B. 18, 3220).—7. Cl.CO.CO₂Et forms oxalyl-di-phenylbiuret, NH CS.NPh C2O2 (v. CHLORO-GLY-OXYLIC ETHER) .- 8. ClCO2Et forms phenyl-thioallophanic ether .- 9. Silicon tetrabromide gives solid SiBr₄(C₇H₈N₂S)₄, sol. hot benzene, but decomposed by boiling alcohol (Reynolds, C. J. 53, 856).—10. An alcoholic solution of allyl bromide forms a base C₁₀H₁₂N₂S (Werner, C. J. 57, 302).—11. Bensyl chloride in alcohol yields CS(N₂H₃Ph)C,H,Cl [112°] (W.).—12. Ethylene bromide forms C₁₀H₁₈N₂S [139°], which gives the salts B"2HBr [214°], B"2HCl [218°], B"H₂PtCl_g, and B"2C₆H₂(NO₂) OH [196°] (Bertram, B. 25, 59).—13. H₂O₂ oxidises it, in alcoholic solution, to di-phenyl-di-amido oiazthiole (Hector, B. 22, 1176).

Salts.—B'₂H₂PtCl₈.—B'₆Cu₂Cl₂ 6aq. [145°] (Rathke, B. 17, 305). Trimetric crystals (from alcohol).

derivative NHPh.CS.NHAc. [173°]. Formed from acetyl sulphocyanide and aniline (Miguel, A. Ch. [5] 11, 318), and by the action of Ac₂O on the product of the action of phenyl thiocarbimide on aldehyde-ammonia (Schiff, B. 9, 570). Tables (from dilute alcohol).

—B'HBr. [270°]. Prisms, v. e. sol. water.

Benzoyl derivative NHPh.CS.NHBz. [149°]. S. (alcohol) 4 at 78°. Needles, insol. water (Miguel).

o-Oxy-bensoyl derivative NHPh.CS.NH.CO.C.H.OH. [192 [192°]. Formed from salicyl-thiocarbimide and aniline (M.).

Di-phenyl-thio-urea C₁₃H₁₂N₂S i.e. CS(NHPh)₂. Mol. w. 228. [151°] (Lellmann, 4. 221, 21; Losanitsch, B. 19, 1821).

Formation. - 1. By boiling aniline with CS. and alcohol (Hofmann, A. 57, 266; 70, 144). 2. By heating aniline with potassium sulphocyanide and H₂SO₄ (Laurent a. Gerhardt, A. Ch. [3] 22, 103; 24, 196).—3. From aniline and phenyl-thiocarbimide.—4. From di-phenyl-cyanamide and H₂S (Weith, B. 7, 1303).—5. By boiling s-di-phenyl-guanidine with CS2 (Hofmann, B. 2, 460).

Properties.—Trimetric crystals (from alcohol), nearly insol. water, v. sol. alcohol and ather. Sol. alkalis and reppd. by acids. Has a very bitter taste.

Reactions.—1. Partially decomposed on distillation into tri-phenyl-guanidine, CS₂, and H₂S.—2. P₂O₅, ZnCl₂, or HCl split it up into aniline and phenyl-thiocarbimide.—3. Boiling alcoholic potash forms di-phenyl-urea.—4. Diphenyl-urea is formed when the alcoholic solution is boiled with HgO.-5. A cold solution of nitrous ether in alcohol forms tri-phenyl-guanidine, phenyl thiocarbimide, and S (Claus, B.4, 143).-6. Iodine, added to its boiling alcoholic solution, forms tri-phenyl-guanidine and phenyl thiocarbimide. - 7. Aniline forms, on heating, tri-phenyl-guanidine and H₂S.-8. Sodiumamalgam reduces it to aniline, H₂S, and CH₄ (Merz a. Weith, Z. [2] 4, 612).—9. When heated with HgCl₂ it yields tri-phenyl-guanidine (Buff, 2, 498).—10. Fuming H₂SO₄ decomposes it on heating, forming aniline, COS, and SO₂ (Fleischer, B. 9, 993).—11. COCl₂ forms C₁₄H₁₀N₂SO [87°], crystallising in prisms and split up by heat into COS and di-phenyl-cyanamide (Will, B. 14, 1486).—12. CSCl₂ in benzene forms C₁₄H₁₀N₂S₃, crystallising in yellow needles [156°], sl. sol. alcohol. In presence of ether CSCl2 forms NPh:C < NPh > CS [79°] (Freund a. Wolf, B. 25, 1459).-13. H2S at 170° forms aniline and CS (Weith, B. 7, 1304).—14. Alkyl iodides form alkyl derivatives, which may perhaps be formulated as derivatives of ψ -thio-ures. Thus, MeI forms NHPh.C(SMe):NPh [110°], which is decomposed by heat into MeSH and C(NPh), and by alco-holic potash into MeSH and CO(NHPh). In like manner EtI forms C₁₅H₁₆N₂S [79°] (v. Dr-PHENYL-ETHYL- ψ -THIO-UREA), and ethylene bromide gives $C_{15}H_{14}N_2S$ [136°] (Will, B. 14, 1490).— 15. Allyl bromide in alcohol yields O₁₈H₁₇N₂SBr, crystallising in trimetric prisms [171°] (Werner, C. J. 57, 303).—16. Benzyl chloride in alcohol forms C₂₀H₁₂N₂SCl, crystallising in prisms [153°] (W.).-17. Phenyl cyanate in presence of benzene at 180° forms di-phenyl-urea and phenyl sulphocyanide (Goldschmidt a. Meissler, B. 23, 271).—18. Ac₂O yields acetanilide and some phenyl-thiocarbimide (Werner, C. J. 59, 398). 19. Chloro-glyoxylic ether forms di-phenyl-di-thio-parabanic acid C₁₈H₁₁N₂S₂O₂ [215°] and C_{2.}H₁₇N₂S₂O [281°], which by boiling with alco-hol and AgNO₂ is converted into di-phenylparabanic acid, aniline, and H.S. while by boiling alcoholic ammonia it is converted into Ing accounts almost it is converted in the converted in the converted by funing HNO, forms C₁,H₁,N₂O₂ [285°]. The compound C₂,H₁,N₃S₂O is converted by fuming HNO, into O₁,H₂N₄SO₃ [285°] (Stojentin, J. pr. [2] 32, 3).

Tri-phenyl-thio-ures NHPh.CS.NPhr [152°] Formed by heating phenyl-thiocarbimide with diphenylamine at 280° (Gebhardt, B. 17, 2092, 8036). Long white needles, v. sol. hot alcohol.

Tetra-phenyl-thio-urea CS(NPh2)2. [195°] Formed by heating tetra-phenyl-guanidine with CS₂ at 270°; the yield being 60 p.c. of the theoretical amount (Bernthsen a. Friese, B. 15, 1530). Formed also from NHPh, and CSCl, (Bergreen, B. 21, 340). Long needles, sol. hot alcohol, insol. alkalis. Yields diphenylamine on heating with diluted (70 p.c.) H₂SO₄.

Di-phenyl-thio-ures v. Di-PHENYL-DI-PHENYL-

References .- AMIDO-, BROMO-, CHLOBO-, IODO-, NITRO-, and OXY- PHENYL-THIO-UREA.

PHENYL-THIO-UREA CARBOXYLIC ACID

v. m-Amido-benzoic acid, Reaction 7.

DI-PHENYL-TOLENYL-AMIDINE C20H18N2 i.e. C.H.Me.C(NPh).NHPh. [168°]. Got by warming C.H.Me.C(OEt):NH2Cl with aniline and alcohol (Glock, B. 21, 2656). Plates (from benzene), v. sol. alcohol.

PHENYL-TOLUENE v. METHYL-DIPHENYL. DI-PHENYL-TOLUIC ALDEHYDE C2H16O i.e. CPh₂H.C_eH₄.CHO. [-15°]. (190°-195° at 46 mm.). Formed by heating terephthalic aldehyde with benzene and conc. H₂SO, at 100° (Oppenheimer, B. 19, 2028). Oil. Reduces ammoniacal AgNO, being itself oxidised to triphenyl-methane carboxylic acid. KMnO4 gives tri-phenyl-carbinol p-carboxylic acid. Reacts with phenyl-hydrazine. Yields C20H16ONaHSO3,

a crystalline pp., sl. sol. water, insol. alcohol.

PHENYL-o-TOLUIDINE C₁₂H₁₃N i.e.

[1:2]C₆H₄Me.NHPh. [41°]. (305° i.V.) at 728

mm. (Graebe, A. 238, 863). Formed, together with NHPh, and ditolylamine by heating o-toluidine with aniline hydrochloride at 280° (Girard a. Willm, Bl. [2] 25, 248). Coloured violet-blue

by HNO₃.

Phenyl-m-toluidine. (300°-305°). Got by heating [1:3:5] C₆H₈Me(NHPh).OH with zincdust (Zega a. Buch, J. pr. [2] 83, 542). Oil.

Coloured green by H₂SO₄ containing HNO₂.

Phenyl-p-toludine C,H,NHPh. [87°].

(318° i.V.) at 728 mm. (Graebe, A. 238, 363).

Formed by distilling tri-tolyl-rosaniline acetate (Hofmann, A. 132, 291). Obtained also, together with diphenylamine and ditolylamine by heating p-toluidine (1 mol.) with aniline (1 mol.) and HCl (1 mol.) at 230° (De Laire, Girard, a. Chapoteaut, Bl. 1867, i. 360; A. 140, 347). Prepared by heating phenol with p-toluidine and ZnCl2, or p-cresol with aniline and ZnCl, at 200°, or with SbCl, (Buch, B. 17, 2634). Crystals. Coloured blue by HNO₂. With benzoic acid and ZnCl₂ at 260° it yields phenyl-methyl-acridine C20H15N (Bonna, A. 239, 60). Bromine vapour forms a tetra-bromide [135°], a hepta-bromo-derivative [185°], and a compound NC13H12Br, or NC13H Br, [254°]. Excess of Br at 810° forms NC, H. Br. [296°] (Bonna). Bromine in HOAc added to an alcoholic solution forms a tetra-bromo-derivative [156°]

Salt.-B'HCl: plates. Acetyl derivative C₁,H₁,AeN. [51°]. Yields C₁,H₁₁BrAcN [72°], crystallising in plates. Bensoyl derivative O.H.,NO. Orystals. Yields orystalline C.H.(NO.).NBz.C.H.(NO., Nitrosamine C.H.(NPh.NO. [82]. Yellow

acedles (from ligroin). Br forms C₁₃H₁₁BrN₂O | [166°] (Bonna, A. 239, 56).

HENYL-TOLYL-ACETAMIDINE C, H, N2 i.e. CH₃.C(NC₂H₂).NHPh. [76°]. Formed from acetyl p-toluidine by successive treatment with PCl, and aniline (Wallach a. Fassbender, A. 214, 206).—B'2H2PtCl6.

PHENYL-p-TOLYL-ACETIC ACID

C.H.,CHPh.CO₂H. [115°]. Formed from abromo-phenyl-acetic acid, toluene, and zincdust (Zincke, B. 10, 996; Neure, A. 250, 149). Plates (from water), v. sol. alcohol.—KA' 4aq. NaA' 6aq.—CaA'22aq: needles (from dilute alcohol).

Ethyl ether EtA'. [34°]. Plates. Amide C₁₅H₁₅NO. [151°]. Nitrile C,H,CHPh.CN. [59°]

Phenyl-di-tolyl-acetic acid C₂₂H₂₀O₂ [78°-83°]. Got by oxidising (C,H,)2CPh.CO.C,H, (Thörner, A. 189, 123). Amorphous powder, insol. water, v. sol. alcohol.

PHENYL-TOLYL-AMINE v. PHENYL-TOLU-DINE

PHENYL-p-TOLYL-BENZYL-BIURET

NHPh.CO.N(CH₂Ph).CO.NHC₆H₄Me. [c. 100°]. Formed from p-tolyl-benzyl-urea and phenyl cyanate (Kühn a. Henschel, B. 21, 505). Needles (from dilute alcohol).

PHENYL-p-TOLYL-BENZYL-UREA

 $C_{21}H_{20}N_{2}O$. [113°]. Got from p-tolyl-benzylcarbamic chloride and aniline (Hammerich, B. 25, 1823). Dimorphous: prisms or plates, sol. ether and alcohol.

PHENYL-DI-p-TOLYL-BIURET C22H21N3O2 i.e. NHPh.CO.N(C_eH₄Me).CO.NHC_eH₄Me. [140°]. Formed by heating di-p-tolyl-urea with phenyl cyanate at 170° (Kühn a. Henschel, B. 21, 505). Crystals (from alcohol).

Di-phenyl-tolyl-biuret

NHPh.CO.NPh.CO.NHC, H, Me. [216°]. Got by heating di-phenyl-urea with p-tolyl cyanate at 170° (K. a. H.).

PHENYL-p-TOLYL-CARBINOL C_{1.}H_{1.1}O i.e. C₂H₄Me.CPhH.OH. [53°]. Got by reducing phenyl tolyl ketone in alcoholic solution by sodium-amalgam (E. a. O. Fischer, A. 194, 265). Stellate groups of needles (from ligroin).

Di-phenyl-tolyl-carbinol $C_{20}H_{18}O$ i.e. C.H.Me.CPh.OH. [150°]. Formed by oxidising di-phenyl-tolyl-methane with CrO, and HOAc (F.). Crystals (from ligroin). May be distilled.

PHENYL-p-TOLYL-CARBINOL o-CARB-OXYLIC ACID. Lactone v. Tolyl-Phthalide.

Di-phenyl-tolyl-carbinol carboxylic acid. $C_{e}H_{s}Me < CO^{CPh_{2}}>0.$ Lactone (above 360°). Formed by oxidising di-phenylm-xylyl-methane with chromic acid mixture (Hemilian, B. 16, 2361; 19, 3063). Prisms.

 $\begin{bmatrix} 4_1^2 \end{bmatrix} C_a H_a M_0 < \stackrel{CPh_a}{CO} > 0,$ isomerio [179°], (above 360°), is got by oxidation of diphenyl-p-xylyl-methane, and is accompanied by 1:2:5]CPh2(OH).C.H.Me.CO2H, which yields BaA'2 and CaA', and does not form a lactone.

PHENYL-o-TOLYL-CARBINYLAMINE [1:2]C.H.Me.CHPh.NH. (299°) at 721 mm. Formed by reducing the oxim of phenyl o-tolyl ketone dissolved in alcohol by adding sodium-

amalgam and HOAc (Goldschmidt a. Stöcker, B. 24, 2806). Yields an acetyl derivative [124°]. -B'HCl. [249°]. Needles, m. sol. water.

m-Isomeride. (299°) at 724 mm. Yields B'HCl [243°] and an acetyl derivative [97°].

p-Isomeride. (296°) at 723 mm. Yields an acetyl derivative [131°] and the salts B'HCl [252°], B'₂H_{*}PtCl₆ 2aq [119°], B'C₄H_{*}O₆ [157°], B'₂C₄H₆O₆ [73°]. The hydrochloride is constant verted by potassium sulphocyanide into the compound C₆H₄Me.CHPh.NH.CS.NH₂ [101°] and by potassium cyanate into the urea derivative $C_6H_4Me.CHPh.NH.CO.NH_2$ [158°].

PHENYL TOLYL - CARBINYL KETONE CARBOXYLIC ACID C₁₆H₁₄O₃ i.e. C₆H₄Me.CH₂CO.C₆H₄.CO₂H. [112°]. Formed $C_6H_4 < CO \xrightarrow{C(\hat{C}H.C_6H_4Me)} > 0$ by heating with KOHAq at 100° (Heilmann, B. 23, 3160). Vitreous prisms.—AgA'. Converted by heating with an alcoholic solution of hydroxylamine hydrochloride into C₆H₄<CO ____ON

[134°].

PHENYL-o-TOLYL-CYANAMIDE

NPh:C:NC,H,Me. Carbophenyltolylimide. (320°-325° uncor.). Formed by adding HgO to à boiling solution of phenyl-o-tolyl-thiourea in dry benzene, the yield being 65 p.c. of the theoretical. Oil. On keeping, it polymerises to a brittle glassy solid [68°-71°]. V. sol. benzene, sl. sol. ether. By boiling with dilute alcohol it is converted into phenyl-o-tolyl-urea. passed into its boiling solution in dry toluene converts it into phenyl-o-tolyl-thiourea. Heated with CS₂ at 180°-200° it yields a mixture of phenyl- and o-tolyl-thiocarbimides. bines with o-toluidine, forming u-phenyl-di-otolyl-guanidine (Huhn, B. 19, 2410)

Phenyl-p-tolyl-cyanamide NPh:0:NC,H,. (325°-380° uncor.). Resembles the preceding isomeride in mode of preparation and properties.

DI-PHENYL-TOLYLENE-TETRA-AMIDO. METHANE C20H20N, i.e.

 $C_1H_1 < NH > C(NHPh)_2(?)$. [161°]. Formed by heating C(NPh)₂ with tolylene-o-diamine at 135° (Dahm a. Gasiorowski, B. 19, 3057). Needles (from benzene).—B'23HCl. [174°].—B'H2SO.

DI-PHENYL-m-TOLYLENE DIAMINE C₁₀H₁₀N₂ i.e. [1:3:5]C₀H₃Me(NPhH)₂. [105°]. Formed by heating oroin (1 pt.) with aniline (4 pts.) and a mixture of ZnCl₂ and CaCl₂ at 220° (Zega a. Buch, J. pr. [2] 33, 542). White needles (from glacial HOAc).

Di-acetyl derivative [160°]. Di-benzoyl derivative [1900] Dinitrosamine C₁₉H₁₆N₄O₂. [170°]. Reference. — DI-NITBO-PHENYL-TOLYLENE-DI-AMINE

PHENYL-o-TOLYLENE-GUANIDINE

C,H, NH C:NPh. [167°]. Mol. w. 229 (calc. 223). Formed from o-tolylene-diamine and diphenyl-cyanamide (Keller, B. 24, 2514). Needles (from warm alcohol). Di-p-tolyl-cyanamide

>C:NPh >C(NHC,H,)2 [1939]. forms

while phenyl cyanide gives the compound

>C:NPh NPh [234°].-B'HCl.-

B'2H2PtCl6.—B'2H2SO4: white needles.

Mono-acetyl derivative. [147°]. Di-benzoyl derivative. [222°]. Prisms.

Nitrosamine C, H,2N,O. [125°]. DI-PHENYL - m - TOLYLENE-DI-METHYL-DI-AMINE C₂₁H₂₂N₂i.e. C₆H₃Me(NPhMe)₂. [124°]. Formed from C₇H₆(NHPh)₂, NaOH, and MeI (Zega a. Buch, J. pr. [2] 33, 546). White laminæ

(from HOAc)

DI - PHÉNYL-o-TOLYLENE - DI - THIO - DI-UREA [1:3:4]C₆H₃Me(NH.CS.NHPh)₂. [c. 150°]. Formed from tolylene-o-diamine and phenylthiocarbimide in alcohol (Lellmann, A. 221, 19). Thin plates (from dilute alcohol), v. sol. warm NaOHAq, sl. sol. water.

Di-phenyl-m-tolylene-di-thio-di-urea [1:2:4]C₆H₃Me(NH.CS.NHPh)₂ [168°]. Formed from tolylene-m-diamine and phenyl-thiocarbimide (Lussy, B. 8, 670; Gebhardt, B. 17,

8046; Billeter a. Steiner, B. 18, 8293; 20, 228). ${f Di-phenyl-}p$ -tolylene-di-thio-di-urea

[1:2:5]C.H.Me(NH.CS.NHPh)₂. [181°]. Formed from tolylene-p-diamine and phenyl-thiocarbimide in alcohol (Lellmann a. Würthner, A. 228, Plates, nearly insol. alcohol. DI-PHENYL-TOLYLENE-DI-UREA

Formed $C_1H_6(NH.CO.NHPh)_2$. [above 300°]. by adding phenyl cyanate to an ethereal solution of tolylene-diamine [99°] (Kuhn, B. 18, 1477). White pp. Insol. water, alcohol, ether, and benzene, v. sl. sol. acetic acid. By boiling with aniline it yields s-di-phenyl-urea.

PHENYL-p-TOLYL-ETHANE C₁₅H₁₆, i.e. (286° uncor.). [27°]. CH₂Ph.CH₂.C₆H₄Me. Formed by reduction of tolyl benzyl ketone by HI and P (Mann, B. 14, 1646). White plates.

ĆH₃.CHPh.C₆H₄Me. Phenyl-p-tolyl-ethane (279°). Formed from C₆H₅.C₂H₄Br, toluene, and zinc-dust (Bandrowski, B. 7, 1016). Oil. Yields

p-benzoyl-benzoic acid on oxidation.

PHENYL-p-TOLYL-ETHYLENE C1.H14, i.e. CHPh:CH.C. H.Me. Methyl-stilbene. [120°]. Formed by boiling p-tolyl-benzyl-carbinol with dilute H₂SO₄ (Mann, B. 14, 1646), and also by distilling p-tolyl cinnamate (Anschütz, C. J. 47, 898; B. 18, 1945). Pearly plates with blue fluorescence, v. sl. sol. alcohol, v. e. sol. ether. Yields

C_{1s}H₁,Br₂ [187°].
DI - PHENYL - DI - p - TOLYL - ETHYLENE

OXIDE ? C_{2s}H₂₄O. Phenyl tolyl (a)-pinacolin.

[215°]. Formed by the action of zinc and HClAq on phenyl p-tolyl ketone in alcohol, and got also by heating a mixture of the ketone with phenyltolyl-carbinol, alcohol, and ZnCl₂ (Thörner a. Zincke, B. 11, 71; A. 189, 104). Minute needles, m. sol. hot alcohol. Conc. HClAq at 150° m. sol. hot alcohol. Conc. HClAq at 150° changes it into the isomeric (β)-pinacolin

 C_6H_4 .CO.CPh $(C_6H_4$ Me $)_2$ [137°] p-TOLYL ETHYLENE DISUL-PHENYL PHONE C.H., SO.C.H., SO.C.H., [162°]. Got by boiling C.H., SO.C.H., Cl with an alcoholic solution of solium toluene p-sulphinate (Otto, J. pr.

[2] 80, 199).

PHENÝL - TOLYL - ETHYLENE - ψ - THIO-UREA NPh:C N(C,H₁) CH₂. [128°]. Formed S.CH₂ by boiling the methylo-iodide of the ethylene derivative of tolyl-di-thio-carbamic acid with

aniline (Will a. Bidschowski, B. 15, 1315). Silky plates.

PHENYL - p - TOLYL - ETHYL - THIO-UREA C16H18N2S, i.e. NPhEt.CS.NHC,H7. [90°]. Formed from p-tolyl-thiocarbimide and ethyl-aniline (Gebhardt, B. 17, 2091). Colourless crystals. s-PHENYL-DI-o-TOLYL-GUANIDINE

CaHaN:C(NHC,H,)2. [102°]. Formed by the action of an alcoholic solution of aniline upon di-o-tolyl-thiourea in presence of PbO. Got also by the combination of C₆H₃.N:C:N.C₇H₇ with aniline (Huhn, B. 19, 2412). Felted needles. V. sol. warm alcohol, ether, and benzene.—B'HCl.—B'₂H₂Cl₂PtCl₄: orange-yellow tables.

u-Phenyl-di-o-tolyl-guanidine C₇H₇N:C(NHC₆H₅)(NHC₇H₇). [112°]. from o-toluidine and C₇H₇N:C:NC₆H₅ Formed (Huhn). Long needles, v. sol. alcohol. Salts.—B'HCl.—B'₂H₂PtCl₆: orange-yellow tables.

s-Di-phenyl-p-tolyl-guanidine C,H,N:C:(NHC,H,s)2. [121°]. Formed by the action of an alcoholic solution of p-toluidine upon di-phenyl-thiourea in presence of PbO (Hofmann, B. 2, 459; Huhn, B. 19, 2412). Felted needles. V. sol. alcohol and ether.— Felted needles. B'HCl.—B'₂H₂Cl₂PtCl₄: reddish-yellow pp. *u*-Di-phenyl-p-tolyl-guanidine C₂₀H₁₉N₃, *i.e.* C₈H₂N:C(NHC₈H₂)(NHC₂H₃). [127°].

Formation.—1. By heating phenyl-p-tolylthiourea with an alcoholic solution of aniline in presence of PbO.—2. By the combination of $C_6H_5N:C:NC_7H_7$, with aniline (Huhn, B. 19, 2409).

-B'HCl.—B'2H2Cl2PtCl: sparingly soluble pp. PHENYL TOLYL-IMIDO-BENZYL KETONE C.H. CO.C(NC,H.).C.H. Formed by heating o-toluidine with benzil at 100° (Bandrowski, M. 9, 689). Yellow plates (from alcohol).

PHENYL O-TOLYL KETONE C11H12O i.e. C₈H₈.CO.C₆H₁Me [1:2]. (314°) at 735 mm· (Smith, B. 24, 4046); (295°) at 722 mm. (G. a. S.). Formed by the action of AlCl, on a mixture of o-toluic chloride and benzene (Ador a. Rilliet, B. 12, 2301; Goldschmidt a. Stöcker, B. 24, 2805). Formed also, together with a much larger quantity of the p- isomeride, from BzCl, toluene, and AlCl, (Elbs, J. pr. [2] 35, 466). Oil. Yields anthracene on long boiling or on heating with zinc-dust.

syn-Oxim C,H,.C.C,H,. [69°].

HO.N anti-Oxim C,H,.C.C,H,. [105°] (S.). HO.K

Phenyl-m-tolyl ketone C_6H_5 . $CO.C_6H_4Me$ [1:3]. (315° i.V.) at 745 mm. S.G. 176 1.088. Formed by the action of AlCl, on a mixture of m-toluic chloride, benzene, and AlCl, (A. a. R.; G. a. S.). Got also by cautious oxidation of phenyl-m-tolylmethane (Senff, A. 220, 252). Oil, miscible with Reduced by HIAq at 200° to C14H14 alcohol. (269° at 725 mm.).

Oxim. [101°]. Reduced by sodium-amalgam in presence of HOAc to m-toluic anilide and

C.H.Me.CPhH.NH. Phenyl-p-tolyl ketone C.H.CO.C.H.Me [1:4]. [60°]. (827° i.V.).

Formation.—1. Together with the o-isomeride by heating benzoic acid with toluene and P₂O₃ at 200° (Kollarits a. Merz, B. 6, 446; Thörner, A. 189, 83). — 2. By oxidising C.H.Me.CPhH. (Zincke a. Plascuda, B. 7, 982).—8. By distilling a mixture of calcium p-toluate and benzoate (Radziszewski, B. 6, 810).—4. By heating BzCl with toluene and zinc at 190° (Grucarevic a. Merz, B. 6, 1243).—5. By the action of AlCl, on a mixture of p-toluic chloride and benzene (A. a. R.).-6. From BzCl, toluene, and AlCl.

Properties.-Prisms, m. sol. alcohol, v. sol. ether. Dimorphous. Yields p-benzoyl-benzoic acid on oxidation. Reduced to C,H,Me.CH,Ph by distillation over zinc-dust. Sodium-amalgam forms phenyl-tolyl-carbinol. Zinc and HClAq acting on the alcoholic solution form two pinacolins $C_{28}H_{24}O$. The (α)-pinacolin [215°] may be converted into the (B)-isomeride [137°] by heating with HOAc.

anti-Oxim $C_6H_5.C.C_6H_4Me.$ [154°] HO.N

(Hantzsch, B. 23, 2325, 2776; 24, 58; cf. Wegerhoff, A. 252, 11). Converted by HOAc and HCl into p-toluic anilide. Yields an acetyl derivative[124°] and a benzoyl derivative [85°] (Auwers, $B.\ 23,\ 399).$

syn-Oxim C_6H_5 .C. C_6H_4 Me. [116°]. Con-N.OH

verted by HOAc and HCl into benzoyl-toluidine and some toluic anilide. Yields an acetyl derivative [118°-122°] and a benzyl derivative [51°

Phenyl p-tolyl diketone C,H,.CO.COC,H, Formed by heating di-bromo-deoxybenzoin with water at 183° (Bucher, B. 22, 2819). Yellow oil, solidified by cold.

References .-- DI-AMIDO-, NITRO-, OXY-AMIDO-, and Oxy- PHENYL-TOLYL-KETONE.

PHENYL TOLYL KETONE CARBOXYLIC

ACID v. TOLUYL-BENZOIC ACID. Phenyl p-tolyl ketone dicarboxylic acid C₁₆H₁₂O₅ i.e. C₆H₅.CO.C₆H₂Me(CO₂H)₂ [1:4:2:6]. Benzoyl-uvitic acid [245°]. Formed by oxidising phenyl mesityl ketone with dilute HNO, (Elbs, J. pr. [2] 35, 489). Needles.—Ag₂A".

An isomeric acid, got by oxidation of phenyl ψ-cumyl ketone C₆H₅.CO.C₆H₂Me₈ [5:1:2:4], yields

Ag₂A".

 $\begin{array}{lll} \textbf{PHENYL-o-TOLYL-METHANE} & \textbf{C}_{14}\textbf{H}_{14} & \textbf{i.e.} \\ \textbf{C}_{6}\textbf{H}_{14}\textbf{C}\textbf{H}_{2}\textbf{C}_{6}\textbf{H}_{4}\textbf{Me} \ [1:2]. & o-Benzyl-toluene. & \textbf{Mol.} \end{array}$ w. 182. (284°). Formed, together with the p-isomeride, by heating benzyl chloride with toluene and zinc-dust (Zincke, B. 6, 906; Senff, A. 220, 249). Got also from C₆H₄Me.CH₂Cl, benzene, and zinc-dust (Barbier, B. 7, 1544). Yields anthracene when passed through a redhot tube.

Phenyl-m-tolyl-methane

 $C_6H_5.CH_2.C_6H_4Me$ [1:3]. (275° i.V.). S.G. 178 ·997. Got by reducing the ketone with HI and P (Ador a. Rilliet, B. 12, 2300). Prepared by heating ω -chloro-m-xylene (1 pt.) with benzene (7 pts.) and AlCl_s (Senff, A. 220, 230; cf. Barbier, C. R. 79, 660). Oil, sol. alcohol and ether. Gives a di-nitro- derivative [141°].

Phenyl-p-tolyl-methane C_eH_s.CH_s.C_eH₄Me [1:4]. (280° i.V.). Formed, with the o-isomeride, by heating benzyl chloride with toluene and zinc-dust (Zincke, B. 7, 1153; A. 161, 93). Formed also by the action of zincdust on benzyl chloride (Prost, Bl. [2] 46, 248), and by heating phenyl p-tolyl ketone with ammonium sulphide and S at 320° (Willgerodt, B. 20, 2470). Oil. Yields p-benzoyl-benzoic acid on oxidation. Does not form anthracene when Vol. IV.

passed through a red-hot tube. Yields a disulphonic acid [88°], which gives $K_2A'' 3\frac{1}{2}aq$, BaA'' $8\frac{1}{2}aq$, and CuA'' $4\frac{1}{2}aq$.

Phenyl-di-tolyl-methane CHPh(C,H,)2. [56°]. Got by heating $(C_1H_1)_2CPh.CO._2H_3$ with sodalime at 300° (Thörner a. Zincke, B. 11, 70).

u1 - pnenyi - o-tolyl - methane C_{.0}H₁₈ i.e. (C₈H₃)₂CH.C₈H₄Me. [59·5°]. (354°). Formed from leuconiline by from leucaniline by diazotising and boiling with alcohol (Fischer, A. 194, 282; Rosenstiehl a. Gerber, A. Ch. [6] 2, 342). Spherical groups of prisms (from MeOH), v. sol. ether. Oxidised by CrO_{\bullet} and HOAc to $(C_{c}H_{\bullet})_{s}C(OH).C_{c}H_{\bullet}Me$ [150°].

Di-phenyl-m-tolyl-methane. [62°]. (above 360°). Formed by distilling its carboxylic acid with baryta (Hemilian, B. 16, 2368). Long thin needles (from alcohol), v. sol. ether. The crystals emit light when powdered. Its dilute

solutions fluoresce blue.

Di-phenyl-p-tolyl-methane. [71°]. (above 360°). Formed by distilling its o-carboxylic acid, and also by the action of P2O, on a mixture of toluene and di-phenyl-carbinol or of benzene and phenyl-p-tolyl-carbinol (Hemilian, B. 7, 1209; 19, 3066; Fischer, A. 194, 263). Needles or prisms, v. sol. hot alcohol.

References .- DI-AMIDO-, CHLORO-, NITRO-. NITRO-AMIDO-, and OXY- PHENYL-TOLYL-METHANE.

PHENYL - p - TOLYL - METHANE -o- CARB-OXYLIC ACID C₆H,Me.CH₂.C₆H₄.CO₂H. [134°]. Formed by warming C₆H,Me.CO.C₆H₄.CO₂H with NH3Aq and zinc-dust (Gresly, A. 234, 236). Needles (from alcohol), nearly insol. water .-BaA'2: plates, v. sol. water.

Phenyl-tolyl-methane dicarboxylic acid CO₂H.C₆H₄.CH(C,H₇).CO₂H. [154°]. (above 300°). Got by saponifying the nitrile, which is prepared by treating an alcoholic solution of CN.C_sH₄.CH₂.CN with benzyl chloride and KOH (Eichelbaum, B. 21, 2679). Prisms, sol. alcohol.

Amide. [224]. Small plates.

Imide C₆H₄ CO.NH [176°]. (above 300°). Got by heating the amide with

conc. HClAq. Needles, v. sol. alcohol.

Nitrile. [110°]. (above 300°). Plates. Di-phenyl-p-tolyl-methane carboxylic acid C.H.Me.CHPh.C.H., CO.H. [155°]. Formed by warming p-tolyl-phthalide with benzene and

AlCl, (Gresly, A. 234, 242). Needles (from alcohol).—BaA', 23aq: sl. sol. water.

Di-phenyl-p-tolyl-methane carboxylic acid CHPh₂.C₆H₃Me.CO₂H[4:2:1]. [217°]. Formed from di-phenyl-p-tolyl-carbinol carboxylic acid, NaOHAq, and zinc-dust (Hemilian, B. 16, 2363). Transparent tables, v. sol. alcohol.—BaA', 4aq. AgA': minute needles, insol. water.

Di-phenyl-p-tolyl-methane carboxylic acid $CHPh_2.C_6H_3Me.CO_2H[4:1:2].$ [203°]. Formed from di-phenyl-methyl-phthalide, NaOHAq, and zinc-dust (Hemilian, B. 19, 3064). Needles or tables, sol. hot alcohol.—BaA'₂3aq.—AgA'.
PHENYL-p-TOLYL DI-METHYLENE DI.

SULPHONE KETONE C16H16S2O, i.E. C₆H₄.SO₂.CH₂.CO.CH₂.SO₂.C₆H₄Mc. Formed from CH.Br.CO.CH, SO₂.C.H.Me by treatment with C.H.SO₂Na and obtained also from C.H. SO. CH. CO.CH.Br and C.H. Me. SO. Na (Otto, J. pr. [2] 36, 427). Plates, m. sol. alcohol, v. sol. CHCl₃. H

PHENYL - 0 - TOLYL - METHYL - PYRROLE | a. Ebenhardt, B. 23, 3271).

NC,H, CPh:OH . [44°]. (327°). Formed by heating its carboxylic acid [199°] (Lederer a. Paal, B. 18, 2596). Plates, v. e. sol. alcohol.

 ${\bf Phenyl-} p{\bf -tolyl-methyl-pyrrole.}$ [91°]. (above 350°). Got by heating its carboxylic acid

[227°]. Tables (from ligroïn).

PHENYL - 0 - TOLYL - MÉTHYL - PYRROLE CARBOXYLIC ACID C.H.MeN CPh:CH

[199°]. Got by saponifying its ether, which is formed by boiling acetophenone-acetoacetic ether with o-toluidine and HOAc (Lederer a. Paal, B. 18, 2596). Small prisms (from alcohol).

p-Isomeride. [227°]. Made in like manner from p-toluidine. Plates, v. sol. alcohol.

Ethyl ether EtA'. [115°]. Crystals. PHENYL-0-TOLYL-METHYL-THIO-UREA C₁₅H₁₆N₂S i.e. NMePh.CS.NHC,H, [121°]. Formed from methyl-aniline and o-tolyl-thiocarbimide (Gebhardt, B. 17, 2091, 3035). Plates.

Phenyl-p-tolyl-methyl-thio-urea. Formed in like manner from p-tolyl-thiocarbimide (G.). Small trimetric tables. By distillation with steam it is resolved into the parent substances, which slowly recombine in the dis-

PHENYL-TOLYL-NAPHTHYL-GUANIDINE C24H21N3 i.e. $CN_3H_2Ph(C_7H_7).C_{10}H_7.$ 60°]. Formed by boiling an alcoholic solution of phenyl-tolyl-thio-urea and naphthylamine with PbO (Tiemann, B. 3, 6). Brittle resin, forming a crystalline hydrochloride.

PHENYL-DI-TOLYL-PHOSPHINE

PHENYL-DI-TOLYL-PHOSPHIAE

(C₆H₄Me)₂PC₆H₅. [57°]. Formed from p-bromotoluene (2 mols.), C₆H₅PCl₂ (1 mol.), and Na (Dörken, B. 21, 1512). Crystals, v. sol. ether.

Di-phenyl-tolyl-phosphine C₆H₄MeP(C₆H₅)₂.

[68°]. Formed from (C₆H₅)₂PCl, p-bromotoluene, and sodium (Dörken, B. 21, 1511).

Small prisms. Yields (C₆H₅)₂PO(C₆H₄Me)

[180°] and (C H₅)₂PS(C.H.Me) [189°]. [130°] and (C₆H₅)₂PS(O₆H₄Me) [139°].
PHENYL-TOLYL-(a)-PINACOLIN C₂₈H₂₄O v.

DI-PHENYL-DI-p-TOLYL-ETHYLENE OXIDE.

Phenyl-tolyl-(8)-pinacolin C₆H₂,CO.OPh(O,H₂)₂. [137°]. Formed by the action of AcCl on phenyl-tolyl-(a)-pinacolin or on phenyl-tolyl-pinacone (Zincke a. Thörner, B. 10, 1477; 11, 65, 1396; A. 189, 110). Prepared by boiling phenyl p-tolyl ketone with alcohol, HClAq, and zinc. Small dimetric tables, sl. sol. cold alcohol. On heating with soda-lime at 800° it yields phenyl-di-tolyl-methane.

PHENYL-TOLYL-PINACONE C₂₈H₂₈O₃ i.e. C₄H₄, CPh(OH).CPh(OH).C,H₄. [165°]. Formed by treating an alcoholic solution of phenyl ptolyl ketone with zinc and H₂SO₄ (Zincke a. Thörner, B. 10, 1476). Minute needles (from alcohol). Decomposed by fusion and by boiling alcoholic potash into phenyl tolyl ketone and phenyl-tolyl-carbinol. Converted into phenyltolyl-(8)-pinacolin by heating with dilute H₂SO₄ at 160°, with HOAc at 190°, with AcCl, or with HClAq. Alcoholic HCl in the cold forms

phenyl-tolyl-(a)-pinacolin.

PHENYL-TOLYL-PROPANE $O_{16}H_{18}$ CH2.CHPh.CH2.C6H4Me. The o- (817°), m- (812°), and p-(808°) compounds are formed by the action of cone. H₂SO₄ on a mixture of styrene and o-, m-, and p-xylene respectively (Kraemer, Spilker,

They are oils, miscible with alcohol and ether.

PHENYL-TOLYL-PROPIONIC ACID

CH_Ph.OH(O_H_Me).CO_H. o-[95.5°], m-[80°], Formed from the corresponding **p**- [105°]. nitriles, which are got from the tolyl-acetonitriles by treatment with NaOEt and benzyl chloride (Päpcke, B. 21, 1331). The o-nitrile is an oil $(840^{\circ}-858^{\circ})$; the *m*-nitrile $[58^{\circ}]$ $(350^{\circ}-860^{\circ})$, and the *p*-nitrile $[79^{\circ}]$ are crystalline.

Di-phenyl-p-tolyl-propionic acid. Nitrile CH₂Ph.CPh(C,H,Me).CN. [121°]. Formed from phenyl-p-tolyl-acetonitrile, NaOEt, and benzyl chloride (Neure, A. 250, 150). Needles.

NO,H, CPh:CH. The o- [115°] (above 300°) and p- [203°] compounds are formed by distilling the corresponding carboxylic acids with lime (Baumann, B. 20, 1492; Paal a. Braikoff, 22,

Both crystallise in white needles

DI-PHENYL-TOLYL-PYRROLE C2.H1.N i.e.

DI-PHENYL-TOLYL-PYRROLE CARBOXY-LIC ACID N(O,H,) < CPh:C.CO,H. The o-acid [227°] and its p- isomeride [206°] are got by saponification of the ethers, which melt at 135° and 145° respectively, and are formed by boiling the corresponding toluidine with phenacyl-benzoyl-acetic ether CH₂Bz.CHBz.CO₂Et and HOAc (Paal a. Braikoff, B. 22, 3088). Both acids are crystalline and sl. sol. ether.

Di-phenyl-p-tolyl-pyrrole dicarboxylic acid $N(C_1H_1) < C(C_2H_4,CO_2H):CH \\ C(C_4H_4,CO_2H):CH \\ C(C_4H_4,CO_2H):CH \\ C(C_4H_4,CO_2H):CH \\ C(C_4H_4,CO_2H):CH \\ C(C_4H_4,CO_2H):CH \\ C(C_4H_4,CO_2H_4) \\ C($ heating C₂H₄(CO.C₂H₄.CO₂H)₂ with p-toluidine on a water-bath (Baumann, B. 20, 1489).

Yellowish needles, v. sol. alcohol and CS₂.

PHENYL p-TOLYL SULPHIDE Ph.S.C,H,. An oil got by the action of NaSPh on p-diazotoluene chloride (Ziegler, B. 23, 2471).

Phenyl p-tolyl disulphide Ph.S2.C,H,. Formed by adding Br to an ethereal solution of PhSH and C,H,SH. Got also, together with Ph₂S₂, by heating toluene p-sulphinic acid with PhSH (Otto a. Rössing, B. 19, 8133).

oil, insol. water, scarcely volatile with steam.

PHENYL p-TOLYL SULPHONE C₁₃H₁₂SO₂
i.e. C₄H₄.SO₂.C₄H₄Me. [125°]. S. (alcohol) 1·62 at
20°. Formed by the action of P₂O₃ on a mixture
of toluene and benzene sulphonic acid, or of benzene and toluene p-sulphonic acid (Michael a. Adair, B. 11, 116). Formed also from benzene sulphonic chioride, toluene, and AlCl, (Beckurts a. Otto, B. 11, 2068). Prisms or tablets. PHENYL-o-TOLYL-SEMI-THIOCARBAZIDE

 $C_{14}H_{15}N_{2}S$ i.s. NHPh.OS.NH.NH. $C_{6}H_{4}Me$. [146°]. Formed from phenyl-thiocarbimide and o-tolyl-hydrazine (A. E. Dixon, C. J. 57, 260). Pearly prisms, v. sl. sol. water, v. sol. hot alcohol.

Isomeride C.H.MeNH.CS.NH.NHPh. [163°] Formed from o-tolyl-thiocarbimide and phenyl-

hydrazine. Vitreous prisms, v. sl. sol. water. PHENYL o-TOLYL-THIO-UREA C₁₄H₁₄N₂S i.e. NHPh.CS.NHC.H.Me. [140°]. Formed from o-tolyl-thiocarbimide and aniline (Staats, B. 13, 137) and by passing H2S into a boiling solution of phenyl-tolyl-cyanamide NPh:C:NC,H, in dry benzene (Huhn, B. 19, 2411). Long needles, sl. sol. water. Boiling HClAq (34 p.c.) splits it up into aniline, o-toluidine, phenyl-thiocarbimide, and o-tolyl-thiocarbimide (Mainzer, B. 15, 1419).

HgO, added to its solution in boiling benzene,

forms phenyl-tolyl-cyanamide.

Phenyl-p-tolyl-thio-urea NHPh.CS.NHC₆H₄Me. [137°] (S.); [141°] (F. a. W.). Formed from p-tolyl-thiocarbimide and aniline (S.), and from phenyl-p-tolyl-cyanamide and H₂S (H.). Plates, sl. sol. water. Split up by HClAq in the same way as the o-isomeride (M.), and decomposed by HgO in like manner. COCl₂ passed into its solution in toluene forms NPh:C $\stackrel{N}{\subset}$ $\stackrel{N(C_7H_7)}{\longrightarrow}$ CO [89°], crystallising in white needles (Freund a. Wolf, B. 25, 1466).

Tetra-ω-phenyl-di-o-tolyl-thio-urea C₂₀H₂₂N₂S i.e. (CHPh₂·C₆H₄·NH)₂CS. [123°]. Formed from CHPh₂·C₆H₄·NH₂ and CS₂ (Fischer a. Fränckel, A. 241, 368). White needles, insol. cold alcohol.

References.—Nitro- and Oxy- Phenyl-tolyl-

THIO-UREA

PHENYL - p - TOLYL - THIO - UREA-ω-CARB-OXYLIC ACID. Nitrile

NHPh.Cs.NH.C₈H₄.CH₂.CN. [141°]. Formed by warming phenyl-thiocarbimide with amidophenyl-acetonitrile (Freund a. Immerwahr, B. 23, 2856). With furfuraldehyde it forms NHPh.Cs.NH.C₈H₄.CCy:CH.C₄H₂O [160°] crystallising from alcohol.

PHENYL-o-TOLYL-UREA C₁₄H₁₄N₂O *i.e.* NHPh.CO.NHC₆H₄Me. [212°]. Formed by boiling NPh:C:NC₇H, with dilute alcohol (Huhn,

B. 19, 2410). Slender needles.

Phenyl-m-tolyl-urea. [165°]. Formed from phenyl cyanate and m-toluidine in ether (Buchka a. Schachtebeck, B. 22, 840). Needles.

Phenyl-p-tolyl-urea. [211°] (H.); [218°] (Freund a. Wolf, B. 25, 1467). Formed by boiling phenyl-p-tolyl-cyanamide with dilute alcohol (Huhn, B. 19, 2408).

Phenyl-di-p-tolyl-urea C₂₁H₂₀N₂O. [136°]. Formed from (C₅H₄Me)₂N.COCl and aniline (Hammerich, B. 25, 1821). Needles, sol. alcohol.

Di-phenyl-p-tolyl-urea NPh₂.CO.NHC,H₁, [180°]. Formed from NPh₂.COCl and p-toluidine

(Michler, B. 9, 713). Needles.

References.—OXY-PHENYL-TOLYL-UREA.

(β)-PHENYL-UMBELLIFERONE $C_{1a}H_{1o}O_{3}$ i.e. $\begin{bmatrix} 4\frac{1}{2} \end{bmatrix} C_{a}H_{1o}(O) + CO - CO$. [244°]. Prepared by the action of $H_{2}SO_{4}$ on a mixture of benzoylactic ether and resorcin (Pechmann a. Duisberg, B, 16, 2126). Plates (from dilute alcohol). PHENYL - URAMIDO - ACETO - NITRILE

PHENYL - URAMIDO - ACETO - NITRILE C. H. OH(CN).NH.CO.NH. Phenyl-aceto-nitrile-urea. [178°] with decomposition. Formed by heating equimolecular quantities of benzaldehyde-cyanhydrin and urea at 100° (Pinner a. Lifschütz, B. 20, 2355). Prisms. V. sol. alcohol, m. sol. hot water. By boiling with dilute HCl it is converted into di-oxyaland heat at the converted into di-

phenyl-glyoxaline CHPh.N C.OH.

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PHENYL-a-URAMIDO-CROTO-NITRILE C_zU_z.CH:CH.C(CN)H.NH.CO.NH_z. Phenyl-croto-nitrile-urea, [160° with decomposition]. Prepared by heating equivalent quantities of cinnamicaldehyde-cyanhydrin and urea to 96° for several hours; the yield is 40 p.c. of the cyanhydrin (Pinner a. Lifschütz, B. 20, 2353). Needles.

M. sol. hot alcohol. By boiling with dilute HCl it is converted into di-oxy-styryl-glyoxaline CO < NH.CH.CH:CHPh.

PHENYL - URAMIDO - PHENYL - ACETIC ACID NHPh.CO.NH.CHPh.CO.H. [154°]. Formed by the action of potash on di-phenylthiohydantoin (Kossel, B. 24, 4153). Sol. water. Ethyl ether Eth'. [165°]. Formed from

Ethyl ether EtA'. [165°]. Formed from phenyl-amido-acetic ether and phenyl cyanate (K.). Crystalline powder, sol, hot alcohol

(K.). Crystalline powder, sol. hot alcohol.

PHENYL-URAMIDO-PROPIONIC ACID

(1₁CH₁N₂O₃ i.e. C₂H₅NH.CO.NH.CH₂.CH₂CQ-H.

(172°)]. Formed by heating β-amido-propionic acid with phenyl-urea at 140° (Hoogewerff a. van Dorp, R. T. C. 9, 49). Tablets and needles, m. sol. cold alcohol. AcCl forms C₁₀H₁₀N₂O₂ (234°) and C₁₀H₂AcN₂O₂ [138°]. KOBr and HCl form C₁₀H₁₀Br₂N₂O₃ [202°] and C₁₀H₂Br₃N₂O₃ [220°].—KA'.—CaA'₂.—AgA'.

Ethyl ether EtA'. [85°]. Needles.

DI-PHENYL-URAZURE

Ethyl ether ÉtA'. [85°]. Needles.
DI-PHENYL-URAZINE C₁₄H₁₂N₄O₂ i.s.

NPh\(^{O}_{O}.NH\) NPh. [264°]. Formed by heating phenyl semicarbazide at 160° (Pinner, B. 21, 2329) or phenyl-carbazic ether at 240° (Heller, A. 263, 282). White needles, v. sol. warm HOAC. Yields the acetyl derivatives C₁₄H₁₁AcN₄O₂ [173°], C₁₄H₁₀Ac₂N₄O₂ [153°] and the ether C₁₄H₁₁EtN₄O₂ [137°].

PHENYĽ-ÚRAZÓĽE v. ĎI-OXY-PHENYL-TRI-AZOLE.

PHENYL-UREA C,H₈N₂O i.e. NHPh.CO.NH₂. [147°]. Formation.—1. By passing cyanic acid vapour into cooled aniline or from aniline sulphate and potassium cyanate (Hofmann, A. 53, 57; 57, 265; 70, 130; 74, 14; Weith, B. 9, 810). 2. From aniline and moist cyanogen chloride.—3. From phenyl cyanate and NH₃.—4. From mercuric fulminate and aniline (Steiner, B. 8, 518). 5. From benzamidoxim, NaOH, and benzene sulphonic chloride (Pinnow, B. 24, 4171).

Properties.—Monoclinic needles, v. sol. alcohol and boiling water, sol. ether. Decomposed at 150° into s-di-phenyl-urea, CO₂, and NH₃.

Reactions.—1. Conc. KOHAq yields aniline, NH_s, and CO₂.—2. Baryta-water yields phenylbiguanide (Emich, M. 12, 16).—3. Aniline at 190° forms s-di-phenyl-urea.—4. Fuming sulphuric acid forms [1:4] C₆H₄(NH₂).SO₂H and SO₂H.C₆H₄,NH.CO₂H (Hentschel, B. 18, 978).—5. ClCO.CO₂Et forms phenyl-allophanic ether and phenyl-parabanic acid.

Acetyl derivative NHPh.CO.NHAc. [183°]. Needles (McCreath, B. 8, 1181; Kühn,

B. 17, 2880; Pinnow, B. 24, 4171).

Propionyl derivative. [137°]. Prisms.

Benzoyl derivative NHPh.CO.NHBz.
199°]. Silky needles, v. sol. alcohol.

s-Di-phenyl-ures $C_{12}H_{12}N_2O$, i.e. $CO(NHPh)_2$. Mol. w. 212. Carbanilide. [235°]. (260°).

Formation.—1. From aniline and phenylcyanate (Hofmann, A. 57, 266; 74, 15).—2. From
aniline and COCl₂ (Hofmann, A. 70, 138; Hentschel, J. pr. [2] 27, 499).—3. By heating diphenyl-thio-urea with alcoholic potash.—4. By
distilling phenyl-urea (H.; Pinnow, B. 24, 4172).
5. By dry distillation of aniline oxalate (Hofmann, Pr. 15, 335).—6. By heating urea (1 pt.)
with aniline (8 pts.) at 160° (Baeyer, A. 131, 251)
or phenyl-urea (1 mol.) with aniline (1 mol.) at

185° (Weith, B. 9, 821).—7. Together with NH₃, benzamide, and benzamilde, by heating dibenzoyl-urea with aniline at 180° (Holleman, R. T. C. 10, 72).—8. By the action of an ethereal solution of ClCO₂CCl, on aniline (Hentschel, J. pr. [2] 36, 310).—9. From NHPh.CO₂Et and aniline at 160° or NaOPh at 220°.—10. From CO(OPh)2 and aniline at 150°-180° (Eckenroth, B. 18, 516). 11. From C(OEt), and aniline at 280° (Bender, B. 13, 699).—12. By heating carbamic ether with aniline (3 mols.) at 180°-185° (Smolka, M. 11, 200).—13. From benzamidine hydrochloride, phenyl cyanate, and NaOHAq (Pinner, B. 22, 1607).

Properties.—Prisms (from alcohol), v. sl. sol. water, v. sol. alcohol and ether. May be distilled.

Reactions .- 1. Conc. H2SO4 forms CO2 and C₆H₄(NH₂)SO₃H.—2. Alcoholic NH₃ forms urea and aniline (Claus, B. 9, 693).—3. PCl₃ forms phenyl cyanate (Weith, B. 9, 810). ClCO2CCl3 also forms phenyl cyanate on heating.-4. P2Os forms phenyl cyanate and aniline.-5. Dry NaOEt at 220° forms aniline and tri-phenylguanidine. - 6. ClCO.CO.Lt forms di-phenylparabanic acid.

Acetyl derivative NHPh.NPhAc. [115°]. Laminæ (from water) (McCreath, B. 8, 1181). u-Di-phenyl-urea NPh₂.CO.NH₂. [189

[189°]. Formed from NPh2.COCl [85°] and alcoholic NH3 at 100° (Michler, B. 8, 1665; 9, 396, 715). Needles. Gives a blue colour with H_2SO_4 . Split up by distillation into cyanic acid and diphenylamine.

Tri-phenyl-urea NPh2.CO.NHPh. [136°]. Formed by heating a solution of NPh₂ COCL dissolved in chloroform, with aniline at 130° (Michler, B. 9, 396, 715). White needles. Resolved by heat into diphenylamine and phenyl

Tetra-phenyl-urea $CO(NPh_2)_2$ [183°]. Formed by heating NPh_2 -COCl with NPh_2 H and zinc-dust (Michler, B. 12, 1166). Got also from diphenylamine and $COCl_2$ (Grard a. Willim, BL[2] 25, 248). Resolved by HClAq at 250° into diphenylamine and CO2.

References. — AMIDO-, BROMO-, CHLORO-, CHLORO-, NITRO- and OXY- PHENYL-UREA.

PHENYL-UREA m-CARBOXYLIC ACID

NHPh.CO.NH.C₆H₄.CO₂H. [270°]. Formed by heating m-amido-benzoic acid with phenyl cyanate at 100° (Kühn, B. 17, 2882). Prisms, sol. alcohol, sl. sol. ether.

Di-phenyl-urea di-m-carboxylic acid $\mathbf{C}_{15}\mathbf{H}_{12}\mathbf{N}_{2}\mathbf{O}_{5}$, i.e. $\mathbf{CO}(\mathbf{C}_{6}\mathbf{H}_{4}.\mathbf{NH}.\mathbf{CO}_{2}\mathbf{H})_{2}$.

Formation.—1. By heating m-uramido-benzoic acid (Griess, Z. 1868, 650; B. 9, 796).—2. By heating a mixture of m-uramido-benzoic acid with m-amido-benzoic acid at 175° (Traube, B. 15, 2122).—3. From m-amido-benzoic acid and COCl₂ (Sarauw, B. 15, 44). — 4. By boiling CS(C₂H₄.NH.CO₂H), with HgO and KOHAq (Griess, A. 172, 169).—5. By heating urea with m-amido-benzoic acid at 130° (T.).

Properties.—Minute needles, almost insol. water, alcohol, and ether.

Salts.—BaA" Saq.—PbA".—Ag₂A".

Ethyl ether Et₂A". [162°]. Needles.

Amide CO(C₆H₄.NH.CO.NH₂)₂. Got by heat-Ing urea with m-amido-benzamide at 140° (Schiff, A. 232, 140). White powder, insol. water, decomposing above 270°.

Di-phenyl-urea di-p-carboxylic acid. Formed when urea is heated with ρ -amido-benzoic acid (Griess, J. pr. [2] 5, 370). Small needles.—BaA".

TETRA - PHENYL - UVINONE C₃₄H₂₀O₄.

Formed in small quantity in the preparation of di-phenyl-furfurane by heating di-phenyl-furfurane dicarboxylic acid (Perkin a. Schlosser, C. J. 57, 956). Thin yellow needles, not melting at 280°. H2SO, forms a dark-green solution with brick-red fluorescence.

DI-PHENYL-VALERAMIDINE C₁₇H₂₀N₂, i.e. C₄H_p.C(NPh).NHPh. [111°]. Formed by heating isovaleric acid with aniline and PCl_s at 150° (Hofmann, J. 1865, 416). Crystalline, nearly insol, water.

β-PHENYL-n-VALERIC ACID

CHPhPr.CO₂H. [52°]. Formed by heating its nitrile with HClAq at 185° (Rossolymo, B. 22, 1235). Needles.

Nitrile CHPhPr.CN. (261°). Formed from phenyl-acetonitrile, PrI, and NaOH. Oil.

 β - Phenyl - iso - valeric acid $C_{11}H_{14}O_2$ i.s. CH2Ph.CHEt.CO2H. (272°). Formed by reduction of phenyl-angelic acid with sodium-amalgam Baeyer a. Jackson, B. 13, 118). Formed also by heating benzyl-ethyl-acetoacetic ether with conc. KOHAq (Anschütz, A. 261, 306). Oil.—AgA'.

Chloride (c. 147° at 24 mm.). Oil. Amide C₁₇H₁₈NO. [89°]. Mond Monoclinic . crystals; a:b:c = .817:1:1.189; $\beta = 70^{\circ} 28'$.

 γ -Phenyl-valeric acid CH₂Ph.CHMe.CH₂.CO₂H. (176.5° at 15 mm.). Got by potash-fusion from 'diethyl carbo-benzonic' acid, which is a product of the action of alcoholic potash on deoxybenzoin (Anschütz, A. 261, 302). Oil, yielding benzoic acid on oxidation.

Ethyl ether C₁₃H₁₈O₃. (146° at 15 mm.). Chloride C₁₁H₁₃OCl. (131° at 11 mm.). Anilide C17H19NO. [102°]. Trimetrio needles; a:b:c = 685:1:608.

δ-Phenyl-valeric acid

CH₂Ph.CH₂.CH₂.CH₂.CO₂H. [59°]. Formed by reducing CHPh:CH.CH₂.CH₂.CO₂H (Baeyer a. Jackson, B. 13, 122). Leaflets, sl. sol. water.

(330°-340°). S.G. 174 Benzyl ether. 1.027. Got by heating benzyl butyrate with Na at 130° (Conrad a. Hodgkinson, A. 193, 318). Liquid.

Isomeride v. Benzyl-Isobutyric ACID.

Reference.—Amido-, Di-Bromo-, Di-Bromo-AMIDO-, NITRO- and OXY- PHENYL-VALERIC ACIDS.

PHENYL-VINYL- v. STYRYL-

DI-PHENYL-VINYL-DIAMINE $C_{14}H_{11}N_2$ i.e. NHPh.CH2.CH:NPh. [105°]. Formed by heating CH₂Cl.CH:NPh with aniline (Berlinerblau, M. 8, 187). Sol. alcohol and ether.

PHENYL VINYL KETONE CARBOXYLIC ACID v. BENZOYL-ACRYLIC ACID.

DI-PHENYL-VINYL NITRITE C14H11NO2 i.e. CPh₂:CH.NO₂. [87°]. Formed by gradually adding HNO₂(1 pt.) to a hot solution of di-phenyl-ethane (1 pt.) in HOAc (10 pts.) (Anschütz a. Romig, A. 233, 327). Needles, v. sol. alcohol and ether.

TRI - PHENYL VINYL TRISULPHONE C₆H₅.SO₂.CH₂.CH(SO₂C₆H₅)₂. [86°]. Formed by oxidising the product of the action of NaSPh on CHCl₂.CH₂Cl (Otto, B. 24, 1885). Crystalline. PHENYL-XYLENE v. DI-METHYL-DIPHENYL

and PHENYL-TOLYL-METHANE.

PHENYL-XYLIDINE C14H15N i.e. $C_0H_3M_{\frac{3}{2}}$.NHPh. [52°]. (173° at 15 mm.). Formed by heating xylidine with aniline hydrochloride (Girard a. Vogt, Bl. [2] 18, 67). Sol. alcohol.

PHENYL-0-XYLYL-CARBINOL C15H16O i.e. C_eH_b,CH(OH).C_eH₃Me₂[1:3:4]? [68°]. (336° i.V.) at 744 mm. V.D. 107.7 (for 106). Formed from the ketone, KOH, and zinc-dust (Elbs, J. pr. [2] 35, 469). White radiating needles (from alcohol). Somewhat decomposed by distillation.

Phenyl-m-xylyl-carbinol $C_eH_s.CH(OH).\check{C_eH_gMe_g}[1:2\cdot4]?$ [57°]. (331° i.V.) at 744 mm. V.D. 102·7 (for 106). Formed from the corresponding ketone by reduction (Elbs,

J. pr. [2] 35, 472).

Phenyl-p-xylyl-carbinol C₆H₅.CH(OH).C₆H₈Me₂[1:2:5]. [88°]. Formed from the ketone, KOH, and zinc-dust (Elbs). Prisms (from alcohol); v. sl. sol. water, m. sol.

HOAc, v. e. sol. alcohol and ether.

PHENYL-p-XYLYL-p-CYMYL-METHANE [2:5:1] Me₂C₆H₃.CHPh.C₆H₃MePr[1:2:5]. Formed from phenyl-p-cymyl-carbinol, p-xylene, and P2Os (Elbs, J. pr. [2] 35,498). Oil.

DI - PHENYL - o - XYLYLENE - DIAMINE $C_{20}H_{20}N_2$ i.e. $C_6H_4(CH_2.NHPh)_2$. [172°]. Formed by boiling di-ω-bromo-o-xylene with an alcoholic solution of aniline (Lesser, B. 17, 1825). Small plates (from alcohol).

PHENYL O-XYLYL KETONE C15H14O i.e. C₆H₅.CO.C₆H₃Me₂ [1:3:4]? [48°]. (340° i.V.) at 744 mm. V.D. 102·2 (for 105). Formed from BzCl, o-xylene, and AlCl,; the yield is 80 p.c. (Elbs, J. pr. [2] 35, 467). Groups of snowy needles (from alcohol). Insol. water, sl. sol. cold HOAc, v. sol. alcohol.

Phenyl m-xylyl ketone C_6H_5 .CO. C_6H_3 Me₂ [1:2:4]. (321° i.V.) at 744 mm. V.D. 102·3 (for 105). From m-xylene, BzCl, and AlCl, (Söllscher, B. 15, 1682; Elbs, J. pr. [2] 35, 469). Oil. Partly converted by long boiling into (B. 2)-methylanthraquinone. In presence of a little of the corresponding phenyl - xylyl - carbinol, (B. 2)-

methyl-anthracene is formed.

anti-Oxim C₈H₉.C.C₈H₅.N.OH. $[126^{\circ}].$ verted by PCl_s into C_sH_g.CO.NHPh (Smith, B. 24, 4048). Yields an acetyl derivative [91°], crystallising in flat prisms.

 $syn\text{-}Oxim \stackrel{C_8H_9.C.C_6H_5}{\text{HO.N}}$. [152°]. Formed, as well as the preceding body, by the action of an alcoholic solution of hydroxylamine on the ketone. PCl, at -20° forms C, H, CO.NHC, H,

Yields an acetyl derivative [103°].

Phenyl - p - xylyl ketone C_aH_a.CO.C_aH_aMe₂
[1:2:5]. [36°]. (317° i.V.) at 744 mm. V.D.
106'2 (for 105). From p-xylene, AlCl_a, and BzCl (Elbs, B. 17, 2847; J. pr. [2] 35, 472); the yield is 65 p.c. Transparent prisms (from alcohol), insol. water, m. sol. HOAc, v. e. sol. ether and alcohol. Very slightly volatile with steam. Yields methyl-anthracene when boiled for a long time.

Reaction.—1. H₂SO₄ has no action in the cold, but on warming HOBz is split off.—2. H₂S₂O₇ forms a disulphonic acid $C_6H_3Me_2.CO.C_6H_3(SO_3H)_2$, whose salt BaA"2aq is v. e. sol. water. -8. HNO₃ (S.G. 1·15) at 180° dicarboxylic forms benzophenone acid $C_{e}H_{s}.OU.C_{s}H_{3}(CO_{2}H)_{2}$ [1:2:5].

PHENYL-0-XYLYL-KETONE o-CARB OXYLIC ACID

[4:3:1] C₆H₃Me₂.CO.C₆H₄.CO₂H₄ [2:1]. Xylene-phthaloylic acid. [162°]. Formed by the action of AlCl, on a mixture of phthalic anhydride and o-xylene (F. Meyer, B. 15, 636). Minute prisms (containing aq). Yields benzoic and (4,8,1)-dimethyl-benzoic acids when fused with potash.

Phenyl-m-xylyl-ketone o-carboxylic acid [4:2:1] $C_bH_bMe_2$. $CO_cG_bH_b$. Formed in like manner from m-xylene (M.). Needles, sl. sol.

water, sol. alcohol.

Phenyl-p-xylyl-ketone o-carboxylic [5:2:1] $C_0H_3Me_2.CO.C_0H_4.CO_2H$ [1:2]. Formed in like manner from p-xylene. Amorphous solid, insol. water, sol. alcohol and benzene.

Phenyl-m-xylyl-ketone (a)-carboxylic acid $C_6H_5.CO.C_6H_2Me_2.CO_2H$ [4:5:3:1]. [160°]. Formed, together with the following isomeride, by oxidising benzoyl-mesitylene with chromic acid mixture (Louise, Bl. [2] 44, 418; A. Ch. [6] 6, 218). Needles, v. sol. ether, insol. cold water.—BaA', 2aq.—MgA', 6aq.—AgA': needles.

Phenyl-m-xylyl-ketone (B)-carboxylic acid C₆H₅.CO.C₆H₂Me₂.CO₂H_[2:5:3:1]. [185°]. Formed as above. The Mg salt is more soluble than that of the isomeric acid. Efflorescent needles, v. sl. sol. hot water. Converted by P2O5 into di-methyl-anthraquinone [158°].—AgA': needles.

dicarboxylic Phenyl-xylyl-ketone C.H. CO.C. HMe₂(CO₂H)₂. Benzoyl-cumidic acid. [85°]. Formed by oxidation of phenyl duryl ketone (F. Meyer a. Ador, J. 1879, 562). Melts at 85°, becomes solid, and melts again at 173°.

-BaA'. $2\frac{1}{2}$ aq : long silky needles. PHENYL - DI - p - XYLYL - METHANE $C_6H_5.CH(C_6H_3Me_2)_2[1:2:5]_2$. [93°]. (above 360°). From phenyl p-xylyl carbinol and p-xylene (Elbs, J. pr. [2] 35, 476). Or from di-p-xylyl carbinol, beazene, and P₂O₅. Prisms (from ligroin). Its solutions show blue fluorescence.

Di-phenyl-o-xylyl-methane $C_{24}H_{20}$ i.e. $(C_8H_5)CH.C_8H_3Me_2[1:3:4].$ [68.5°]. (above 360°). Formed from di-phenyl-carbinol, o-xylene, and P_2O_3 (Hemilian, B. 19, 3070). Needles, v. sol.

alcohol and ether.

 ${f Di-phenyl-}m$ - ${f xylyl-methane}$ C_eH₅.CH₂.C_eH₈Me₂[1:2:4]. [61·5°]. (above 360°). Formed by boiling di-phenyl-carbinol with m-xylene and P₂O₅ (Hemilian, B. 19, 3061). Prisms, v. sol. alcohol and ether. Oxidised by chromic acid mixture to methyl-di-phenyl-

phthalide and di-phenyl-phthalide carboxylic Di-phenyl-p-xylyl-methane. [92°]. (above

360°). Formed by digesting di-phenyl-carbinol with p-xylene and P2O5 (Hemilian, B. 16, 2360; Bl. [2] 34, 326; Petrieff, Bl. [2] 41, 816). Monoclinic crystals, v. sol. alcohol and ether. Oxidised by chromic acid mixture to di-phenylmethyl-phthalide [179°], di-phenyl-tolyl-carbinol m-carboxylic acid [c. 253°] and di-phenyl-

Reference.—NITRO-DI-AMIDO-PHENYL-DI-XYLYL-

METHANE. o-CARB-PHENYL-DI-XYLYL-METHANE OXYLIC ACID [4:2:1] C₆H₂Me₂.CH₂.C₆H₄.CO₂H. [158°]. Formed by reducing phenyl-m-xylylketone carboxylic acid with zinc-dust and NH,Aq (Greely, A. 234, 237). Small needles (from alcohol).-BaA'aq: plates (from dilute alcohol).

phthalide carboxylic acid [245°].

PHENYL-p-XYLYL-(β)-PINACOLIN C_{so}H_{ss}O 4.6. C.H. C(O.H.Me.), CO.C.H. [146°]. Formed by the action of zinc and HClAq on phenyl p-xylyl ketone (Elbs, J. pr. [2] 35, 477). Clumps of prisms (from ligroin), m. sol. alcohol and ether. Soda-lime at 320° splits it up into benzoic acid and phenyl-di-xylyl-methane.

PHENYL-XYLYL-PROPANE C17H20 ie. C.H. CHMe.CH. C.H. Me. (324°). Formed from ψ-cumene, styrene, and conc. H₂SO₄ (Kraemer,

Spilker a. Ebenhardt, A. 23, 3273).
PHENYL-XYLYL-PROPIONIC ACID C.H.Mez.CHPh.CHz.COzH. Formed from cinnamic acid, m-xylene, and H2SO4 (Liebermann s. Hartmann, B. 25, 959). Amorphous, v. sol. warm benzene. When allocinnamic acid is used it is accompanied by another acid [220°].

DI - PHÉNYL - XYLYL - PYRRÖLE OXYLIC ACID O.H.N CPh:C.CO.H. [254°].

Formed by saponifying its oily ether, which is got from phenacyl-benzoyl-acetic ether and (4,2,1)-m-xylidine (Paal a. Braikoff, B. 22, 3090). Small needles, m. sol. hot alcohol and benzene.

PHENYL XYLYL SULPHONE C14H14SO2, i.e. C₆H₅·SO₂·C₆H₂Me₂. [80°]. Formed from benzene sulphonic chloride, *m*-xylene, and AlCl₃ (Beckurts

a. Otto, B. 11, 2069). Yellowish needles. PHENYL m-XYLYL-UREA C₁₅H₁₆N₂O i.e. C.H.NH.CO.NHPh. [131°]. Formed by mixing m-xylidine with phenyl cyanate (Brömme, B. 21, 2703). White matted needles, sol. alcohol.

PHENYTHRONIC ACID v. PHENYL-METHYL-

FURFURANE DICARBOXYLIC ACID.

PHILLYRIN C₂₇H₂₄O₁₁ 1½aq. [160°]. S. 08 at 9°; S. (alcohol) 2·5 at 9°. A glucoside in the bark of Phillyrea latifolia (Bertagnini, A.92, 109; 118, 124). Crystalline, insol. ether. Split up by dilute acids into glucose and crystalline phillygenin $C_{21}H_{24}O_{6}$, v. sol. ether. A glucoside $C_{26}H_{22}O_{11}$ ([184°]; S. 05 in the cold, 12·5 at 100°) A glucoside in Olea fragrans is perhaps identical with phillyrin. It is split up by acids into glucose and

 $C_{20}H_{22}O_{e}$ [70°] (Eykman, R. T. C. 5, 127). PHLEIN $C_{30}H_{62}O_{31}$. [215°]. S.G. 1·48. S. 3·26 at 10°. [α]₀ = -48·5. Occurs in the bulbs of cat's-tail grass (Phleum pratense) and in the roots of Baldigera arundinacea (Ekstrand a. Johanson, B. 20, 3310; 21, 597). Carbohydrate resembling starch, but not coloured blue by iodine. Reduces AgNOs, but not Fehling's solu-

PHLOBAPHENE Cas H26O17. Named from φλοιόs, bark, and βαφή, colour. Occurs in oak bark, and formed by boiling quercitannic acid with dilute acids (Hofstetter, A. 51, 63; Grabowski, A. 145, 3; Oser, J. 1876, 903; Böttinger, A. 202, 270; 240, 338). Reddish-brown powder, insol. water, ether, and cold alcohol. Alkalis form a reddish-brown solution, which absorbs oxygen from the air. Turned black by FeCl. Potash-fusion yields protocatechuic acid (Procter, C. J. 36, 979). Yields $C_{ss}H_{10}Ac_{r}O_{11}$, $C_{ss}H_{10}Bz_{r}O_{11}$, and $C_{ss}H_{ss}K_{s}O_{11}$, which is insol. alcohol, v. sol. water. Bromine yields $C_{ss}H_{1s}Br_{1o}O_{17}$, and C₂₈H₂₀Br₆O₁₇, which forms C₂₈H₁₃Br₆Ac₇O₁₇. A substance C₂₆H₄₆O₂₅ resembling phlobaphene occurs in hops (Etti, A. 180, 223; D. P. J. 228,

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PHLORAMINE C.H., NO., i.e.

C₆H₂(OH)₂(NH₂). Amido-resorcin? Formed by passing NH₃ over phloroglucin (Hlasiwetz a. Pfaundler, A. 119, 202). Thin plates (from water), sl. sol. cold water, v. sol. alcohol, insol. ether. Turns brown in air. Decomposed by alkalis. Its solutions are not coloured by FeCl, -B'HCl. $-B'_2H_2SO_12aq: long yellowish needles.$

PHLOREIN C₁₈H₁₁NO₇. Formed by passing nitrous acid gas into an ethereal solution of phloroglucin (4 g.) containing HNO₃ (4 c.c. of S.G. 1.25) (Benedikt, B. 7, 445; A. 178, 93). Lustrous dark-green powder, insol. water, v. sol. alcoholand ether, forming a dark-brown solution. Alkalis form a purple solution. Yields phloro-glucin when fused with potash. Zinc and dilute H₂SO₄ form a colourless body, re-oxidised to phlorein by air.

PHLORETIC ACID v. p-Oxy-a-PHENYL-PRO-PIONIC ACID. It yields a crystalline di-bromo-

derivative (Hlasiwetz, A. 102, 145).

PHLORETIN C₁, H₁₄O₅. [255°]. Formed by boiling phlorizin or glycyphyllin with dilute acids (Stas, A. 30, 200; G. Roser, A. 74, 178; Hlasiwetz, A. 96, 118; Schiff, A. 156, 2; 172, 357; 229, 374; Rennie, C. J. 49, 860). Small laminæ, with sweet taste, v. sl. sol. hot water and ether, v. e. sol. alcohol. Inactive to light. Bromine forms C₁₅H₁₀Br₄O₅ [205°-210°] (Schmidt a. Hesse, A. 119, 103). Alkaline solutions absorb oxygen and turn orange in air; boiling conc. KOHAq splits it up into phloretic acid and phloroglucin. Aniline at 170° forms $C_{21}H_{19}NO_{43}$ a scarlet powder. AcCl gives amorphous $C_{15}H_{12}Ao_2O_5$ aq. Salts.— $C_{15}H_{11}(NH_4)_3O_5$: amorphous. Gives off NH_5 in air.— $C_{30}H_{18}Pb_5O_{10}$ 5aq.— $C_{15}H_{13}AgO_5$:

unstable pp

PHLORIZIN C21H24O10 2aq. Named from φλοιός, bark, and ρίζα, root. [109°] (when anhydrous). S.G. 19 1.43. S. 1 in the cold. $[a]_D = -49$ at 15°. Occurs in the root-bark of the apple, pear, plum, and cherry tree, from which it may be extracted by dilute alcohol (Stas a. De Koninck, A. 15, 75; 30, 193; A. Ch. [2] 69, 367; Mulder, Rev. Scient. 3, 50; Roser, A. 74, 178; Strecker, A. 74, 184; Rennie, C. J. 51, 635). Silky needles with slightly bitter taste, sl. sol. cold water, v. e. sol, hot water and alcohol, insol, ether. Melts at 109°, becomes solid, and melts again at 171°. Lævogyrate. Decomposed by dilute acids into glucose and phloretin. Gives pps. with baryta and MeOH, with lime-water, and with lead subacetate. Air and ammonia form reddish-brown amorphous phlorize $\Omega_{21}H_{30}N_{2}O_{13}$. Gives a violet colour on boiling with $ZnSO_{4}$ and KNO_{2} (Nickel, Fr. 28, 248). FeCl, gives a brownish-red colour. Aniline at 180° forms C₂₃H₂₄N₂O₈, a yellow powder, yielding a mono- and a tri- acetyl derivative.

Acetyl derivatives C21 H23 AcO10 2aq: needles (from water). — $C_{21}H_{21}Ac_3O_{10}$. C21H18Ac5O10 aq: amorphous solid.

Tri-bensoyl derivative. Powder. Isophlorizin $O_{21}H_{24}O_{10}$. [105°]. Occurs in the leaves of the apple-tree (Rochleder, Z. [2]4,741). Silvery needles. Its ammoniacal solution turns brown in air. Its solution is ppd. by lead sub-acetate. Dilute $\mathbf{H}_2\mathbf{SO}_4$ splits it up into glucose and isophloretin $\mathbf{O}_{10}\mathbf{H}_{14}\mathbf{O}_{5}$, which is v. sol. ether, and yields phloroglucin when heated with conc. KOHAq.

PHLORQBROMINE C. Br. HO. [152°]. Formed by the action of excess of bromine on a very dilute aqueous solution of phloroglucin (Benedikt, A. 189, 165; C. J. 34, 499). Dimetric prisms; a:b=1:1·2. Insol. water. Not attacked by potash or HNO₃. Warm alcohol decomposes it, forming penta-bromo-acetone [76°]. NH₃Aq forms CBr₃H and C₅Br₅H₄N₂, crystallising from water in colourless laminæ [124°].

PHLOROGLUCIN C.H.O. i.s. C.H.(OH), [1:8:5]. Mol. w. 126. [219°] (Baeyer, B. 19, 2186). H.F. 153,348 (Stohmann, J. pr. [2] 83, 471).

Formation.—1. By boiling phloretin with conc. KOHAq (Hlasiwetz, A. 96, 118).—2. By potash-fusion from quercetin, maclurin, catechin, scoparin, gamboge, dragon's blood, limettin, and bergaptene (Hlasiwetz, A. 112, 96; 119, 199; 127, 357; 134, 118, 283; 138, 190; Zwenger, A. 123, 154; Gautier, Bl. [2] 33, 583; Tilden a. Beck, C. J. 57, 323; Pomeranz, M. 12, 387).—3. By fusing resoroin with a large excess of NaOH, the yield being 65 p.c. (Barth a. Schreder, B. 12, 503).—4. By soda-fusion from phenol, benzene, trisulphonic acid, orcin, and naringenin (Barth a. Schreder, B. 12, 422; M. 3, 649; Will, B. 20, 297).—5. By fusing its tricarboxylic ether with potash (Baeyer, B. 19, 3458).—6. By fusing s-di-bromo-phenol with potash (Blau, M. 7, 632).

Preparation.—By soda-fusion from resorcin (Tiemann a. Will, B. 14, 954; 18, 1323).

Properties.—Trimetric crystals (containing 2aq); a:b:c=825:1:3·417. Melts at 200°-209° when slowly heated. V. sol. water, alcohol, and ether; sl. sol. NaClAq. Tastes sweeter than sugar. May be sublimed. Its solution is ppd. by lead subacetate. FeCl_s gives a bluish-violet colour. Reduces Fehling's solution and ammoniacal AgNO_s. Its alkaline solution absorbs oxygen and turns brown in air. Colours acidified pine-wood red. An alcoholic solution gives a red colour with HClAq and vanillin (Lindt, Fr. 26, 260), cil of cloves, or oil of pimento (Ihl, Chem. Zeit. 18, 264).

Chem. Zeit. 13, 264). Reactions.—1. Bromine forms crystalline tribromo-phloroglucin and finally phlorobromin.

2. Chlorine passed into its aqueous solution forms tri-chloro-phloroglucin C₆Cl₃(OH), 3aq [136°], which soon decomposes into tetra-chloroacetone hydrate, and di-chloro-acetic acid (Webster, C. J. 47, 423; Zincke a. Kegel, B. 22, 1476). Chlorine passed into its solution in HOAc forms CHCl₂.CO.CHCl₂4aq [49°]. Chlorine passed into a cooled solution of dry phloroglucin in chloroform forms CO CCl₂·CO CCl₂ [48°], (269°), decomposed by water into CO2, di-chloroacetic acid, and CO(CHCl₂)₂.—3. Dilute HNO₃ forms nitro-phloroglucin. -4. Ammonia produces phloramine. -5. Nitrous acid in its ethereal solution forms phlorein. In acetic acid solution it gives tri-nitroso-phloroglucin (vol. iii. p. 619) .-6. Aqueous HI at 140° forms nearly tasteless scales of phloroglucide $C_{12}H_{10}O_3$ 2aq, sl. sol. warm water. Phloroglucide is also got by heating phloroglucin alone or with POCl₂.—7. Heated with salicylic acid it forms two compounds of the form $C_eH_4 < CO > C_eH(OH) < CO > C_eH_4$, one of these (described by Kostanecki and Nessler) which vields an acetyl derivative C. H.O. (OAc)

[218°]; the other isomeride [826°] crystallises from HOAc in green-yellow tables.—8. With o-amido-bensoic aldehyle and NaOHAq it forms red C₁₃H₂NO₂, possibly di-oxy-acridine C₆H₂(OH)₂ CH C₆H₄, which forms B'₂H₂PtCl₄ and C₁₃H₃Bz₂NO₂ (Eliasberg a. Friedländer, B. 25, 1752).—9. Phenyl cyanate unites, forming C₆H₄(O.CO.NHPh)₃, a yellowish powder [128°] (Goldschmidta. Meissler, B. 23, 269).—10. Bensone sulphonic chloride added to the slightly alkaline solution forms C₆H₂(O.SO₂C₆H₅)₃ [117°] (Georgesen, B. 24, 418).—11. Aniline at 210° forms triphenyl-tri-amido-benzene [193°] (Minunni, B. 21, 1984).—12. Phloretic acid at 170° forms C₃₃H₃₂O₁₄, crystallising from water in laminæ (Hlasiwetz, A. 119, 199).

Tri-oxim C_cH_s(NOH)_s. Formed from phloroglucinand aqueous hydroxylamine (Baeyer, B. 19, 159). Colourless crystals, v. sl. sol. water and alcohol, sol. alkalis and acids. Explodes at

Phenyl-hydrasine derivatives. The salt C₆H₆O3N₂H₃Ph [78°-83°] is got from phlorogluoin (1 mol.) and phenyl-hydrazine (3 mols.) in alcoholic solution (Baeyer a. Kochendörfer, B. 22, 2190). Nodules, sol. alcohol and ether. Cold NaOHAq sets free phenyl-hydrazine. In alcoholic solution it changes on keeping to C₆H₃(OH)(N₂H₂Ph)₂ [144°], which crystallises from toluene in needles, and yields a pentabenzoyl derivative [176°].

Tri-acetyl derivative C_eH₄(OAc)₄. [106°]. Got by heating phloroglucin with AcCl.

Di-benzoyl derivatives C.H. (OH) (OBz). Two isomerides [165°] and [191°-195°] are got from phlorogluoin with BZCI. The compound [165°] is more sol. benzene than the other (Skraup, M. 10, 391, 722).

Tri-bensoyl derivative C₆H₃(OBz)₃. [174°]. Formed, together with C₁₂H₄(OBz)₄, [199°], by warming phlorogluoin with BzCl and NaOH (S.). Plates, sol. benzene.

Tri-methyl ether C₈H₃(OMe)₃. [52°]. (256°). Formed by passing HCl into a solution of phloroglucin in MeOH, the resulting C₆H₃(OH)(OMe)₂ being treated with MeI and KOH (Will, B. 21, 603). Crystals, insol. water and alkalis, v. sol. alcohol and ether. Conc. HNO₃ forms a blue solution. Bromine gives C₆Br₃(OMe)₃ [145°]. When phloroglucin is treated with KOH and MeI the products are C₆H₃Me₂O₃ [184°], C₆H₂Me₄O₃ [114°], and C₆Me₂O₃ [80°] (Margulies, M. 9, 1052; 10, 459). When tetra-methyl-phloroglucin is heated in sealed tubes with HClAq it yields isopropyl ketone, isobutyric acid, HOAc, and CO₂. Hence it would appear to be CMe₂CO.CMe₂CO.CH (Spitzer, M. 11, 104, 287). Hexamethyl-phloroglucin heated with LIAq at 200° produces isoutyric acid and C,H_{1e}, and, on oxidation, yields di-isopropyl ketone. Hence it would appear to be CMe₂CO.CMe₂

Di-ethyl ether C₀H₃(OEt)₃(OH). [75°]. Made by passing HCl into an alcoholic solution of phloroglucin or its carboxylic acid (Will a. Albrecht, B. 17, 2106). Long white needles. May be distilled.

Tri-ethyl ether C.H. (OEt). [43°]. Got by heating the di-ethyl ether with Etl and alco-

holic potash (W. a. A.).

Penta-ethyl derivative v. vol. ii. p. 505. Tri-phenyl ether C,H3(OPh),? [175°]. A product of the action of Na on phenyl acetate (Hodgkinson, C. J. Proc. 2, 188). V. sl. sol. Aq. Вкомо-рт-горо-, References .- TRI-BROMO-,

TRI-CHLORO-, NITROSO-, and NITRO- PHLOROGLUCIN.
PHLOROGLUCIN CARBOXYLIC ACID v.

TRI-OXY-BENZOIC ACID.

Phloroglucin tricarboxylic ether C15H18O9 i.e. $C_0(OH)_3(CO_2Et)_3$. [106°]. Formed from malonic ether and ZnEt₂ (Lang, B. 19, 2937). Prepared by dissolving Na (14·4 g.) in malonic ether (200 g.) at 100° and heating the product for six hours at 145° (Baeyer, B. 18, 3457; Bally, B. 21, 1766). In this preparation an anhydride $C_6(OH)_2(CO_2Et)_2 < \stackrel{Q}{<} [170^\circ]$ is also formed. Needles (from dilute alcohol), insol. water. Does not react with nitrous acid gas.

Reactions.-1. Potash-fusion yields phloroglucin.—2. Bromine in CS₂ ppts. bromo-phloro-glucin dicarboxylic ether [128°].—3. Chlorine gives tri-chloro-acetamide. 4. Phenyl cyanate and C_sH_s at 200° form C_s(O.CO.NHPh)_s(CO₂Et)_s [195°] (Goldschmidt a. Meissler, B. 23, 270)

Trì-acetyl derivative C₆(OAc)₃(CO₂Et)₃. [76°]. Needles (from alcohol or ether). $Oxim C_6H_s(NOH)_s(CO_2Et)_s$. [171°].

PHLOROGLUCIN-PHTHALEIN C20H12O7. Formed by heating phloroglucin with phthalic anhydride at 170° (Link, B. 13, 1652). orange needles, sl. sol. water. Its alkaline solutions are orange-red, without fluorescence. Zincdust and NaOHAq reduce it to phloroglucinphthalin C20H14O2, an amorphous reddish-yellow

PHLOROGLUCIN SULPHONIC ACID

C₆H₆SO₆ i.e. C₆H₂(OH)₃.SO₃H. Formed by mixing phloroglucin with H2S2O, (Schiff, B. 6, 26; A. 178, 191). Yields a crystalline K salt. POCl. converts it into the anhydrides C12H10S2O11, v. e sol. water, and C12H8S2O10, sl. sol. water, and C24H18S2O15

PHLOROL v. o-ETHYL-PHENOL.

PHLORONE v. XYLOQUINONE

PHORONE C, H₁,O *i.e.* CO(CH:CMe₂)₂. Mol. w. 188. [28°]. (197°) at 743 mm. S.G. $\frac{9}{2}$ ·885. $\mu_D = 1.500$ at 20° (Brühl, A. 235, 15). R_{20} 70.93 in a 14.5 p.c. benzene solution. Formed, together with mesityl oxide, by leaving acetone in contact with quicklime (Fittig, A. 110, 82). Formed also by the action of conc. HClAq on acetone (Baeyer, A. 140, 301), and by heating nitroso-triacetonamine with potash (Heintz, A. 187, 250). Yellowish prisms. Yields acetone, oxalic acid, and CO₂ on oxidation with KMnO₄ (Pinner, B. 15, 591). Slowly combines with NaHSOs, forming di-isobutyl ketone di-sulphonic acid. Yields \u03c4-cumene on heating with P2Os and mesitylene on heating with conc. H2SO4. Yields medityl oxide on distilling with dilute H₂SO₄ (Claisen, A. 180, 18). Bromine in CS, forms CoH14Br4O [88°]. Zinc and H2SO4 reduce it, in alcoholic solution, to deoxyphorone C18H28O [108°]. HI forms C.H., I.O [13°] (Kasaneff, B. 8, 485). Phorone does not react with benzoic aldehyde (Claisen, B. 14, 852).

Oxim C,H₁₄:NOH. [48°]. (218°). Tables, v. sol. alcohol (Nägeli, B. 16, 496).

Isophorone. The substance to which this name has been given is a mixture (Laycock, A. 258, 230).

Camphor-phorone C.H. O i.e.

CH2.CH2 OMe: \mathbb{CPr} (Königs a. Eppens, B. 25, CO

260). (208°) (Kachler, A. 164, 79). Got by distilling calcium camphorate (Laurent, A. Ch. [2] 65, 329; Gerhardt a. Liès-Bodart, A. 72, 293) and by heating camphor (1 pt.) with H₂SO₄ (4 pts.) at 100° (Schwanert, A. 123, 298). Colourless oil with aromatic odour. Inactive to light. PCl_s yields C₀H_{1s}Cl (205°). Yields amethyl-glutaric and acetic acids on oxidation. Na followed by MeI yields $C_9H_{12}MeO$ (225°-230°). Na followed by AcCl yields $C_{18}H_{22}AcO_2$ (230°-240°). Bromine in CS_2 forms $C_9H_{13}Br_3O$ $[52^{\circ}]$

PHORONIC ACID C₁₁H₁₈O₃. [184°]. Formed by saponification of its nitrile, which is got by boiling the product of the action of gaseous HCl on acetone with alcoholic potash (Pinner, B. 14, 1071; 15, 585). Large prisms (from dilute alcohol), sl. sol. water. -KHA" 1 aq: needles, v. sol.

water.—CaA" 3aq: prisms.—Äg,A" aq: pp. $Ethyl\ ether\ Et_2A''$. [125°]. Amide. [above 300°]. Prisms.

Anhydride C, H, 60. [138°]. Got by heating the acid at 190°

Imide C₁₁H₁₈O₃(NH). [205°]. Formed from the anhydride and alcoholic NH,

Nitrile C₁₁H₁₈N₂O₂. [above 320°].

PHOSENE v. SYNANTHRENE.

PHOSGENE GAS. Another name for COCl₂;

v. Carbon Oxychloride, vol. i. p. 692.

PHOSPHAM. (?)PN.H. When dry NH₃ is passed into well-cooled PCl₃, a solid colourless mass is obtained, said by H. Rose to contain PCl_s and NH_s in the ratio PCl_s:5NH_s (P. 24, 308; 28, 529). This substance is probably a mixture of various compounds; other substances are formed if the PCl₃ is not cooled. If PCl₅ is treated with NH, at the ordinary temperature, 5NH₃ are absorbed by PCl₅ (H. Rose, l.c.); from this product ether dissolves out N₃P₃Cl₆ (v. NI-TROGEN PHOSPHOCHLORIDE, vol. iii. p. 570); if the product is heated, HCl and NH₄Ol are evolved, and a white, loose powder remains, to which Gerhardt (A. Ch. [3] 18, 188; 20, 225) gave the composition PN2H and the name phospham. Liebig a. Wöhler (A. 11, 139) obtained this compound by passing PCl₂ or PCl₃ over heated NH₄Cl, and washing and heating the product; they supposed it to be PN₂. The same substance was obtained by Pauli (A. 101, 41) by heating an intimate mixture of P₂S₃ and NH₄Cl, also by heating red P with S and NH₄Cl, and by heating powdered Ca phosphide with S and NH₄Cl, and washing the residue with an acid. Pauli's analyses showed less N than is required by the formula PN₂H. By treating POl, with NH₂, washing for a long time with water, digesting with HClAq, then with KOHAq, washing with water and then with ether, Salzmann (B.7, 494)obtained a substance resembling phospham and approximately agreeing with the composition PsNsH4; Salzmann regarded this substance as probably a mixture of several very similar compounds of P, N, and H. Besson (C. R. 114, 1264) says that PCl₃ absorbs 8HN₃, and that on heating PN₂H is formed.

Phospham is a white, loose powder; insoluble in ordinary solvents; melts and vaporises when heated in absence of air; oxidised by heating in air; yields HPO₂ and HN₃ when moistened with water and heated (Gerhardt, *l.c.*); fused with KOH or BaO₂H₂ forms NH₃ and ortho-

phosphate, with incandescence.

Phospham may be regarded as one of the nitriles of H₃PO₄: thus PO.OH.(ONH₁)₂-4H₂O = PN₂H; the other nitriles would be PON, which is known (v. Phosphorus oxynitride, p. 144) [PO(OH)₂ONH₄-3H₂O = PON], and PN₃H₄, which has not been isolated [PO(ONH₄)₃-4H₂O = PN₃H₄]. Mendelejeff (B. 23, 3472 note) thinks it likely that phospham is a polymeride of PN₂H, inasmuch as it is analogous to N₃H (hydrazoic acid, vol. iii. p. 559) and both P and compounds show a greater readiness to polymerise than N and compounds of N. M. P. M.

PHOSPHAMIC ACIDS AND ALLIED COM-POUNDS. Two smic acids may be derived theoretically from PO(OH), viz. PONH₂(OH)₂ and PO(NH₂)₂OH; both acids have been isolated. The compound PO(NH₂), is described as Phosphamide, and PO.NH.NH₂ as Phosphamido-IMIDE (infra). The two thiophosphamic acids, PS.NH2.(OH)2 and PS(NH2)2OH are probably formed by the action of NH, on PSCI, (v. THIO-PHOSPHORYL CHLORIDE, Reaction 7, p. 148). The three amic acids (or salts of these) derivable from H.P.O, are known: P2O3NH2(OH), P₂O₃(NH₂)₂(OH)₂, and P₂O₃(NH₂)₃OH. By heating P₂O₃(NH₂)₃OH the NH₄ salt of the acid P2O3(OH)N is obtained. There are also several acids (or their salts) known, which may be derived from hypothetical $P_4O_7(OH)_6[2P_2O_3(OH)_4-H_2O]$ $= P_4O_7(\hat{OH})_a]$ by replacing \hat{OH} by \hat{NH}_2 or \hat{NH} . Imidophosphoric acid, PO.NH.OH, perhaps exists.

PHOSPHAMC ACID PO.NH₂(OH)₂. (Amidophosphoric acid.) Obtained by Stokes (Am. 15, 198 [1893]) by suspending the lead salt in a little ice-water, decomposing by excess of H₂S, and filtering into alcohol. White microscopic crystals; easily sol. water; does not give NH₃ with caustic alkalis; heated to 100° is slowly changed to an NH₄ salt, perhaps NH₄.PO₃; aqueous solution soon changes to NH₄.PO₄. Acid and normal salts have been prepared. The lead salt is formed by adding lead acetate to a solution of PO.NH₂.OH.OK, which salt is produced by adding alcoholic NH₃ to an alcoholic solution of PO.Cl(OPh)₂, and saponifying the PO.NH₂(OPh)₂ thus formed by KOHAq.

PHOSPHODIAMIC ACID PO.(NH₂)₂OH. (Diamidophosphoric acid.) Stokes (B. 27, 565 [1894]) prepared this acid by saponifying the product of the action of NH₂Aq on POCl₂(OPh). The acid gives salts PO(NH₂)₂OM, and also salts of an acid P(NH₂)₂(OH)₃ (Abstract in C. J. 66 [11], 188 [1894]).

Pyrophosphamic acid P₂O₃(NH₂)(OH)₃. This is said by Gladstone (C. J. 17, 229) to be the chief product of the action of NH₃ on P₂O₅; Schiff (A. 103, 168) formulated this product as PO.NH.OH, imido-phosphoric acid; Stokes (l.c.) says the product is a mixture. For details v.

G. (l.c.; also C. J. 3, 135; 21 64); also G. a. Holmes (C. J. 17, 225).

Pyrophosphodiamic acm $P_2O_a(NH_2)_2(OH)_2$ (G., C. J. 19, 290; G. a. H., Lc.). Formed by action of water or alkalis on $N_3P_3Ol_a$, by treating POOl₂ with cone. NH_3Aq , and in many other ways. A white amorphous solid, easily soluble in water and alcohol. The acid is dibasic. The salts are difficult to obtain pure, as they easily change into pyrophosphamates; salts of Ba, Ag, and Zn are described.

PYROPHOSPHOTRIAMIC ACID P₂O₃(NH₂)₃OH (G., C. J. 19, 1; 21, 64). Dry NH₂ is passed into POCl₃ until saturated, the product is heated to 220° and boiled for a short time with water, the residue is washed with cold water, and then with a little dilute alcohol. A white, amorphous, tasteless powder, nearly insoluble in water; slowly acted on by cold water, more quickly by hot water, yielding P₂O₃(NH₂)₂(OH)₂Aq and NH₃Aq; boiled with HClAq gives H₂PO₄ and NH₃. Pyrophosphotriamates of NH₄, Ba, Cd, Co, Cu, Cr, Fe, Pb, Mg, Mn, Hg, Ni, Pt, K, Ag, Tl, and Zn are described; the acid is said to be tetrabasic.

Pyrophosphonitrilio acid $P_2O_3(OH)N$. The NH₄ salt of this acid is said to be formed by heating $P_2O_3(NH_2)_3OH$, and the K salt by heating $P_2O_3(NH_2)_3OK$ (G., C. J. 22, 19).

TETRAPHOSPHAMIC ACIDS AND ALLIED COMPOUNDS. By the reaction of NH₃ with POCl₃ under different conditions, Gladstone (C. J. 21, 261; 22, 15), obtained various compounds which may be regarded as (1) tetraphosphodiamic acid P₄O₇(NH₂)₂(OH)₃, (2) P₄O₇(NH₂)₄(OH)₂ tetraphosphotetramic acid, (3) ? tetraphospho-imidotetramide P₄O₇(NH₂)₁NH (called by G. tetraphosphopentazotic acid), (4) ? tetraphosphodimido-diamide P₄O₇(NH₂)₂(NH)₂ (called by G. tetraphosphotetrimic acid). Some of the above acids are regarded by Mente (A. 248, 232) as imido-acids.

M. M. P. M.

PHOSPHAMIDES. I. PHOSPHORIC TRIAMIDE O.P(NH.)s. (Phosphoryl triamide) [derived from O.P(OH)s=HsPOs]. This compound is said by Schiff (A. 101, 300) to be obtained by passing NHsgas into POCls, and treating the product with water. It is described as a snow-white, amorphous solid; heated out of air gives NHs and PON; fused with KOH gives NHs and K phosphate; scarcely acted on by boiling water, KOHAq, or dilute acids; soluble in hot conc. HsO,; slowly attacked by boiling conc. HNOs or HClAq. Gladstone (C. J. 22, 18) could not obtain this compound.

II. PHOSPHOROUS DIAMIDE OH.P.(NH₂)₂ [derived from OH.P(OH)₂ = H₂PO₃]. The white powder produced by passing NH₃ into P₄O₄ in benzene or ether has probably this composition (Thorpe a. Tutton, $C.\ J.\ 59$, 1027). It dissolves in water with incandescence; HClAq reacts violently, forming non-inflammable PH₃, P, NH₄Cl, H₂PO₃Aq, and H₃PO₄Aq.

THIOPHOSPHAMIDE IS described under THIO-

PHOSPHORYL CHLORIDE, Reactions, No. 7, p. 148.
M. M. P. M.

PHOSPHAMIDO-IMIDE PO.NH.NH₂ (Phosphoryl imido amide. Formerly called Phosphodiamide) (Gerhardt, A. Ch. [3] 18, 188; Schiff, A. 101, 300). Obtained by saturating PCl₄ with NH₄, boiling the product with water as long as HCl passes into solution, then boiling

with potash, then with HNO, Aq or H2SO, Aq, washing, and drying over H₂SO₄. A white powder, insoluble in alcohol, water, or turpentine; heated out of air gives NH, and PON; heated in moist air gives NH, and HPO;; fused with KOH gives K phosphate and NH,; slowly oxidised by fusion with KNO, but resists action of most oxidisers. M. M. P. M.

PHOSPHATES. Salts of the phosphoric acids. Besides the salts directly derived from the three phosphoric acids HPO_s , H_sPO_4 , and $H_4P_2O_7$ (v. Phosphoric acids, p. 124), phosphates exist which are probably derived from hypothetical di-, tri-, or meta- phosphoric acid nHPO; and a few salts have been isolated which are perhaps derived from the hypothetical acids P₄O₇(OH)₆ and P₁₀O₁₉(OH)₁₂.

Composition.—Orthophosphates are de-

rived from H,PO4; they may be divided into classes according to the replacing powers of the metals: (1) monovalent metals form MH₂PO₄, M₂HPO₄, and M₂PO₄; (2) divalent metals form MH₄P₂O₈, MHPO₄, and M₅P₂O₈; (8) trivalent metals form MH₃P₂O₈ and M₂P₂O₈; (4) tetravalent metals form MH₂P₂O₈ and M₃P₄O₁₈.

Pyrophosphates are derived from H,P,O,: (1) salts of monovalent metals M2H2P2O, and M₄P₂O₇; (2) salts of divalent metals MH₂P₂O₇ and M₂P₂O₇, &c. Metaphosphates are derived from HPO₃:

(1) salts of monovalent metals MPO,; (2) salts of divalent metals MP2O6, &c. Various modifications of Na (and other) metaphosphates are known, all having the empirical formula MPO, but differing both in physical and chemical properties from one another; these salts were classified by Fleitmann a. Hanneberg (A. 65, 304; F.,P. 78, 233, 338) as: (1) dimetaphosphates, salts of $H_2P_2O_6$; (2) trimetaphosphates, salts of $H_2P_3O_6$; (3) tetrametaphosphates, salts of $H_4P_4O_{12}$; and (4) hexametaphosphates, salts of $H_6P_6O_{18}$. Tammann (Z. P. C. $\hat{6}$, 122) has example $\hat{6}$. mined the electrical conductivities, and the depressions of freezing-points, of aqueous solutions of several di-, tri-, and hexa-metaphosphates; he comes to the conclusion that Fleitmann's dimetaphosphates should be represented in solution as M₃(PO₃), (i.e. as trimetaphosphates), and that his tri-salts when in solution are really $M_2(PO_2)_2$ (i.e. dimetaphosphates). Tammann's results point to the existence in aqueous solution of 3 isomeric hexametaphosphates which may be represented as (i.) $M_2.M_4(PO_3)_6$ e.g. $Na_2.Na_4(PO_3)_6$, $K_2.Na_4(PO_3)_6$; (ii.) $M_4.M_2(PO_3)_6$.c.g. $Na_4.Na_2(PO_3)_6$, $Ag_4.Na_2(PO_3)_6$; (iii.) $M_5.M(PO_3)_6$ e.g. $Na_5.Na(PO_3)_6$; $(NH_4)_5.Na(PO_3)_6$. There are also, according to Tammann, three different Na metaphosphates insoluble in water (v. also T., J. pr. [2] 45, 417).

Tetra- and deka-phosphates. Salts of the compositions Na_pP₄O₁₂ and Na₁₂P₁₀O₃₁ are obtained by fusing together Na₄P₂O₇ and Ns. (POs); these salts may be regarded as derived from the hypothetical condensed acids $H_0P_4O_{12}$ (* $2H_4P_2O_7-H_2O$), as (= $5H_4P_2O_7-4H_2O$), respectively. and H,2P,0O,1

The composition of all the phosphates may be represented empirically by the expression nMO.mP.O.xH.O, where MO stands for a basic oxide.

Occurrence.—Phosphates of Al and Ca occur

in large quantities in many rocks; phosphates of Fe, Pb, Mg, NH4, &c. are also widely distributed minerals. Coprolites (the fossilised excrements of former land-animals), and guano (the excrement of certain birds) consist very largely of Ca phosphate. Phosphates occur in some kinds of coal, in the ashes of plants, and in the bones of animals.

Formation.—Some of the orthophosphates are obtained by reacting on H₃PO₄Aq with metallic oxides or carbonates; others are formed by double decomposition from the alkali phosphates; when a metaphosphate is fused with a metallic oxide or hydroxide a normal orthophosphate is generally formed. Many pyrophosphates are obtained by double decomposi-tion from Na₄P₂O₄, which is formed by strongly heating Na₂HPO₄; other pyrophosphates are found by neutralising H₄P₂O₄Aq by basic oxides or hydroxides. Metaphosphates are generally formed by strongly heating orthophosphates MH,PO, or MRHPO, when M or R is a metal whose oxide is volatile; many are produced by double decomposition from NaPO₃; some are formed by heating Cu₂(PO₃), with a metallic sulphide and filtering from CuS.

The processes by which any one of the three classes of phosphates is formed from another is perhaps best understood by representing the

phosphates thus:

meta pyro P₂O₅.2M₂O P2O3.M2O $P_2O_5.3M_2O$ Metaphosphate becomes pyro- by gaining M.O. and becomes ortho- by gaining 2M2O, and so on. Metaphosphate fused with metallic oxide, hydroxide, or carbonate yields pyrophosphate or orthophosphate, according to the quantities used; thus (1) 2NaPO₃ + Na₂CO₃ = Na₄P₂O₇ + CO₂; (2) 2NaPO₃ + 2Na₂OO₂ = 2Na₃PO₄ + 2CO₂. The metaphosphate of a heavy metal yields an orthophosphate and H3PO, by boiling with water; thus $3AgPO_8 + 8H_2O = Ag_3PO_4 + 2H_3PO_4$. change from a pyro- to an ortho- phosphate is often effected by boiling with water, the product being an acid salt, thus Na₄P₂O₇+H₂O = 2Na₂HPO₄; or the pyrophosphate is fused with hydroxide or carbonate, thus

 $Ba_2P_2O_7 + BaO_2H_2 = Ba_3(PO_4)_2 + H_2O_4$ Acid orthophosphates with one atom of H yield pyro-phosphates when strongly heated, e.g. $2Na_2HPO_4-H_2O=Na_4P_2O_7$; and those with two atoms of H yield metaphosphates, e.g.

 $NaH_2PO_4 - H_2O = NaPO_2$ Similarly a double orthophosphate of the form M_2RPO_4 yields a pyrophosphate on heating if R forms a volatile oxide, e.g. $2Na_2(NH_4)PO_4 = Na_4P_2O_7 + H_2O + 2NH_3$. The removal of M_2O from a normal orthophosphate with formation of a metaphosphate is often effected by fusing with SiO₂ or B₂O₃ or other anhydride which forms a salt with M.O. e.g. Na, PO, + SiO, = NaPO, + NagSiO,

The metaphosphates formed by the processes sketched above belong to several series of salts all of which have the empirical formula nMPO₃. According to Fleitmann a. Henneberg (A. 65, 304; F., P. 78, 233, 338), these salts belong to five series.

Monometaphosphates MPO: Not many of these have been obtained with certainty; they are produced by heating alkaline oxides with

equivalent quantities of H₂PO₄ to redness until the residue ceases to give an acid reaction (the temperature not being high enough to melt the mass), and washing the residue with cold water; some of the salts—e.g. NaPO₃ and KPO₃—are formed by heating MH₂PO₄ or M₂H₂P₂O₇ to redness (c. 315°) till the residue is almost wholly insoluble in cold water (Maddrell, G. S. Mem. 3,

Dimetaphosphates M₂P₂O₆. The salts of Cu, Zn, and Mn are prepared by heating equivalent quantities of MO and H₂PO₄ to redness. The Na and NH₄ salts are obtained by decomposing the Cu salt by Na₅SAq or NH₄ sulphide solution, filtering, and ppg. by alcohol. These salts yield others of the series by double decomposition. The salts of this series are soluble in water and crystallisable. Tammann (Z. P. C. 6, 122) thinks these salts are really trimetaphos-

phates (v. Properties and Reactions).

Trimetaphosphates M₂P₂O₆. The salts of this series are crystalline. The Na salt is prepared by slowly heating NaNH₄HPO₄ with frequent stirring, till the residue has a faintly acid reaction, treating with cold water, filtering, and allowing the filtrate to crystallise. The salt is also formed by heating NaNH₄HPO₄ till a glassy mass remains, and cooling very slowly. The other salts are obtained from the Na salt by double decomposition. Tammann (l.c.) regards these salts as dimetaphosphates (v. Properties and Reactions).

Tetrametaphosphates M₄P₄O₁₂. PbO is heated with H₂PO₄ to redness, as in making the Cu, Zn, and Mn dimetaphosphates (v. supra); the crystalline salt which is formed is insoluble in water, and much more easily decomposed by acids than the Pb salt got from Na₂P₂O₆. The Na salt is obtained by decomposing the Pb salt by Na₂SAq; and other salts are obtained from the Pb salt by double decomposition. Salts of Bi and Cd are obtained similarly to the Pb salt. These phosphates are uncrystallisable. Tammann (J. pr. [2] 45, 417) says that CuNa₂P₄O₁₂ is the only salt which certainly belongs to this series.

Hexametaphosphates M₆P₆O₁₆. NaNH₄HPO₄ is heated till a glassy mass remains, which is cooled rapidly (Graham, T. 1833.253). The Na salt yields others by double decomposition. These salts are non-crystallisable, and dry to resinous-like masses. Tammann's experiments (l.c.) seem to show that Na₆P₆O₁₈ is a mixture of several isomeric salts (v. p. 118).

Tammann (l.c.) describes salts belonging to the series of penta-, okto-, deka-, and tetrakai-

deka- metaphosphates.

Tetraphosphates M₆P₄O₁₃. A few of these salts have been prepared, e.g. Na₆P₄O₁₃ by fusing 2NaPO₂ + Na₄P₂O₇, and Ba₃P₄O₁₂ by fusing the Na salt with BaCl₂ (F. a. H., A. 65, 304; Uelsmann, A. 118, 99).

Delaphosphates M₁₂P₁₀O₃₁. The Na salt was obtained by fusing Na₄P₂O₇ + 8NaPO₂ (F. a. H., l.c.); some other salts were obtained from the Na salt by double decomposition (U., l.c.).

Properties and Reactions.—The orthophosphates MH_PO₄ dissolve in water, forming acid solutions; the dimetallic salts M_HPO₄ dissolve in water when M is an alkali metal, the solutions are feebly alkaline. Of the normal salts M_PO₄,

only those of the alkali metals are soluble in water; the solutions have an alkaline reaction, and are decomposed very easily, even by CO2 forming M₂HPO,Aq. Most of the heavy metals form only M₂PO₄. Solutions of alkali orthophosphates give a yellow pp. (Ag₃PO₄) with AgNO₂Aq, and a yellow pp. on warming with NH₄ molybdate solution. The orthophosphates M₃PO₄ are not decomposed by heating strongly, unless they are salts of volatile bases; M2HPO4 give M₂P₂O₇ and H₂O, and MH₂PO₄ give MPO₃ and H₂O, when strongly heated. M₃PO₄, M= alkali or alkaline earth metal, are not changed by heating with charcoal, but M₂HPO₄ and MH₂PO₄ give M₃PO₄ and P. When M is a heavy metal, the salts M,PO, generally give metallic phosphides by heating with C. Orthophosphates heated with K or Na yield alkali phosphide. Many insoluble orthophosphates are decomposed by fusion with alkali carbonate, but those of the alkaline earths are only partially decomposed. H₂SO₄ decomposes all orthophosphates. When the salts M_sPO₄ are heated with SO₂Aq under pressure, M₂HPO₄ or MH₂PO₄ are formed (Pitter, Chem. Indust. 1878. 398; Gerland, J. pr. [2] 4,

Of the pyrophosphates, the normal alkali salts are soluble in water, with feebly alkaline reactions; the others are generally insoluble but dissolve in Na, P₂O, Aq, forming double salts (H. Rose, P. 76, 13; Schwarzenberg, A. 65, 133; Persoz, A. 65, 163). Solution of pyrophosphates give orthophosphates when boiled with the stronger acids, or heated with water to 280° (Reynoso). M.P.O. are not changed by heat unless they are salts of bases decomposed by heat; M₂H₂P₂O₇ give MPO₃ and H₂O when strongly heated. All the pyrophosphates are changed to orthophosphates by heating strongly with sufficient alkali or alkali carbonate. Pyrophosphates of metals whose oxides are not reduced by H give orthophosphates and PoO. when heated in H; those which contain metals whose oxides are reduced by H, but not by heat alone, give metallic phosphide when heated in H; and those containing metals the oxides of which are reduced by heat alone, yield metal, H_2O , H_3PO_4 , &c., when heated in H (Struve, J. pr. 79, 845). Pyrophosphates in solution give a white pp. with AgNO, Aq; they do not give an immediate pp. with warm NH, molybdate, and they do not coagulate albumen when acidified by acetic acid.

Some metaphosphates are soluble, others are insoluble, in water; some are crystalline, others are amorphous (v. supra: Formation of different classes of metaphosphates). Solutions of these salts in water are changed to orthophosphates on long-continued boiling; they give a white pp. with AgNO₂Aq, no immediate pp. with NH₄ molybdate, and they coagulate albumen after acidification by acetic acid.

Tammann (Z. P. C. 6, 123) has examined the electrical conductivities of solutions of Fleitmann's di- and tri- metaphosplates of Na; his results indicate that the ions of the salts in solution are Na_x and (PO_s)_x. He has also measured the depression produced in the freezing-point of water by dissolving these salts therein; and, taking the results along with the electrical conductivities, he concludes that Fleitmann's di-

salts are tri-salts, and F.'s tri-salts are really di-salts. By the reaction of AgNO3Aq on the Na_eP_eO₁₈ of Graham and Fleitmann, Tammann obtained a crystalline salt Ag, P,O18, and two salts Ag₄Na₂P₆O₁₈ and Ag₅NaP₆O₁₈, and from these he formed three Na salts which he formulates as Na₆P₆O₁₈, Na₄.Na₂.P₆O₁₈, and Na. Na. P.O18. Measurements of the electrical conductivities of these salts confirm the above formulæ, and they tend to show that the ions of the salts in solution are Na, and Na, PoO18, and Na, and NaP,O,, respectively. A fuller examination of Graham's salt led T. to conclude that it contains another isomeric hexametaphosphate Na2.Na4.P6O18

For detection and estimation of phosphates a

manual of analysis must be consulted.

ORTHOPHOSPHATES. Salts derived from H₃PO₄. (For thioxyorthophosphates v. Phos-

PHORIC SULPHIDE, p. 147.)

Aluminium orthophosphates. Many occur in minerals, e.g. angelite, fischerite, wavellite, turquois. The pps. obtained by adding Na₂HPO₄Aq to alum solutions vary in composition. may be represented as xAl_2O_3 , yP_2O_3 , x being chiefly 1, 2, 3, and y being 1 and 2 (Munroe, A. 159, 638; Wittstein, J. B. 27, 167; Fuchs, S.

24, 121; Millot, C. R. 82, 89).

The normal salt, AlPO, xH2O, is obtained by gradually pouring neutral alum solution into excess of Na₂HPO₄Aq; a white loose powder; soluble mineral acids, soluble organic acids, soluble NH, citrate solution (Erlenmeyer, B. 14, 1869), soluble Al salt solution (Fleischer, Fr. 6, 28), soluble in potash; infusible at red heat, fused with Fe and SiO₂ gives Fe phosphide (Schlösing, C. R. 67, 1247); partially decomposed by fusing with K₂CO₃ (Rammelsberg, P. 64, 405; cf. Vauquelin, A. Ch. [2] 21, 138). When air-dried x = 4 to 5; dried over H_2SO_4 , x = 3 (Rammelsberg, Millot, Wittstein). For other phosphates of Al v. Rammelsberg (l.c.), Schweitzer (B. 3, 310; 6, 28), Ludwig (Ar. Ph. [2] 59, 19), Hautefeuille a. Margottet (C. R. 106, 135), Erlenmeyer (Z. 3, 572).

Double salts with phosphates of Ca, Li, and Mg have been described (v. Gm.-K., i. 2,

Ammonium orthophosphates. The normal salt, Am, PO. 3H,O, crystallises in short pyramids, obtained by treating Am2HPO4 with conc. NH, Aq (Berzelius, Gm.-K. i. 2, 527). Loses two-thirds NH, by boiling water (Kraut, C. C. 1855. 894). Sestini (B. 11, 1927) obtained a salt with 5H₂O. The di-ammonium salt, Am₂HPO₄, is formed in monoclinic prisms by allowing NH, or (NH₄)₂CO₃ in H₃PO₄Aq to evaporate; soluble four parts cold water, solution is alkaline; on boiling NH, is evolved; S.G. 162-167 (Rammelsberg, B. 3, 278; Schiff, A. 112, 88; Buignot, J. 1861. 15; Proust a. Brooke, Gm.-K. i. 2, 528). The mon-ammonium salt, AmH2PO4, crystallises in quadratic forms, isomorphous with KH_PO, by adding NH_Aq to H_PO_Aq till BaCl_Aq ceases to give a pp. S.G. 1.758 (Schiff, *l.c.*; Mitscherlich, *P.* 19, 385).

Barium orthophosphates. The normal salt, Ba₃P₂O₈, is a heavy white solid, obtained by ppg. BaCl₂Aq by Na₃PO₄Aq (Graham; Rotondi, B. 15, 1441). Two acid salts have been obtained: (1) BaHPO, by boiling the normal salt with |

H₂PO₄Aq, or by ppg. BaCl₂Aq by Na₂HPO₄Aq; soluble in 10,000 to 20,000 parts water, easily soluble if NH, salts, BaCl, or NaCl is present; soluble in H.PO.Aq or dilute HNO.Aq (Berzelius, Ludwig, Rose, Brett, Wittstein, Bischoff, Malaguti, Gm.-K. i. 2, 272; Skey, C. N. 22, 61; Erlenmeyer, J. 1857. 145; Setschenoff, C. C. 1875. 97; Gerland, J. pr. [2] 4, 125); (2) $BaH_4P_2O_8$, from a solution of $BaHPO_4$ in H_3PO_4 ; white triclinic crystals; decomposed by much water to H₃PO₄Aq and BaHPO₄ (Berzelius).

Double salts.—(1) 3Ba,P₂O₄,BaOl₂ (Rose, P. 8, 211; Ludwig, Ar. Ph. [2] 56, 265; Deville

a. Caron, A. Ch. [3] 67, 451);

(2) 4BaHPO₄.BaCl₂ (Erlenmeyer, J. 1857. 145); (3) BaNaPO₄.10H₂O (Rose, Guyton, Berzelius, Gm.-K. i. 2, 313; de Schulten, Bl. [2] 39, 500; Joly, C. R. 104, 1702); (4) BaKPO.10H2O (de S., l.c.).

Beryllium orthophosphates. The normal salt, Be, P2O8, is a white pp., obtained by adding Na₂HPO₄Aq to solution of a salt of Be. An salt in H₃PO₄Aq (Vauquelin, *Qm.-K.* i. 2, 269).

Double salts. Be.NH₄ and Be.K phosphates are known (Rössler, *Fr.* 1878. 148;

Ouvrard, C. R. 110, 1333).

Bismuth orthophosphate BiPO. White powder, obtained by adding Bi₂O₃ xH₂O to H,PO,Aq; little known (Wenzel, Gm.-K. i. 2, 832; Chancel, C. R. 50, 416).

Cadmium orthophosphates. The normal salt, Cd₃P₂O₈, is obtained as an amorphous pp. by adding Na_sPO₄Aq to a Cd salt solution (de

Schulten, Bl. [3] 1, 473).

Acid salts (de S., l.c.): (1) Cd₅H₂P₄O₁₆.4H₂O, by ppg. hot CdCl₂Aq by Na₂HPO₄Aq.

(2) CdH₂P₂O₈.2H₂O, by dissolving the foregoing salt in cold dilute H.PO.Aq and allowing to evaporate at ordinary temperature.

Double salt 3Cd₃P₂O₅.CdCl₂ (de S., l.c.). Calcium orthophosphates. Ca phosphates

are widely distributed minerals.

Normal salt Ca,P2O,. Obtained, as a white amorphous powder, by adding NH,Aq to solution of bone-ash in HClAq or HNO, Aq; by adding Na,PO,Aq, or a mixture of Na,HPO,Aq and NH,Aq, to CaCl,Aq. For details of preparation from phosphorite, or bone-ash, &c. v. DICTIONARY OF APPLIED CHEMISTRY. The pp. from CaCl2Aq in presence of NH₃ contains 5H₂O according to Wittstein (J. 1866, 168; v. also Ludwig, Ar. Ph. [2] 69, 286). Slightly soluble in water; one litre cold water dissolved 79 mgm. freshly ppd., and 31 mgm. strongly heated, Ca₃P₂O₈ (Völcker, J. 1862. 131). Decomposed slowly by boiling water into CaH₄P₂O₈ and 2Ca₃P₂O₈.CaO₂H₂ (Warington, C. J. [2] 11, 983). Solubility in water is increased by small quantities of alkali salts (v. Wöhler, P. 4, 166; Liebig, A. 61, 128 Lassaigne, J. 1847. 341; Thomson, Ann. Philos. 17, 12; Volcker, J. 1862. 131; Fresenius, Fr. 10, 133; Hilnefeld, Delkeskampf, Morveau, Gm.-K. i. 2, 367; Terreil, Bl. [2] 35, 548; Erlenmeyer, B. 14, 1253). Presence of starch and various organic substances increases solubility in water (v. Wöhler, A. 98, 143; Hayes, N. Ed. P. J. 5, 378). Soluble in CO₂Aq; CaCO, and CaHPO, separate on standing (Liebig, A. 106, 196; Warington, C. J. [2] 4, 296; Dusart a. Pelouze, C. R. 66, 1327; von Georgievics, M. 12, 566).

Partially decomposed by boiling with NaOHAq, or by fusing with Na2CO3 (Wöhler, A. 51, 437; Rose, P. 95, 437; Fresenius, Fr. 10, Easily changed to CaC₂O₄ by boiling

with (NH₁)₂O₂O₄Aq (Fresenius, *l.c.*).

Strongly heated with C and SiO₂, gives Ca silicate, P, and CO; heated in CO, all P is set free (Schlösing, C. R. 59, 384; Anderson, B. 5, 1065). Heated with Na, K, or Mg, metallic phosphide is formed (v. Gm.-K. i. 2, 366).

Dissolved by SO₂Aq; on standing, CaHPO₄ and CaSO₃ separate (v. Gerland, C. N. 20, 268; Aldred, C. N. 42, 177; Rotondi, B. 15, 1441). Dilute H2SO, Aq forms CaHPO, or CaH4P2O, or CaSO, and H.PO, according to the quantities of H_2SO_4 and $Ca_3P_2O_8$, the temperature, and the time of action (v. Gm.-K. i. 2, 369; Kolb, C. R. 78, 825; Joulie, C. R. 76, 1288; Crum, A. 63, 394). For the (similar) reactions with HClAq and HNO, Aq v. Crum, l.c.; Piccard (J. 1886. 168); Bischof (S. 67, 39); Birnbaum a. Packard (Z. [2])7.137); Tissier (C. R. 38, 192); Warington (C. J.

[2] 4, 302).

Double salts. (1) Ca₃P₂O₈.Ca₂SiO₄ (Carnot a. Richard, C. R. 97, 316). (2) 3Ca₃P₂O₈ CaCl₂by fusing Ca₃P₂O₈ with NaCl, or heating to 250° with CaCl2Aq, or passing PCl3 vapour over glowing lime (Forchhammer, P. 91, 568; Debray, A. Ch. [3] 61, 424; Daubrée, Ann. M. (4) 19, 684; Deville a. Caron, A. Ch. [3] 67, 451). (3) 3Ca₃P₂O₈.CaF₂; occurs native as apatite (with varying quantities of CaCl₂); obtained by fusing Na₂HPO₄, CaF₂, and CaCl₂ (D. a. C., l.c.; Briegler, A. 97, 95). (4) Ca(NH.)PO, xH.O., (Herzfeld a. Fcuerlein, Fr. 1881. 191). (5) CaKPO, (Rose, P. 77, 291). (6) CaLiPO, (Rose, P. 77, 298). (7) CaNaPO, (Rose, P. 77, 291; Bunge, A. 172, 18).

I. Dicalcium Acid salts. phosphate Ca2H2P2O8. Obtained by adding Na,HPO4Aq, or NaNH, HPO, Aq, to excess of CaCl₂Aq; pp. contains a little Ca₃P₂O₈: also by the action of CO₂Aq on CaCO, and (NH4)2HPOAq (Becquerel, C. R. 34, 573); and by acting on powdered CaCO, with H_3PO_4Aq (Debray, A. Ch. [3] 61, 424); by passing CO₂ into H₂O, holding freshly ppd. Ca,P₂O₈ in suspension (Percy, P. M. [3] 26, 194; Reichardt, Ar. Ph. [3] 2, 236). Obtained in crystals by allowing solutions of (NH4)2HPO4 and CaCl2 to mix slowly through a porous cell (Vohl, A. 88, 114), or through a layer of water (Drevermann, A. 87, 120); also by allowing a solution of amorphous CaHPO, in a weak acid to evaporate (Bödeker, A. 69, 206; Baer, P. 75, 152; Vorbringer, Fr. 9, 457; v. also Causse, J. Ph. [5] 21, 544). CaHPO, ppd. at low temperatures and dried quickly at 100° contains 2H₂O, which is lost by long drying at 100° in a stream of dry air. Forms Ca,P,O, when strongly heated; decomposed to Ca,P,O, and CaH,P,O, by long boiling with water. For description of various hydrates &CaHPO.yH2O v. Vorbringer (Fr. 9, 457), Gerland (J. pr. [2] 4, 104), Millot (Bl. [2] 33, 194), Skey (C. N. 22, 61), Dusart a. Pelouze (C. R. 66, 1827), Davies (C. N. 64, 287). Soluble in boiling NH, ClAq (Kraut, Ar. Ph. [2] 111, 102); soluble SO, Aq (Gerland, J. pr. [2] 4, 123), in HClAq and HNO, Aq (Berzelius, Baer, Gm.-K. i. 2, 372). Decomposed by boiling alkali carbonate solutions (Malaguti, Gm.-K. i. 2, 378)

II. Monocalcium phosphate CaH, P,O, H,O.

This salt is the chief constituent of artificial 'superphosphate of lime' prepared by treating crushed bones, phosphorite, &c., with H.SO. (v. DICTIONARY OF APPLIED CHEMISTRY). Prepared by dissolving Ca₂H₂P₂O₈ in 31 p.c. H₃PO₄Aq, crystallising, washing with absolute alcohol, then with ether, and drying (Stoklasa, Listy Chem., 13, 203, 240, 278; abstract in C. J. 58, 695). This salt is also formed by evaporating Ca₂H₂P₂O₈ with HNO₃Aq + H₂SO₄Aq. Also by leaving a solution of the commercial honey-like monocalcium phosphate in contact with Ca, P,O, for some time, and evaporating the solution formed at a gentle heat; the salt is thus obtained in crystals (Pointet, Bl. [3] 5, 254). Not hygroscopic; the hygroscopic nature of the preparation made by Birnbaum and others is said by Stoklasa (l.c.) to be due to traces of H,PO,. Thin rhombic plates (v. Haushofer, Z. K. 7, 263); S.G. 2.02 (Schröder, B. 11, 126). Loses H₂O at 100°, decomposes at 200° to mixture of CaP₂O₄ and Ca₂P₂O₇ (Birnbaum, B. 6, 898). Decomposed by H₂O. Stoklasa (l.c.) represents the decomposing action of water as

xCaH₄P₂O₂.H₂O+H₂O+H₂O =(x-1)CaH₄P₂O₂.H₂O+CaHPO₄+2H₂O+H₂PO₄; the greater the quantity of water the more salt is dissolved without decomposition; with 200 parts H2O to 1 part salt a clear solution is formed without any free H₃PO₄; above this limit CaH₄P₂O₈ is re-formed from the products of its decomposition (for older observations v. Birnbaum, l.c.; Erlenmeyer, N. J. P. 7, 225; Warington, Fr. 1880. 243). According to Reynoso (C. R. 34, 795), water at 280° decomposes the salt, forming H₃PO, and crystalline Ca₃P₂O₈. Various salts produce Ca₂H₂P₂O₈ from solutions of CaH₄P₂O₈ (v. Debray, A. Ch. [3] 61, 424; Dusart a. Pelouze, C. R. 66, 1327; Vorbringer,

Fr. 9, 457; Erlenmeyer, l.c.; Birnbaum, l.c.).
Cerium orthophosphate CePO. By fusing Ce₂O₃ with KPO₃; also by digesting Ce₂K₃(PO₄)₃ with KClAq. Monoclinic prisms; S.G. 3.8 (Ouvrard, C. R. 107, 37). The double salt Ce₂K₃(PO₄)₃ is formed by fusing K₂PO₄ or K₄P₂O₇.

Chromium orthophosphates. The normal chromic salt, $Cr_2P_2O_8$, is known in two forms, green and violet. Violet form, $Cr_2P_2O_8$.12H₂O, a violet crystalline pp. by adding Na₂HPO₄Aq to excess of Cr alum solution; loses 7H₂O at 100°, becoming green; changed to green by conc. HNO, Aq, H, SO, or PCl, (Rammelsberg, P. 68, 383; Etard, C. R. 84, 1091). Green form, Cr₂P₂O₈.6H₂O, by adding excess of Na₂HPO₄Aq to Or alum; also by ppg. a slightly acid solution of a Cr salt by Na₂HPO₄ and Na acetate (Rose, P. 77, 291, 298; Carnot, Bl. [2] 37, 482; Vauquelin, Gm.-K. i. 2, 304). Soluble mineral acids, also KOHAq (v. Dowling a. Plunkett, Chem. Gazette, 1858. 220; Kämmerer, Fr. 12, 375). Gives alkali chromates by fusion with alkali sulphates (Grandeau, C. R. 95, 921). Forms a double salt with Ca (Dingler, D. P. J. 212, 532; Plessy, Rep. China app. 1862. 453; Köthe, D. P. J. 214, 59). The acid salt CrH₃P₂O_{8.8}H₂O is probably formed by dissolving Cr₂O₃.xH₂O in H₂PO₄Aq (Vauquelin, Gm.-K. i. 2, 804; Haushofer, Z. K. 7, 263).

Chromous orthophosphate Cr.P.O. H.O. a blue pp. by adding Na,HPO,Aq to OrCl,Aq; quickly becomes green in air (Moissan, A. Ch. |

Cobalt orthophosphates. The normal salt Co.P.O.xH2O is formed by heating CoHPO.3H2O with water, or with Co(NO₂)₂Aq, to 250° in a sealed tube (Debray, A. Ch. [3] 61, 438); also by treating Co(NH₄)PO₄.6H₂O with boiling water (Chancel, Precis d'analyse [Paris, 1862] 1, 263). In this salt x=2. A salt with x=8 is said to be formed by ppg. CoH, P,O,Aq by alcohol (Reynoso, C. R. 84, 795; Reitler, Vierteljahr. Pharm. 7, 246). Two acid salts are known: (1) CoHPO.3H.O, by boiling CoCO, with H.PO.Aq, or boiling solution of a Co salt with CaH.P.O. (Debray, J. Ph. [3] 46, 121); a salt with 5H.O was obtained by Bödeker (A. 94, 357). (2) CoH₁P₂O₈, by heating Co pyrophosphate with water to 280° (Reynoso, C. R. 34, 795). Double salts: NH₄ salts v. Dirvell, C. R. 89, 903; Popp, Z. [2] 6, 305. For luteo- and roseo- cobaltamine phosphates v. Gibbs a. Genth, Am. S. [2] 21, 86; 23, 234, 319; Braun, Gm.-K. i. 8, 463; Porumbaru, C. R. 93, 842. For K and Na double salts v. Ouvrard, C.R. 106, 1729.

Copper orthophosphates. Normal salt Cu₂P₂O₈3H₂O, obtained by adding a little Na₂HPO₄Aq to excess of a Cu salt solution (Mitscherlich, Gm.-K. i. 3, 614); also by heating CuCO₃ with H₂PO₄Aq to 70° (Debray, A. Ch. [3] 61, 437). A blue-green powder; decomposed by hot water under pressure (Debray; v. also Reynoso, C. R. 34, 1795; Jörgensen, Gm.-K. i. 3, 615). Basic salts and acid salts are known; composition doubtful (v. Rammelsberg, P. 68, 383; Metzner, A. 149, 67). For double salts with NH, v. Metzner, l.c., and Schiff, A. 123, 41; with Na, v. Weineck, A. 156, 57; with K, also with Na, v. Ouvrard, C. R. 111, 177. For an account of the salts produced by the reaction of Na₂HPO₄Aq with CuSO₄Aq and Cu(NO₈)₂Aq v. Steinschneider, C. C. 1891, ii. 51 (abstract in C. J. 60, 1423).

Didymium orthophosphate DiPO.: double salts with alkali metals (Ouvrard, C. R.

107, 37).

Iron orthophosphates. Ferrous salts. The normal salt Fe₃P₂O₈.8H₂O occurs native as vivianite; it is obtained by ppg. FeSO,Aq with excess of Na₂HPO₄Aq, and digesting at 60°-80° for eight days. Small monoclinic crystals (Rammelsberg, P. 64, 251, 405), nearly colourless but becoming blue in air (Debray, C. R. 59, 40). A salt with one H₂O is obtained by heating Fe₂H₂P₂O₈.4H₂O with water at 250° (Debray, A. Ch. [3] 61, 437). Various acid salts are obtained by dissolving Fe in H,PO,Aq; if the materials are air-free, and evaporation is effected in CO₂, the salt Fe₂H₂P₂O₃.4H₂O is formed (Erlenmeyer, Debray, Rammelsberg, *l.c.*). For double salts with NH, v. Otto, J. pr. 2, 409; and Debray, C. R. 59, 40.

The normal salt FePO, is Ferric salts. obtained by ppg. FeCl₃Aq by Na₂HPO₄Aq; dried at 50° contains 8H₂O, at 100° contains 4H₂O, dehydrated by heating strongly; also formed by oxidation in air of Fe₃P₂O₃ (Debray, Rammelsberg, Wittstein, *l.c.*). Whether the salt ppd. from FeCl, Aq in presence of acetic acid is the normal or a basic salt is undecided (v. Winkler, J. prakt. Pharm. 5, 337; Heydenrich, C. N. 4, 158; Struve, N. Peters. Acad. Bull. 1, 465;

Wackenroder a. Ludwig, Ar. Ph. [2] 53, 1; Mohr, Fr. 2, 520). FePO, xH₂O is sol. c. 1500 parts boiling water; but is decomposed (Lachowicz, M. 13, 357). Acid salts are obtained by dissolving Fe₂O_{1,2}±H₂O in H₃PO₄Aq: Erlenmeyer got FeH₂P₂O₁₂ by evaporating quickly (v. also Waine, C. N. 36, 132; Wagner, Chem. Zeitung, 1881. 247). Hautefeuille a. Margottet (C. R. 106, 1857) obtained various ferric phosphates by dissolving Fe₂O₃.xH₂O in glacial phosphoric acid. Lanthanum orthophosphate LaPO₄: also

double salts with K and Na (v. Ouvrard, C. R.

107, 37).

Lead orthophosphates. The normal salt, Pb_sP₂O_s, is ppd. as a white powder by adding Na_HPO, Aq to Pb acetate solution (Berzelius; Mitscherlich, Gm. 3; v. also Heintz, P. 73, 119; Gerhardt, A. Ch. [3] 22, 505; Haushofer, Z. K. 7, 264). A double salt with PbCl, is obtained by action of Na₂HPO₄Aq on PbCl₂ (Debray, A.Ch. [3] 61, 419; Manross, A. 82, 348; Deville a. Caron, A. Ch. [3] 67, 451). By ppg. boiling Pb2NO, Aq by H,PO, Aq, Heintz (P. 73, 119) obtained the acid salt Pb,H,P,O,s.

Lithium orthophosphates. The normal

salt, Li₂PO₄.H₂O(?2Li₂PO₄.H₂O). By heating Na₂HPO,Aq with solution of a salt of Li (Mayer, A. 98, 183; v. also Berzelius, P. 4, 245; Rammelsberg, C. N. 38, 240, also B. 15, 283; Fresenius, Fr. 1, 42; Kraut, A. 182, 165). Loses H₂O at 100°. Obtained in rhomboidal tables, S.G. 2·41 at 15°, by dissolving in fused LiCl (de Schulten, Bl. [3] 1, 479). Soluble in c. 2,540 parts water. Acid salts: LiH2PO4 and LiH₅P₂O₈·H₂O, by dissolving normal salt in

H₃PO₄Aq (Rammelsberg, l.c.)

Magnesium orthophosphates. The normal salt, Mg₃P₂O₈xH₂O, is obtained with 4H₂O by ppg. Mg salts by alkali phosphate and drying at 100° (Graham; Rose, P. 76, 24; Gregory, A. 54, 98; Stein a. Tollens, A. 187, 79; Fresenius). Forms monoclinic crystals (Haushofer, Z. K. 6, 137). Very slightly soluble water (Völker, J. 1862. 131). For solubility in salt solutions v. Liebig (A. 106, 185), and in citric acid v. Erlenmeyer (B. 14, 1253). Soluble SO₂Aq (Gerland, J. pr. [2] 4, 126; Rotondi, B. 15, 141). Crystals with 6H₂O separate from MgSO₄Aq + Na₂HPO₄Aq at 36° and upwards and the contractive and the c and at lower temperatures a salt with 14H2O is obtained (v. Percy, P. M. [3] 26, 194; Reischauer, N. R. P. 12, 43; 14, 57).

Acid salts. 1. MgHPO, xH₂O (x = 6,7,8,9);

by dissolving magnesia alba in H.PO.Aq (Bergmann, Fourcroy, Gm.-K. i. 2, 446; Debray, J. pr. 97, 116; Haushofer, Z. K. 7, 257).—2. MgH₄P₂O₂; by boiling the foregoing acid salt with water (Schaffner; Rotondi, B. 15, 141).

salts. Magnesium-ammonium orthophosphate MgNH,PO,6H2O. By adding Na HPO Aq to a Mg salt to which NH ClAq and excess of NH, Aq have been added; or by adding a mixture of H₂PO₄Aq and excess of NH₂Aq to a Mg salt. For experiments on different conditions of formation v. Graham (A. 29, 25); Riffault (A. Ch. [2] 19, 90); Mohr (Fr. 12, 36); Gibbs C. N. 28, 51); Lesieur (C. R. 59, 191); Stein a. Tollens (Gm.-K. 1, 2, 476); Kubel (Fr. 8, 125); Kissel (Fr. 8, 164); Brunner (Fr. 11, 30); Berzelius (P. 4, 275); Heintz (Fr. 9, 16). Obtained in large crystals by separating the reacting liquids by a membrane (Vohl, A. 88, 114; Nonier, C. R. 78, 300; Millot, Bl. [2] 18, 20; Millot a. Maquenne, Bl. [2] 23, 238). Loses NH, in air; strongly heated gives Mg_P_Or. For solubility in various solutions v. Fresenius (A. 55, 109); Ebermayer (J. pr. 60, 41); Liebig (A. 106, 196); Völcker (J. 1862. 131); Kissel (Fr. 8, 173); Gerland (J. pr. [2] 4, 127); Millot (Bl. [2] 18, 20); Ville (C. R. 75, 344). Another Mg-NH, phosphate is described by Gawalovski (C. C. 1885. 721). For other double salts: with MgCl₂, v. Deville a. Caron (A. Ch. [3] 67, 456); with MgCl₂ and MgF₂, v. Bischof (D. P. J. 237, 51, 136); with MgCl₂ and CaF₂, v. D. a. C. (l.c.); with K, v. Haushofer (Z. K. 7, 257), Ouvrard (C. R. 106, 1729); with Na, v. Rose, Berzelius, &c. (Gm.-K.i. 2, 476), Ouvrard (l.c.).

Manganese orthophosphates. Normal manganic orthophosphate, MnPO₄,H₂O₄ a greenish grey pp. by adding conc. Mn(No₈)₂Aq to H₂PO₄ and a little boiling water (Christensen, J. pr. [2] 28, 1). For properties of products of reaction of H₂PO₄ on manganates and permanganates v. Laspeyeres (J. pr. [2] 15, 320); Hermann (P. 74, 303); Rose (P. 105, 289); Barreswill (C. R. 44, 44, 1978).

677); Hoppe-Seyler (J. pr. 90, 303).

Normal manganous orthophosphate, Mn₁P₂O₄.æH₂O. Obtained with 14H₂O by ppg. MnSO₄Aq by Na₂HPO₄Aq; loses 7H₂O over H₂SO₄, and one more H₂O at 100° (Erlenmeyer, A. 190, 208). Salts with 9 to 11 H₂O, 7H₂O, and 3H₂O are obtained by decomposing MnHPO₄.xH₂O by water, and drying under different conditions (E., l.c.). The salt without H₂O is formed by strongly heating the hydrates, also by heating Mn₂P₂O₇ in H (Struve, J. 1860. 73). A white amorphous pp., soluble mineral acids and acetic acid, and various salt solutions (v. Berzelius, Gm.-K. i. 2, 472; Heintz, P. 74, 449; Joulin, A. Ch. [4] 30, 272; Gerland, J. pr. [2] 4, 97; Wittstein, Gm.-K. 2, 472; Braun, Fr. 7, 340; Erlenmeyer, B. 14, 1253).

Various acid salts have been described:—
1. MnHPO, 3H₂O (Bödeker, A. 69, 206; Heintz, P. 74, 449; Erlenmeyer, A. 190, 208).—2.
MnH₂P₂O₈.2H₂O (Heintz, Erlenmeyer, l.c.).—
3. Mn₂P₂O₈.MnHPO₄.xH₂O (Erlenmeyer, l.c.; Haushofer, Z. K. 7, 257). Double salts: with MnCl₂ (Deville a. Caron, A. Ch. [3] 67, 459); with Fe₃P₂O₈ and MnCl₂, and with Fe₃P₂O₈ and MnF₂ (D. a. C., l.c.); with NH₄ (Otto, S. 66, 288; Heintz, l.c.); with K and with Na (Ouvrard, C. B. 160, 1700)

C. R. 106, 1729).

Mercury orthophosphates. Normal mercurous orthophosphate, Hg₃PO₄, by ppg. excess of Na₂HPO₄Aq by HgNO₅Aq (Gerhardt, J. 1849. 283). Brooks (P. 66, 63) describes a basic salt A double salt with HgNO₅ is described by Gerhardt (l.c.; v. also Gm.-K. 3, 918). Normal mercuric orthophosphate, Hg₃P₂O₅, is obtained by heating Hg₃PO₄ to low redness, also by adding strongly acidified Hg(NO₃)₂Aq to Na₂HPO₄Aq (v. Gm.-K. 3, 754, 918; also Haack, C. C. 1890. [2] 736). For a double salt Hg.NH₂Hg₂O.PO₄ v. Hirzel (Gm.-K. 3, 820).

Nickel orthophosphates. The normal salt Ni₂P₂O₈.7H₂O is obtained as a green floculent pp. by adding Na₂HPO₄Aq to a Ni salt; heated strongly loses 7H₂O (Rammelsberg, P. 68, 383; Struve, Gm.-K. 3, 543). Double salts: with NH₄ (Debray, C. R. 59, 40); with Mg (Rose,

Gm.-K. 3, 568); with Na and with K (Debray, l.c.; Ouvrard, C. R. 106, 1729).

Potassium orthophosphates. Normal salt K₃PO₄. A white solid, obtained by adding excess of K₂CO₅ to H₂PO₄Aq, and evaporating; slightly soluble cold water, more soluble hot water. Acid salts: 1. K₂HPO₄ (Graham, Funke, Gm.-K. 2, 30). 2. KH₂PO₄; by dissolving K₂CO₅ in so much H₃PO₄Aq that blue litmus paper is turned red but goes blue again on drying. Colourless tetragonal crystals; very soluble water; at red heat gives KPO₃; S.G. 2·29 to 2·4 (v. Gm.-K. 2, 30; also Senarmont, A. Ch. [3] 33, 391; Schiff, A. 112, 88; Buignet, J. 1861. 15).

Silver orthophosphates. Normal salt Ag₂PO₄, a yellow solid, obtained by adding an alkali phosphate to AgNO₃Aq (Wetzlar, Gm.-K. 3, 918; Skey, C. N. 22, 61; Lassaigne, J. Ph. [3] 16, 289; Joly, C. R. 103, 1071). The acid salt Ag₂HPO₄ forms, in white hexagonal crystals, from a solution of the normal salt in H₂PO₄Aq (Berzelius, P. 2, 163; Hurtzig a. Geuther, A. 111, 160; Schwarzenberg, A. 65, 162; Joly, C. R.

103, 1071).

Sodium orthophosphates. The normal salt, Na₂PO₄.12H₂O₇ is obtained by adding excess of NaOHAq to Na₂HPO₄Aq and evaporating. Sixided pyramids; melt at 77°; S.G. 1.618 (Schiff, 4.112, 18); soluble in 2 pts. water at 15°; solution absorbs CO₂ from air, giving Na₂HPO₄Aq. The anhydrous salt, Na₂PO₄, is obtained by heating the salt with 12H₂O, or by heating Na₂HPO₄.12H₂O with NaOH in equal molecular proportions (Graham, Mitscherlich, Gm.-K. 2, 163). S.G. 2.511 at 12°, 2.536 at 17.5° (Clarke's Table of Specific Gravities). For experiments on diffusion of the different Na orthophosphates v. van Bemmelen, B. 11, 1675; Hinteregger, B. 11, 1619; Tobias, B. 15, 2452.

Disodium hydrogen orthophosphate
Na₂HPO₄.12H₂O. (Ordinary sodium phosphate.)
Occurs in urine of carnivorous animals. Prepared by adding a slight excess of Na₂CO₄ to H₂PO₄Aq, and crystallising. (For preparation from bone-ash &c. v. DIOTIONARY OF APPLIED CHEMISTRY.)

Colourless, rhombic, prisms; for measurements v. Senarmont, A. Ch. [3] 33, 391. S.G. 1·5235 at 16° (v. Gm.-K. 2, 165); melts at 35° (Kopp, A. 93, 129; v. also Gm.-K. 2, 165); effloresces in air, losing $5\mathrm{H}_2\mathrm{O}$.

Solubility in water (Mulder, Gm.-K. 2, 165). 100 pts. water dissolve Na. HPO...

TOO DID! HOUSE O	1000110 1102111 011
at 0° 2.5 pts.	at 55° 87.7 pts
5 2.8	60 91.6
10 8.9	65 93.8
1 5 5.8	70 95.0
20 9.3	75 95.8
25 15·4	80 96.6
30 24.1	85 97.2
35 39·3	90 97.8
40 63.9	95 98 4
45 74·8	99 98 8
50 82.5	100 99-

Saturated Na₂HPO₄Aq boils at 105°, and freezes at \sim 45° (Büdorff). S.G. of Na₂HPO₄Aq at 19° (Schiff, A. 110, 70); 2 p.c. = 1·0883, 4 p.c. = 1·0166, 6 p.c. = 1·025, 8 p.c. = 1·0332, 10 p.c. = 1·0418, 12 p.c. = 1·0503 (p.c. of Na₂HPO₄.12H₄O). S.G. solution saturated at 15° = 1·0469, at 16° = 1·0511

(Michel a. Kraft, Stolba, Cm.-K. 2, 166). Loses 12H₂O in vacuo over H₂SO₄, or by heating to 100°; a salt with 7H₂O crystallises from Na₂HPO₄A₂ at 33° (Blücher, Mulder, Clark, Gm.-K. 2, 166). For vapour-pressure of watergas when the salt is warmed v. Debray (C. R. 66, 195), Horstmann (A. Suppl. 8, 125), Pfaundler (B. 4, 773). For change of volume on

melting v. Wiedemann (W. 17, 561). Absorbs HCl, probably forming NaH,PO, and NaCl (Thomas, C. J. 33, 27). Na₂HPO,Aq absorbs CO₂ (Pagenstecher, J. B. 21, 124; Liebig, A. 64, 349; 79, 112; Heidenhain a. L. Meyer, A. Suppl. 2, 157; Marchand, J. pr. 37, 321). The solution decomposes NH Cl (Rose, Gm.-K. 2, 166); boiled with S forms Na, PO, Aq with Na₂S₂O₃Aq and Na polysulphides (Filhol a. Senderens, C. R. 94, 649; 95, 343; 96, 1057). For remarks on character of reactions of Na, HPO, Aq with solutions of metallic salts v. Joly, C. R. 103, 1129.

Trisodium diphosphate Na₃H₃P₂O₈.xH₂O; by exactly neutralising H,PO,Aq by NaOHAq, and evaporating in vacuo (Filhol a. Senderens, l.c.).

Sodium dihydrogen phosphate NaH, PO, H, O; by adding H,PO, to solution of Na,H,P,O, till BaCl₂Aq is no longer ppd. (Mitscherlich, Berzelius, Gm.-K. 2, 167). A salt with 4H₂O was obtained by Joly a. Duffet (C. R. 102, 1391) by

cooling a very conc. solution.

Double salts. 1. Sodium ammonium hydrogen orthophosphate Na(NH4)HPO4.4H2O (Microcosnic salt). Occurs in guano. Obtained by dissolving 5 pts. Na₂HPO₄,12H₂O and 2 pts. (NH₄)₂HPO₄ in water, adding a little NH₃Aq, and allowing to crystallise in the cold (Berzelius). Large, clear, monoclinic crystals; S.G. 1.554 (Schiff); very soluble water, solution loses NH₃. The solid effloresces in air, losing all NH, and 4H₂O, and leaving NaH₂PO₄; when strongly heated leaves Na_a(PO_a)_s. For double salts formed by fusing with metallic oxides v. Wallroth, B. 16, 3059.—2. NaKHPO₄,7H₂O Walfold, B. 10, 5053.—2. NakH-V₄, H₂O (Mitscherlich, Liebig, Schiff, Gm.-K. 2, 167).—3. Na(NH₄)₂PO₄4H₂O (v. Gm.-K. 2, 220).—4. Na₂(NH₄)₂PO₄12H₂O (Herzfeld a. Feuerlein, Fr. 1881. 191).—5. Na₂K₃H₆(PO₄)₄:22H₂O, and Na₃(NH₄)₃H₆(PO₄)₄:3H₂O (Filhol a. Sender College (PO₄)₄:200 (PO₄)₄: derens, C. R. 94, 649; 95, 343; 96, 1057).— 6. $2Na_3PO_4.NaF.xH_2O$; x=19 and 22 (Baumgarten, J. 1865. 219; Thorpe, C. J. [2] 10, 660).

Strontium orthophosphates. Normal salt Sr₃P₂O₈, is ppd. by adding Na₂PO₄Aq or Na₂HPO₄Aq + NH₃Aq to excess of SrCl₂Aq; also by adding cold ammoniacal solution of 90 pts. Na₂HPO₄.12H₂O to 100 pts. SrCl₂ in Aq (Barthe, C. R. 114, 1267). By pouring SrCl₂Aq into Na₃PO₄Aq at c. 10° gelatinous Sr₃P₂O₆ is formed, but rapidly changes to crystalline SrNaPO.9H2O (Joly, C. R. 104, 1702.)—Acid salt SrHPO₄; by adding Na₂HPO₄Aq to SrCl₂Aq (Skey, C. N. 22, 61; Berzelius, Brett, Wittstein, Gm.-K. 2, 324; Malaguti, A. Ch. [3] 51, 346; Barthe, l.c.). Barthe (l.c.) describes another acid salt, SrH₂P₂O₈·2H₂O_.

Double salts. 1. 3Sr,P₂O₈.SrCl₂ (Deville s. Caron, A. Ch. [3] 67, 452).—2. SrKPO₄ (Rose, Weber, Gm.-K. 2, 324).—3. SrNaPO₄.9H₂O (R.

W., l.c.; Joly, C. R. 104, 905, 1702).
Thallium orthophosphates. Normal thallous orthophosphate Tl.PO.; by adding H.PO.Aq and

NH, Aq to TINO, Aq, or mixing Tl2SO, Aq and Na₂HPO, Aq with a little NH, Aq (Crookes, C. N. 3, 193, 303; 7, 290; 8, 159; Lamy, Bi. [2] 4, 193). While silky mass, or needles; insoluble alcohol, easily soluble NH, salts (Carstanjen, J. pr. 102, 65, 129). Acid salts.—TlH₂PO, and Tl₂HPO, xH₂O (Lamy, l.c.); according to Rammelsberg (B. 15, 283) the second of these does not exist. Normal thallic orthophosphate TlPO₄.2H₂O, by adding H₃PO₄Aq to Tl(NO₈)₃Aq (Willm, Bl. [2] 2, 89; 5, 354; Strecker, J. pr. 102, 65, 129).

Thorium orthophosphates. Various double salts of Th and K, and Th and Na, are described by Troost and Ouvrard (C. R. 102, 1422; 105,

30).
Tin orthophosphates. The compositions of the stannous salt, obtained by adding Na,HPO,Aq to SnCl₂ in dilute acetic acid, and the stannic salt, by digesting SnO₂ with H_sPO_4Aq , are somewhat doubtful (v. Lenssen, A. 114, 113; Reynoso, J. pr. 54, 261; Reissig, A. 98, 339; Girard, C. R. 51, 468). Ouvrard (C. R. 111, 177) describes various double Sn-K and Sn-Na phosphates.

Titanium orthophosphates. Various double salts of Ti and K, and Ti and Na, are described

by Ouvrard (C. R. 111, 177).

Uranium orthophosphates. For double U-K and U-Na salts, v. Ouvrard (C. R. 110, 1333). Johnson (B. 22, 976) describes a salt UO₃.2P₂O₅.

Zinc orthophosphates. Normal salt Zn₂P₂O₄.4H₂O; by ppg. solution of a Zn salt by Na₂HPO₄Aq; pp. is gelatinous, but becomes crystalline. Insoluble water, easily soluble in acids, NH₂Aq, and NH₄ salts (v. Mischer-Libert County) and NH₄ salts (v. Mischer-Libert County). lich, Gm.-K. 3, 16; Schindler, Mag. Pharm. 26, 62; Debray, Bl. [2] 2, 14; Heintz, A. 143, 356; Graham, A. 29, 23). The acid salt ZnH₄P₂O₈·2H₂O is obtained in large, triclinic crystals by dissolving ZnO in H₃PO, Aq (Demel, B. 11, 1171).

Double salts. 1. With NH, v. Debray (C. R. 59, 40), Bette (A. 15, 129), Heintz (A. 143, 156), Schweikert (A. 145, 57), Rother (A. 143, 356).—2. With Na, v. Scheffer (A. 145, 53), Ouvrard (C. R. 106, 1729).—3. With K, v. Ouvrard (l.c.).

Zirconium orthophosphates. For double salts with K and Na, v. Ouvrard, C. R. 102,

1422; 105, 30.

PYROPHOSPHATES. Salts derived from H₄P₂O₇. (For Thiopyrophosphates v. Phosphoric SULPHIDE, Reactions, No. 7, p. 147; and for Selenopyrophosphates, v. Phosphoric selenide, p. 145.)

Aluminium pyrophosphate Al₄(P₂O₇)₂. White amorphous pp. by adding AlCl₃Aq to Na₄P₂O₇Aq; soluble NH₃Aq, Na₄P₂O₇Aq, and mineral acids (Schwarzenberg, A. 65, 147; Rose, A. 76, 16; Wittstein, Repert. 63, 224). Forms a double salt with Na (Persoz, A. 65, 170; Pahl, Bl. [2] 22, 122).

Ammonium pyrophosphates. Normal salt (NH₄)₄P₂O₇; by adding excess NH₂Aq to H₄P₂O₇Aq and recrystallising from alcohol. Solution boiled gives the acid salt (NH₄)₂H₂P₂O, (Schwarzenberg, A.65, 141).

Barium pyrophosphate Ba₂P₂O₇.xH₂O. White amorphous solid; by adding Na₁P₂O₇Aq to BaCl₂Aq, or H₄P₂O₇Aq to BaOAq. Heated in steam gives off PH3; gives orthophosphate by heating with alkali carbonates (Hesse, P. 80, 71; Schwarzenberg, l.c.; Birnbaum, Z. [2] 7, 139; Struve, J. pr. 79, 348; Rose, P. 76, 20). Forms

a double salt with Na (Baer, P. 75, 166).

Bismuth pyrophosphate Bi₄(P₂O₇)₃ (Chancel,

C. R. 50, 416).

Cadmium pyrophosphate Cd2P2O2.2H2O (at

100°) (Schwarzenberg, A. 65, 153).

Calcium pyrophosphates Normal salt Ca₂P₂O₂AH₂O₃ by adding Na₄P₂O₃Aq to CaCl₂Aq, Calcium or CaOAq to H₄P₂O₇Aq; sol. in much Na₁P₂O₇Aq. Amorphous, but crystallises from solution in acetic acid or SO₂Aq (Schwarzenberg, A. 65, 145; Rose, P. 76, 16, 20, 230). For action of heat, v. Baer (P. 75, 155). Decomposed by water at 280° to H₃PO₄Aq and Ca₃P₂O₈ (Reynoso, C. R. 34, 795). Partially decomposed by boiling with NaOHAq or fusing with alkali carbonate (Rose). The acid salt CaH2P2O2.2H2O is said to be obtained by decomposing the normal salt by a little H₂C₂O₄Aq (Pahl, Bl. [2] 22, 122). Von Knorre a. Oppelt (B. 21, 769) failed to obtain this salt, but prepared the salt 2CaH₂P₂O, Ca₂P₂O, 6H₂O by adding CaCl₂Aq to $Na_2H_2P_2O_7Aq.$ $Double\ salt\ CaNa_2P_2O_7.4H_2O\ (Baer,\ P.\ 75,$

159).

Chromium pyrophosphate $Cr_*(P_2O_7)_s$. A reddish green pp. by adding $Na_*P_2O_*Aq$ to Cr alum solution; soluble $Na_*P_2O_*Aq$, KOHAq, SO_2Aq , and mineral acids (Schwarzenberg, A. 65, 149).

Cobalt pyrophosphates. Salts having the composition Co₃P₄O₁₃ and Co₅P₁₀O₃₁ are said to be formed by strongly heating luteo-cobalt phos-

phate (Braun, Fr. 3, 468; Gm.-K. 3, 463).

Copper pyrophosphates. The normal salt Cu₂P₂O₇.2H₂O, by adding Na₄P₂O, Aq (not excess) to Cu salt solutions and drying at 100°; greenishwhite amorphous pp., becoming crystalline by boiling in SO₂Aq; soluble Na₄P₂O₂Aq, NH₄Aq, and mineral acids (Schwarzenberg, A. 65, 156; Stromeyer a. Gladstone, Gm.-K. 3, 615). Forms double salts with NH., K, and Na (v. Schwarzenberg, l.c.; Schiff, A. 123, 1; Persoz, A. Ch. [3] 20, 315; Fletimann a. Henneberg, A. 65, 387

Iron pyrophosphates. The ferrous salt is obtained by ppg. FeSO, Aq by Na, P2O, Aq; composition not decided (v. Schwarzenberg, A. 65, 153; Struve, J. pr. 79, 348). The ferric salt Fe₁(P₂O₁)_{3.9}H₂O is a yellowish pp., formed by adding neutral FeCl₂Aq to Na₄P₂O₂Aq (acidified FeCl.Aq gives FePO,; Schwarzenberg). Gladstone (C. J. [2] 5, 435) obtained an allotropic form by heating the ordinary salt in dilute H2SO4Aq

Double salts, with Na (Gladstone, l.c.; Persoz, A. Ch. [3] 20, 815; Fleitmann a. Henneberg, A. 65, 390; Rieckher, N. J. P. 23, 1; Milck,

J. 1865. 263).

Lead pyrophosphate Pb₂P₂O₇. A white amorous powder; by ppg. Pb(NO₃)₂Aq by phous powder; by ppg. Pb(NO₃)₂Aq by Na₄P₂O₄Aq. Obtained in colourless, transparent, rhombic prisms by dissolving PbO in molten KPO, (Ouvrard, C. R. 110, 1333). Soluble excess of Na₄P₂O,Aq, insoluble acetic acid or NH₄Aq; decomposed by boiling water (Stromeyer, Schwarzenberg, Gm.-K. 8, 121; Gerhardt, A. Ch. [3] 22, 506). Forms a double salt with Na (Gerhardt, Lc.; Ouvrard, Lc.). Vol. IV.

Lithium pyrophosphate Li,P,O,.2H,O; by adding Na,P₂O,Aq to LiClAq, dissolving in acetic acid, and ppg. by alcohol (Rammelsberg, B. B. 1883, 18; Merling, J. 1879, 1043). A double salt, Li, NaP₂O₁, is formed in prismatic crystals by dissolving Li, CO₂ in molten NaPO₂ (Ouvrard, C. R. 110, 1333).

Magnesium pyrophosphates. The normal salt Mg2P2O,.3H2O formed by adding is MgSO,Aq to Na,P₂O,Aq (Rose, P. 76, 16, 20), or by dissolving MgO in Na₂H₂P₂O,Aq and heating (Schwarzenberg, Gm.-K. 2, 448). Pp. becomes crystalline by remaining under liquid from which it has been ppd. (Popp, Z. [2] 6, 305), also by boiling with SO₂Aq (Schwarzenberg, *l.c.*). The salt is obtained, without H.O. by strongly heating MgNH₄PO₄. Loses 3H₂O at 100°; fused with soda or heated with conc. H.SO, gives Mg.P.O. (Rose; Weber, P. 73, 137). Easily soluble HClAq and HNO₃Aq; ppd. incompletely by NH₃Aq (Fresenius; Weber). An acid salt, Mg₂H₂P₂O₂, combined with NO₂, is said to be formed by evaporating to dryness with HNO, Aq (Luck, Fr. 13, 255; Campbell, P. M. [4] 24, 380). Forms a double salt with Na (Persoz, A. 65, 169; Schwarzenberg, A. 65, 146; Beer, P. 75, 168).

Manganese pyrophosphates. The normal manganous salt Mn,P,O,.3H,O is a white amorphous powder; by ppg. a salt of Mn by Na,P₂O,Aq (Schwarzenberg, A. 65, 133). Soluble Na,P₂O,Aq (Pahl, Bl. [2] 22, 122). Loses one H₂O at 120° (S., l.c.). The salt without H₂O is formed by strongly heating the hydrated salt, or MnHPO.3H.O., or MnNH.PO.H.O (S., Lc.; Heintz, &c., Gm.-K. 2, 475); white powder, S.G. 3.5847 at 20° (Lewis, Am. S. [3] 14, 281); heated to whiteness in H gives Mn.P.O.; reduced by charcoal to Mn phosphide (Struve, J. 1860. 78). The acid salt MnH₂P₂O₂.4H₂O is said to be formed by action of oxalic acid on the normal salt (Pahl, l.c.). Various double salts are described—with K (Pahl, l.c.; Schjerning, J. pr. [2] 45, 515); with Na (Pahl, Rose; Christensen, J. pr. [2] 34, 41); with Na and NH₄ (v. Otto, J. pr. 2, 418). The acid manganic salt MnHP₂O, is said to be formed by heating MnPO₄ with H₂PO₄Aq to 170°-190° (Christensen, J. pr. [2] 28, 1).

Mercury pyrophosphates. The normal mercurous salt Hg₄P₂O₇.H₂O is a heavy white powder obtained by adding Na, P₂O, Aq to HgNO, Aq (Rose, Gm.-K. 8, 754; Schwarzenberg, A. 65, 133). The normal mercuric salt, Hg₂P₂O₂, is obtained by adding Na_4P_4O , Aq to $Hg(NO_8)_2Aq$ (S., *l.c.*; Reynoso, C. R. 84, 795).

Nickel pyrophosphate Ni₂P₂O₇. A green pp. by adding Na₄P₂O₇Aq to NiSO₄Aq; soluble excess Na₄P₂O₇Aq, NH₄Aq, and acids; obtained crystalline by boiling with SO2Aq; decomposed by water

at 280°-300° (S., l.c.; Reynoso, l.c.).

Potassium pyrophosphates. The normal salt, K₄P₂O₇,xH₂O₇ is obtained by strongly heating K₂HPO₄; also formed by reaction of POCl₅ with KOHAq (Gladstone, C. J. [2] 5, 435). Loses one H₂O at 100°, and is dehydrated at 300°. Solution in water does not change to K2HPO4 on boiling (Schwarzenberg), nor on evaporation (Gladstone), but change is effected by heating to 280° (Reynoso, C. R. 34, 795); boiling with KOHAq produces the ortho-salt. The acid salt, K,H,P,O,, is obtained by treating the normal with acetic acid and alcohol, and evaporating over H2SO. (Schwarzenberg, A. 65, 137). Forms a double \hat{s} alt with $2\hat{K}_{1}(\hat{N}\hat{H}_{1})\hat{H}\hat{P}_{2}\hat{O}_{7}$. $\hat{H}_{2}\hat{O}$ (S., l.c.); also with $\hat{C}r_{2}\hat{H}_{4}(\hat{P}_{2}\hat{O}_{7})_{2}$ (Schjerning, J. pr. [2] 45, 515).

Silver pyrophosphates. Normal salt, Ag. P.O.; a white pp. from AgNO, Aq and Na, P2O, Aq (Clark, Stromeyer, Gm.-K.3, 919). S.G. 5.306 at 7.5° soluble NH, Aq, reppd. by HNO, Aq; boiled with Na₂HPO₄Aq gives Ag₃PO₄. Heated with H₃PO₄Aq at 180° gives the acid salt Ag₂H₂P₂O₇ (Hurtzig a. Geuther, A. 111, 160).

Sodium pyrophosphates. The normal salt, Na₄P₂O₇, is formed by strongly heating Na₂HPO₄; a glass-like solid, becoming opaque on cooling (Clark, N. Ed. P. J. 7, 298); gives Na,PO, and PH, when heated in H (Struve, J. pr. 79, 350); heated with NH₄Cl gives NaCl, NaPO₃, and PCl₃ The hydrated salt, Na, P2O, 10H2O, crystallises from Na, P2O, Aq (Clark, l.c.; von Blücher, P. 50, 542) in monoclinic crystals (Rammelsberg; Haidinger, P. 16, 510; Handl, W. A. B. 32, 250; Schacchi, P. 109, 365; Duffet, C. R. 102, 1827). Crystallises unchanged from Aq with NH₃ or NH₄Ol added. Solution boiled with acids gives Na₃PO₄Aq. Poggiale (J. Ph. [3] 44, 273) gives solubility; 100 parts water dissolve-

	Na P.O.	Na P.O. 10H.O
at 0°	3.0	5.41
10	3.95	6.81
20	6.23	10.92
30	9.95	18.11
40	18.5	24.97
50	17.45	83.25
60	21.83	44.07
70	25.62	52.11
80	30.04	63.40
90	35.11	77.47
100	40.26	93.11

The acid salt Na,H,P₂O,; by heating NaH₂PO₄ to 190° – 204° (Graham, Gm.-K. 2, 169), or by addition o' alcohol to an acetic acid solution of Na, P2O, (Schwarzenberg, A. 65, 139). Crystallises from aqueous solution with 6H2O (Bayer, J. pr. 106, 501; Rammelsberg, B. 16, 21; Duffet, C. R. 102, 1327). When strongly heated gives Na H₂P₄O₁₃ (Fleitmann a. Henneberg, A. 65, 328).

Double salts.—1. $Na_2(NH_4)_2P_2O_7.xH_2O_7$ x=5 or 6 (Schwarzenberg, A. 65, 140, 142; Rammelsberg). — 2. Na, \mathbb{K}_2P_2O , 12H₂O, and Na, \mathbb{F}_2O , Na, \mathbb{H}_2P_2O , 2H₂O (S. Lc.; R., Lc.). Wallroth (Bl. [2] 89, 816) has described double salts with pyrophosphates of Al, Be, Bi, Cd, Ca, Ce, Cr, Co, Cu, Di, Er, La, Mn, Mg, Ni, Th, Yb, Y, Zn, and Zr. For double salts with pyrophosphates of Sn and Ti v. Wunder, J. 1870. 359; 1871. 323. Na,P₂O, seems to combine with B₂O₃ by fusing Na,P₂O, with boric acid (Prinvault, C. R. 74, 1249).

Strontium pyrophosphate, $Sr_2P_2O_7$. A crystalline powder; from $Sr(NO_8)_2Aq$ and Na₄P₂O₇A₂; also by heating SrHPO₄ (S., l.c.). Forms acid salts (von Knorre a. Oppelt, B. 21, 769); also a double salt with Na (Baer, P.

75, 166).
Thallium pyrophosphates. Normal thallous salt, Tl,P,O,; by heating Tl,HPO, (Descloizeaux, 4. Ch. [4] 17, 329). S.G. 6.786. Crystallises

from aqueous solution with 2H₂O (D., l.c.). The acid salt Tl₂H₂P₂O₂,H₂O is formed by heating TlH₂PO₄ to 250°-275° (Lamy, Rammelsberg, Gm.-K. 3, 177).

Thorium pyrophosphates. Double salts with K and Na are described by Ouvrard (C. R.

105, 30).

Uranium pyrophosphates. Double salts with K and Na are described by Ouvrard (C. R. 110, 1333).

Yttrium pyrophosphate. Johnson describes Y₄(P₂O₇)₄, formed by heating Y₂3SO₄ with HPO₄ to 316° (B. 22, 976).

Zinc pyrophosphate, Zn₂P₂O₇. By ppg. Zn salts by Na, P2O, (Stromeyer, Gm.-K. 2, 832; Gladstone, ibid. 3, 16; Schwarzenberg). Forms a

double salt with NH, (Bette, A. 15, 129).

Zirconium pyrophosphates. Double salts with K and Na are described by Ouvrard (C. R.

105, 30)

METAPHOSPHATES. Salts derived from HPO₃. The general methods of preparation, properties, and reactions of the metaphosphates are described at the beginning of this article (p. 106); the divisions into mono-, di-, tri-, tetra-, and hexa- metaphosphates, with the processes by which typical salts of each division are prepared, have also been described already (p. 107). It will suffice here to enumerate the salts, with an indication in each case to which division the salt is assigned by Maddrell, Fleitmann a. Henneberg (cf. Tammann's results regarding Na, di-, and tri- metaphosphates p. 107). T.'s paper in J. pr. [2] 45, 417 should also be consulted.

Aluminium dimetaphosphate, Al.P.O.s. By dissolving Al₂O₂.xH₂O in H₂PO₄Aq, evaporating, and heating to 316° (Maddrell, C. S. Mem. 3, 873; Johnson, B. 22, 976). For hexa-salt v. T.,

J. pr. [2] 45, 417.

Ammonium metaphosphates. (NH₄)₂P₂O₆, from Cu salt by (NH₄)₂SAq; at 200°-250° becomes mono-salt (NH₄)PO₅ (Fleitmann, A. 72, 236). For decay and sends a line of the contract of

236). For deca- and penta- salts v. T., l.c.

Barium metaphosphates. (1)? Mono- salt

BaP₂O₆; by heating BaCO₂ with H₃PO₄ to 316°

(M., l.c.; F., l.c.; Rose, P. 76, 5; Birnbaum, Z.

[2] 7, 139). (2) Di- salt, Ba₂P₄O₁₂xH₂O; from Na or NH, salt and BaCl, Aq (F., l.c.). (3) Trisalt, Ba, P, O_{1,0}xH₂O; from Na salt and BaCl₂Aq (F. a. Henneberg, A. 65, 313). (4) Hexa-salt; from Na salt and BaCl₂Aq (Rose; Graham, P. 32, 68).

Bismuth metaphosphate. ? Tetra-salt; by fusing H.PO, with Bi,O,xH,O (F., l.c.); HPO,Aq, Bi3NO, solution and NH, Aq give a pp. (Persoz, Gm.-K. 2, 832).

Cadmium metaphosphate. ? Tetra- salt (F., l.c.; Persoz, Gm.-K. 3, 54). Double salts with Na and K (Ouvrard, C. R. 106, 1729).

Calcium metaphosphates. A salt is obtained by heating CaHPO₄; for action of HPO₅ on CaCO₅ v. Rose (P. 76, 5), Liebig (A. 11, 262). Di-sālt, Ca₂P₄O₁₂xH₂O; from Na salt and CaCl₂Aq (M., l.c.; F., l.c.). Double salts.—CaAm₂P₄O₁₂:2H₂O (F., l.c.); CaNaP₅O₅ (F. a. H., l.c.) l.c.).

Chromium dimetaphosphate, $Cr_aP_aO_{18}$. By heating CrO_aH_a with H_aPO_a to 816° (M., l.c.; Hautefeuille a. Margottet, C. R. 96, 849, 1042:

Johnson, B. 22, 976).

Cobalt metaphosphates. ? Mono- salt CoP2O4; by evaporating CoSO4 with H3PO4Aq, and heating to 316° (M., l.c.; H. a. M., l.c.). Di- salt Co2P4O12; by heating roseo-cobaltic phosphate (Braun, Ammon. Kobaltverbindungen [Göttingen, 1862]). ? Hexa-salt Co, P, 20, 25; by CoCl, Aq and the Na salt (Rose). Okto-salt (T., l.c.). Double salt with Na (M., l.c.).

Copper metaphosphates. Di- salt Cu.P.O., by heating CuO or Cu(NO₃)₂ with H₃PO₄ to 316° (M., l.c.; F., l.c.); obtained with 8H₂O by evaporating CuSO₄Aq mixed with Na₂P₂O₆ (F. l.c.). Hexa-salt; by ppg. CuCl₂Aq (not CuSO₄Aq) by the Na salt (Rose). Double salts with NH, and Na (F., l.c.; T., l.c.).

Iron metaphosphate. Di-ferric salt Fe2PeO18; by heating FeCl, with H,PO, to 316° (M., l.c.; H. a. M., l.c.; Johnson, B. 22, 976).

Lithium hexametaphosphate. A double $salt Li_2Na_4P_6O_{18}$, is described by Tammann (l.c.).

Lead metaphosphates. Di- salt Pb2P4O12; from Pb(NO₃)2Aq and the Na salt; also by heating H₂PO₄ and PbO (F., l.c.). Tri-salt Pb₃P₆O₁₈; from the Na salt and Pb(NO₂)₂Aq (F. a. H., l.c.). Deka-salt, T. (l.c.). Double salt PbAm2P4O12 (F., l.c.).

Magnesium metaphosphates. ? Mono- salt MgP₂O₈, by heating MgO with H₃PO₄ (M., l.c.). Di- salt Mg₂P₄O_{1:}xH₂O; and tri- salt Mg₂P₄O_{1:}xH₂O; from the corresponding Na salt and MgCl₂Aq (F., l.c.). Hexa- salt Mg₂P₁O₂O₈ (Graham, Gm.-K. 2, 214). Double salts, with NH₄ and Na, v. F. (l.c.); F. a. H. (l.c.); M. (l.c.); Graham (l.c.); Ouvrard (C. R. 106, 1729); Tammann (l.c.).

Manganese metaphosphates. Di- salt manganese metaphosphates. Di- salt Mn₂P₄O₁₂; by heating a Mn salt with H₂PO₄ to 316° (M., l.c.; F., l.c.); with 8H₂O by ppg. (NH₄)₂P₂O₂Aq by excess of MnCl₂Aq (F., l.c.). Tri- salt Mn₂P₂O₁₈.11H₂O; from MnCl₂Aq (not MnSO₄) and Na₂P₂O₁₈. Hexa- salt Mn₂P₁₂O₃₈; by using Na₂P₄O₁₈ (Otto, Rose, Gm.-K. 2, 490; by using Na₂P₄O₁₈ (Otto, Rose, Gm.-K. 2, 490; Lorent J. pr. [2] 45, 417). Double salts, with NH, and Na (v. F., l.c.; Rose, l.c.; T., l.c.).

Mercury metaphosphates. White pps. are formed by adding Na tri- and hexa- metaphosphate to $Hg(NO_3)_2Aq$ (F. a. H., l.c.; Rose, Gm-K.

2, 754).

Nickel metaphosphates. Di-salt Ni₂P₄O₁₂; by heating Ni₂O₃ with H₂PO₄ to 316° (M., l.c.; H. a. M., l.c.). Hexa-salt Ni_eP₁₂O₃₆; by adding NiCl₂Aq to Na_eP₆O₁₆ (Rose). Double salt with Na to Na₆P₆O₁₈ (Rose).

(M., l.c.). Potassium metaphosphates. Mono- salt KPO,; by heating KH,PO, to full redness (M., l.c.; Graham, Darracq, Gm.-K. 2, 32).
S.G. 2.2639 at 14.5° (Clarke, Am. S. [8] 14, 281).
Di-(F., l.c.). Hexa-salt K₀P₀O₁₈ (T., l.c.). Double salt with NH₄ (F., l.c.). Compound with B₂O₃ (Prinvault, C. R. 74, 1249).

Silver metaphosphates. Di- salt Ag,P2O; and tri-salt Ag.P.O.; from AgNO.Aq and the corresponding Na salts (F., l.c.; F. a. H., l.c.). Hexa- salt Ag.P.O.; from AgNO, heated with H.PO, to 316°, or by adding Na.P.O.; Aq to AgNO.Aq (Graham; F., l.c.; T., l.c.; Rose, Berzelius). Double salts with Al, Cr, and Fe

(H. a. M., Lc.).

Sodium metaphosphates. For conditions of formation and general reactions v. p. 106. The following salts have been described: Mono- salt NaPO3 (F., l.c.; M., l.c.; F. a. H., l.c.; Graham, Gm.-K. 2, 170; Jamieson, A. 59, 350; Müller, P. 127, 428); di- salt Na, P.O, 2H, O (F., l.c.); tri- salt Na, P.O, 6H, O (F. a. H., l.c.); tetra- salt Na, P₄O₁₂ (F., l.c.); hexa- salt Na, P₆O₁₈ (F., l.c.; T., l.c.; G., l.c.). For double salts with Na and NH, v. F. (l.c.).

Strontium metaphosphates. ? Mono- salt SrP₂O₆; by heating SrCO₂ with H₂PO₄ (M., l.c.; cf. Rose, P. 76, 8; Scheerer, J. pr. 75, 113). For

a deka- salt, v. T. (l.c.).

Thallium metaphosphate. TlPO₃ said to exist in two forms, one insoluble in water (? disalt), and the other soluble (Lamy, Gm.-K. 2,

Thorium metaphosphate ThP₄O₁₂ (? di- salt, Th₂P₈O₂₄); by heating Th(SO₄)₂ with H₂PO₄ to redness (Johnson, B. 22, 976). For double salts with K and Na, v. Troost a. Ouvrard (C. R. 105, 30).

Uranium metaphosphate. Di-salt U.P.O.

(H. a. M., l.c.).

Zinc metaphosphates. Di-salt Zn₂P₄O₁₂; by heating ZnO with H₃PO₄ to 350° (F., l.c.); with 4H₂O by adding ZnCl₂Aq to (NH₄)₂P₂O₆Aq. For double salts with NH₄ and Na, v. F. a. H. (l.c.); Bette (A. 15, 129).

TETRAPHOSPHATES and DEKAPHOS-PHATES. Salts of the hypothetical acids $H_{e}P_{4}O_{13}$ and $H_{12}P_{10}O_{21}$ (v. pp. 106, 107).

Barium tetraphosphate Ba,P,O13. A heavy crystalline powder; by fusing Na, P4O19 with BaCl₂ (F. a. H., A. 65, 313).

Magnesium tetraphosphate Mg₃P₄O₁₃. pared similarly to the Ba salt (F. a. H., l.c.).

Silver tetraphosphate Ag₆P₄O₁₂. From the Na salt and AgNO₂Aq (Berzelius, P. 19, 331).

Sodium tetraphosphato Na₆P₄O₁₇. By fusing Na₆P₆O₁₈ with Na₄P₂O₇ or Na₂PO₄ (F. a. H., l.c.; Uelsmann, A. 118, 101).

Silver dekaphosphate $Ag_{12}P_{10}O_{21}$. From the Na salt and $AgNO_{2}Aq$ (F. a. H., l.c.; U., l.c.).

Sodium dekaphosphate $Na_{12}P_{10}O_{31}$. A glassy mass; by fusing $Na_{6}P_{6}O_{18}$ and $Na_{4}P_{2}O_{7}$ (F. a. H., *l.c.*). H., l.c.).

PHOSPHENYL CHLORIDE v. PHENYL-dichloro-PHOSPHINE.

PHOSPHENYLIC ACID v. PHENYL PHOS-PRITES

PHOSPHENYLOUS ACID C.H.PHO.OH. Benzene phosphinic acid. [70°]. S. 7.23 in the cold; 211 at 100°. Formed by adding water to C₆H₅PCl₂, and boiling (Michaelis, A. 181, 308; B. 10, 816). Leaflets, v. sol. alcohol. Decomposed by heat into phenyl-phosphine, benzene, and HPO, PCl, forms C,H,PCl,-KA' 2aq.-

Hild.—BaA', 4sq.—PbA',: amorphous pp.

Ethyl ether EtA'. Oil.

Di-ethyl ether C₆H₂P(OEt), (235°).

Formed from C₆H₂PCl, and NaOEt. Liquid with

very powerful odour.

PHOSPHIDES. Compounds of P with one other more positive element. The chief phos phides are those of Al, Sb, As, Ba, Be, Bi, Cd, Ca, Cu, Cr, Au, H, Fe, Pb, Mg, Mn, Hg, Mo, Ni, K, Pt, Ag, Na, Th, Sn, Tl, W, Zn, Zr. (Phosphides of H are described as Phosphorus hydrides under Phosphorus.) Many phosphides are formed by heating together P and the more positive element in an atmosphere of CO₂ or N; phosphides are also produced by heating P with several metallic oxides or with salts in solution; also by heating metals or their oxides in PH₂. Phosphides are generally brittle solids; many react with water or dilute acids, evolving P hydrides; some yield basic phosphates when heated in air. For descriptions of the different phosphides v. the individual elements.

M. M. P. M. PHOSPHINE. PH₃ is sometimes known by this name, v. Phosphorus, hydrides of, p. 135.

PHOSPHINES are the phosphorus analogues of amines, and result from the replacement of hydrogen in phosphuretted hydrogen by alkyl radicles. They may be classified, like the amines into primary, secondary, and tertiary bases, and, in addition, quaternary compounds are known analogous to the salts of compound ammoniums.

Methods of preparation. 1. Action of the haloid derivatives of hydrocarbon radicles on metallic phosphides. It was by this reaction that Paul Thénard (C. R. 21, 144; 25, 892) discovered the first members of the phosphine group in 1843-47. He experimented with methyl chloride and calcium phosphide, and apparently isolated trimethyl-phosphine, (CH₃)₂P₄ as an inert solid, and a substance (CH₃)₄P₂ analogous to cacodylas aspontaneously inflammable liquid, boiling at 250°, very explosive, poisonous, and unstable. Thénard recognised the relationship of trimethyl-phosphine to ammonia, and predicted the existence of the then undiscovered organic compounds of nitrogen and antimony.

In 1855 Hofmann a. Cahours (T. 1857; A. Ch. [3] 61, 5) investigated the action of methyl iodide on sodium phosphide, and obtained tetramethyl-phosphonium iodide in addition to trimethyl-phosphine and the phosphorised cacodyl of Thénard. They found that the method was dangerous, and furnished mixtures the separation of which presented enormous

difficulties.

Berlè (J. pr. 66, 73) about the same time obtained very small quantities of triethylphosphine by the action at a very high temperature of ethyl iodide on sodium phosphide obtained by heating sodium and phosphorus together in rock oil.

Cahours in 1859 (C. R. 49, 87; J. 1859, 430) prepared tetrethyl-phosphonium iodide by the action of ethyl iodide on crystallised zinc phosphide (obtained by heating the metal in

phosphorus vapour at 180°).

In 1882 Letts a. N. Collie (Tr. E. 30 pt. i. 181) investigated the action of benzyl chloride on sodium phosphide, and found that tetra-benzyl-phosphonium chloride was produced in abundance.

2. Action of organo-metallic bodies on trichlorids of phosphorus. The action of zinc
alkyls a phosphorus trichloride was first investigated by Hofmann a. Cahours (A. Ch. [3]
51), and was further studied by Hofmann (A. Ch.
[8] 62; 63 · 64). By this method tertiary phosphines are exclusively formed: $3R_2Zn + 2PCl_2$ $= 2R_2P + 3ZnCl_2$. It is necessary to treat the
product of the reaction with caustic potash, in

order to decompose the compound of the phosphine with zinc chloride. By this reaction Hofmann a. Cahours obtained PMe₃ and PEt₄, and showed that they resemble the corresponding amines in many respects, especially in the readiness with which they combine with alkyl iodides to give quaternary compounds. On the other hand, they proved that, unlike the amines, tertiary phosphines readily combine with oxygen to give very stable compounds of the general formula R₃PO.

3. Action of alcohols on phosphonium iodide; and 4. Action of alkyl iodides on phosphonium iodide and oxide of zinc (Hofmann's methods). In the year 1871 Hofmann again took up the study of the phosphines (B. 4, 205), and by the action of alkyl iodides on phosphuretted hydrogen obtained not only tertiary and quaternary compounds, but also primary and secondary bases. Hofmann took advantage of the fact that phosphonium iodide when heated with alcohol yields phosphuretted hydrogen, ethyl iodide, and water. When phosphonium iodide (1 mol.) is heated with alcohol (3 mols.) for 8 hours at 180° the crystalline product is a mixture of PEt,HI and PEt,I, and, on addition of caustic soda, triethyl-phosphine separates as a clear liquid, while the solution gives, on evaporating, beautiful crystals of tetrethyl-phosphonium iodide. If the tubes are heated for four hours only, two layers of liquid are visible on cooling, and the tubes show great pressure when opened. It may be taken for granted, then, that the reaction occurs in two phases, in the first of which EtI is liberated. which then acts upon PH, in the same way as it acts on ammonia. Hofmann employed this method successfully in the methyl (B. 4, 209), ethyl (B. 4, 205), propyl (B. 6, 292), butyl (B. 6, 296), and amyl (B. 6, 297) series. Hofmann Hofmann (B. 4, 372) endeavoured to prepare the primary and secondary bases according to the equations:

 $PH_4I + C_2H_6O = C_2H_5PH_2HI + H_2O$ $PH_4I + 2C_2H_6O = (C_2H_5)_2PHHI + 2H_2O$, by altering the proportions of phosphonium iodide and alcohol, but without success, the tertiary base being produced alone or mixed with the quaternary compound, while with the proportions required for the second equation the

tubes invariably exploded.

In the meantime Drechsel a. Finkenstein (B. 4,352) believed that they had succeeded in obtaining the primary bases by saturating EtI or MeI with PH3, and allowing the solutions to remain for some time at the ordinary temperature, or by heating them at 100°, and also by heating an ethereal solution of ZnI2 saturated with PH, together with MeI. Hofmann (B. 4, 372) repeated these experiments, and showed that only tertiary and quaternary derivatives were formed. The idea then occurred to him of heating the alkyl iodide with phosphonium iodide in presence of a metallic oxide, with the happiest results; for on heating a mixture of phosphonium iodide (2 mols.) with ethyl iodide (2 mols.) and zinc oxide (1 mol.) in sealed tubes at 150° for from six to eight hours, a complete reaction occurred, the tubes when cold containing a crystalline mass consisting exclusively of the hydriodates of the primary and secondary bases, the former being the chief product, while the latter was only formed in small quantities. The addition of water to the product of the reaction liberated the primary phosphine alone, which was distilled off in a hydrogen atmosphere, while the secondary phosphine was subsequently set at liberty by the action of an alkali. By means of this general method Hofmann prepared methyl (B. 4, 430), ethyl (B. 4, 605), propyl (B. 6, 292), butyl (B. 6, 296), amyl (B. 6, 297), and benzyl

(B. 5, 100) phosphines.

Michaelis's methods. Hofmann's methods, although of excellent service for obtaining the phosphines of those radicles which form alcohols, could not be employed in the preparation of phosphines containing purely aromatic radicles (B. 5, 100). Michaelis, on the other hand, not only succeeded in obtaining all the phenyl-phosphines, but also in discovering a fairly general method for the production of primary phosphines. The substance forming the starting-point for the preparation of phosphines by these methods is trichloride of phosphorus. One atom of chlorine is first replaced in that body by one or other of the following processes:-

(A) The mixed vapours of a hydrocarbon and the trichloride are repeatedly passed through a red-hot tube. Thus when benzene is employed 'phosphenyl' chloride is obtained: $PCl_3 + C_6H_6 = (C_6H_5)PCl_2 + HCl$.

(B) A mercury alkyl is heated with the trichloride under pressure : PCl₂ + (C₂H₃)₂Hg

= (C₂H₅)PCl₂+Hg(C₂H₅)Cl. (c) By digesting a hydrocarbon with the trichloride and aluminum chloride, the reaction being the same as (1). Other substances besides hydrocarbons yield substituted phosphorus chlorides when submitted to this reaction. Thus a mixture of acetone, PCl2, and aluminium chloride react spontaneously, according to the

equation:

 $2(CH_s)_2CO + PCl_3 = 2HCl + (CH_3.CO.CH_2)_2PCl.$ By means of these different reactions Michaelis and his pupils have obtained a considerable number of substituted phosphorus chlorides, among which are: Phenyl phosphoruschloride by methods 1, 2, and 3 (B. 6 [1873], 601; 8 [1875], 922; 12 [1879], 1009). Tolyl phosphorus-chloride by method 3 (B. 13, 653). Xylyl phosphorus-chloride by method 3 (A. 212, 203, 209). Ethyl phosphorus-chloride by method 2 (B. 13, 2174). Propyl phosphorus-chloride by method 2 (B. 13, 2174). Naphthyl phosphorus-chloride by method 2 (B. 9, 1051). Acetonyl phosphorus-chloride by method 3 (B. 17, 1273). The substituted chlorides resemble PCl₃ in properties. As a rule they are fuming liquids, combining readily with chlorine to give solid compounds analogous to pentachloride of phosphorus; treated with water, they yield phosphinous (phosphinic) acids. Thus phenyl phosphorus-chloride gives phenyl phosphinous (benzene phosphinic) acid: $(C_4H_5)PCl_2 + 2H_2O = (C_6H_5)PH_2O_2 + 2HCl$. The products of addition which they form with chlorine react with water to give phosphinic (phosphonic) acids: (C₆H₅)PCl₄+3H₂O=(C₆H₃)PH₂O₅+4HCl. Michaelis was unable to obtain more than mere traces of the primary phosphine by the action of nascent hydrogen on the substituted chlorides. When gaseous HI is passed into the chloride,

chlorine is gradually displaced by iodine: $C_dH_sPCl_2 + 3HI = C_cH_sPl_2HI + 2HCl$. From this iodide Michaelis obtained phenyl-phosphine by the action of alcohol: 8C,H,PI,HI+9C,H,O $= C_6 H_a P H_2 + 2 C_6 H_a P O_2 H_2 + 3 H_2 O + 9 C_2 H_a I$. later experiments he employed a simpler method, namely the destructive distillation of the phosphinous (phosphinic) acid which, as already stated, is produced by the action of water or alcohol on the substituted chloride: 3C,H,PO,H, $= C_6H_5PH_2 + 2C_6H_5PO_3H_2$. From phenyl phosphorus chloride Michaelis obtained diphenylphosphine by the following reactions: (1) The chloride is digested with mercury diphenyl at

 $(C_gH_h)PCl_2 + Hg(C_gH_h)_2 = (C_gH_h)_2PCl + HgCl(C_gH_h)_2$ (B. 8, 1304); or it is heated for some time at 280°, when the following reaction occurs: $2(C_6H_b)PCl_2 = (C_6H_b)_2PCl + PCl_3.$

(2) Diphenyl phosphorus chloride when heated with water or dilute soda solution decomposes in the following manner: $2(C_aH_b)_2PCl + 2H_2O = (C_aH_b)_2PH + (C_aH_b)_2HPO_2 + 2HCl$ (Michaelis a. Gleichman, B. 15, 801). He also obtained the tertiary base; at first by acting upon a mixture of phenyl-phosphorus chloride and bromo-benzene with sodium: $(C_oH_s)PCl_2 + 2C_oH_sBr + 3Na$ = $2NaCl + NaBr + (C_oH_s)_3P$, but later this method was modified in a remarkable way, by substituting for phenyl phosphorus chloride, phosphorus chloride alone, the reaction occurring quite easily and very energetically at ordinary temperatures according to the equation:

 $PCl_a+3(C_aH_a)Br+6Na=3NaCl+3NaBr+(C_aH_a)_aP$.

5. Action of alcohols on a mixture of phosphorus and phosphorus iodide. Letts a. Blake (Tr. E. pt. 2) have shown that very remarkable reactions occur when benzyl alcohol acts upon a mixture of phosphorus and phosphorus iodide (in the proportions employed for obtaining phosphonium iodide by the action of water), whereby a number of the phosphorus derivatives of benzyl are obtained. The action occurs spontaneously at ordinary temperatures, and, although its mechanism was found difficult to investigate, the following equations account for the formation of the different products:-

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(1) PI_2 + C_1H_1OH + H_2O = (C_1H_2)H_2PO_2 + HI + I

(2) PI_2 + C_1H_1OH + 2H_2O + I = (C_1H_2)H_2PO_3 + 3HI

(3) PI_2 + 2C_1H_1OH = (C_2H_2)_2HPO_2 + HI + I
(4) PI_2 + 8C_1H_1OH + HI = (C_1H_1)_2PO + 2H_2O + 3I

(5) PI_2 + 4C_1H_1OH + 4HI = (C_1H_1)_2PI + 4H_2O + 5I
                                                           = H_{s}PO_{s} + HI + I
(6) PI_2 + 2H_2O
(7) PI<sub>2</sub> + 4H<sub>2</sub>O + 3I
(8) 2HI + C<sub>2</sub>H<sub>2</sub>OH
                                                           = H_3PO_4 + 5HI
= C_7H_8 + H_4O + I_3
(9)P+L
                                                           - PI.
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All the products indicated by the above equations were obtained in quantity, especially benzyl phosphinic acid, di-benzyl phosphinic acid, and tri-benzyl phosphine oxide, and the method is undoubtedly the easiest and best for the preparation of those substances. Whether other alcohols behave in a similar manner to benzyl alcohol with a mixture of phosphorus and its iodide has not yet been ascertained.

General proporties of the primary phosphines. The following primary bases have been obtained:

Nan	ae	Condition	Boiling-point
Methyl-pho		Gas	-14°
Ethyl	,, 2	Liquid	25°
Iso-propyl	,, 8	,,	41°
Iso-butyl	,, 4	,,	62°
Iso-amyl	,, 5	**	106°
n-Octyl	,, 6	"	184°-187°
Phenyl	,, 7	,,	160°-161°
Benzyl	,, 8	,,	180°-183°
p-Tolyl	" 9	Solid at +4°	178°

- Hofmann, B. 4, 209.
 Hofmann, B. 6, 292.
 Hofmann, B. 6, 297.
 Hofmann, B. 6, 297.
 Michaelis, B. 7, 6, 1688.
 Hofmann, B. 5, 100; Letts a. Blake, Tr. E. 35 pt. 2.
 Michaelis a. Panek, A. 212, 233.

Most of the primary bases are liquids, insoluble in water, but soluble in ether. They fume in the air, and grow very hot, their vapour igniting spontaneously at times. The products of this oxidation appear to have been investigated only in a few cases, and chiefly in the aromatic series. Phenyl phosphine (Michaelis, B. 10, 807) and tolyl phosphine (Michaelis a. Panek, A. 212, 234) both absorb a molecule of oxygen, and are converted into phosphinous (phosphinic) acids, which are monobasic, and therefore probably have the constitution RPH(OH):O. These acids are readily decomposed by heat, giving the primary phosphine, and the corresponding phosphinic (phosphonic) acid 3RPH2O2 = RPH2+2RPH2O3, a reaction analogous to that which gives rise to phosphoretted hydrogen and phosphoric acid when hypophosphorous and phosphorous acids are heated. Letts and Blake (Tr. E. 35, pt. 2) find that benzyl phosphine yields a mixture of benzyl phosphinous (phosphinic) and benzyl phosphinic (phosphonic) acids with some phosphoric acid. Submitted to the action of strong nitric acid, many of the primary phosphines (and probably all) absorb three atoms of oxygen, and are converted into phosphinic (phosphonic) acids. This has been shown by Hofmann (B. 5, 110) to be the case in the methyl, ethyl, isopropyl, isobutyl, and isoamyl series.

The phosphinic acids are solid substances, which, with the lower members of the fatty series. can be distilled unchanged (Hofmann, B. 6, 303). But in some other instances a different reaction occurs: thus phenyl phosphinic (benzene phosphonic) acid when heated slowly to 200° gives a pyro-acid, while when rapidly heated to 250° it decomposes into benzene and metaphosphoric acid: C₈H₅PH₂O₅ = C₆H₆ + HPO₅. The phosphinic (phosphonic) acids are all dibasic, and no doubt have the structure RPO(OH)₂. All the primary phosphines have distinct alkaline properties. They combine readily with hydracids forming crystalline compounds, which can, as a rule, be volatilised (with dissociation more or less complete), and which resemble the compounds of phosphoretted hydrogen in being instantly decomposed by water with liberation of the phosphine. The hydrochlorides combine with chloride of platinum to give chloroplatinates. The salts of the primary bases with oxyacids have been scarcely at all investigated. The action of halogens on primary phosphines has not been sufficiently investigated. Methyl and ethyl

phosphine take fire when they come in contact with chlorine or bromine (Hofmann, B. 4, 433, 609). Benzyl-phosphine yields with bromine a mixture of its own hydrobromide and substitution-products (Letts a. Blake, Tr. E. 35, pt. 2).

Sulphur acts on the primary bases. With the methyl and ethyl derivatives, compounds have been obtained but not investigated (Hofmann, B. 4, 433, 610). With phenyl phosphine sulphur acts slowly in the cold, rapidly at a high temperature (Michaelis, B. 10, 810). Two substances are produced, one, a thick liquid, soluble in ether, having the composition (C,H,)PH,S, the other a crystalline product to which Michaelis assigns the formula $(C_6H_5P)_3S$. The first of these bodies decomposes when heated in the following manner:

 $2C_{6}H_{5}PH_{2}S = C_{6}H_{5}PS + C_{6}H_{5}PH_{2} + H_{2}S.$ When benzyl phosphine is warmed with sulphur the following reaction occurs (Letts a. Blake, l.c.):

 $2C_{7}H_{7}PH_{2} + 6S - (C_{7}H_{7})_{2}P_{2}S_{5}H_{2} + H_{2}S$ and the resulting pyro-benzyl-thiophosphinic acid when boiled with water is decomposed, yielding mono-thio-benzyl phosphinic acid:

 $(C_2H_1)_2^2P_2S_1H_2 + 4H_2O = 2C_1H_1PS_1OH_2 + 3H_2S_1$ In view of the analogies existing between nitrogen and phosphorus, considerable interest is attached to the action of carbonyl chloride and bisulphide of carbon on the primary phosphines. Bisulphide of carbon acts upon both methyl and ethyl phosphine (Hofmann, B. 4, 433, 610), but the products have not been in-Michaelis a. Dittler (B. 12, 338) vestigated. have studied the action of both reagents on phenyl phosphine. When carbonyl chloride is passed slowly into that substance, an energetic reaction occurs in the following manner

 $2COCl_2 + C_6H_5PH_2 = C_6H_5PCl_2 + 2CO + 2HCl.$ Phenyl phosphine and bisulphide of carbon act upon each other when heated in a sealed tube at 150°, and sulphuretted hydrogen is liberated. The product of the reaction is a resinous body (C_sH_sPHCS)₂S, and the reaction itself proceeds according to the equation

 $2C_aH_aPH_2 + 2CS_2 = (C_aH_aPHCS)_3S + H_2S.$ Michaelis a. Dittler were not successful in their attempts to prepare a phosphorised mustard oil from this compound.

They were equally unsuccessful in obtaining a phosphorised carbylamine by the action of chloroform and caustic potash on phenyl phosphine. It is true that a reaction occurs, but its course is completely different from that which takes place with an amine, viz.

 $C_6H_aPH_2 + 4KHO + CHCl_8 =$ $\mathbf{C_6H_3PHKO_2} + 3KCl + CH_3OH + H_2O.$

It thus appears that, in their behaviour with carbonyl chloride and a mixture of caustic potash and chloroform, primary phosphines behave in an entirely different manner from the corresponding amines, though there is a certain degree of analogy as regards the action of both on bisulphide of carbon. This difference is no doubt due to the strong affinity of phosphorus for electronegative elements, such as the halogens, oxygen, and sulphur. Chloracetic and bromacetic acid react with benzyl phosphine, and chloroformic ether appears to form a product of addition (Letts a. Blake, l.c.). Probably other phosphines act in a similar manner with these reagents.

Primary phosphines readily combine with alkyl iodides to give hydriodides of secondary phosphines.

General properties of the secondary phosphines. The following secondary phosphines have been obtained:

Name		Condition	Boiling-point
Dimethyl phosphin	e 1	Liquid	25°
Di-ethyl ,,	2	,,	85°
Di-isopropyl ,,	8	"	118°
Di-isobutyl ,,	4	,,	153°
Di-isoamyl ,,	5	,,	210°-215°
Di-phenyl ,,	6	,,	280° (about) 78°-80°
Methyl-isopropyl,,	7	77	78°_80°
Iso-propyl,			
isobutyl "	8	,,	139°-140°
• "			Decomposes
Di-benzyl "	•	?	when boiled

- ' Hofmann, B. 4, 610.
- Ibid. B. 6, 294
- Ibid. B. 6, 298. Ibid. B. 6, 295.
- · Letts a. Blake, l.c.
- ⁸ Hofmann, B. 4, 433.
- 4 Ibid. B. 6, 296
- Michaelis, B. 15, 801 A.
 Hofmann, B. 6, 300.

All the secondary phosphines obtained as yet are liquids, having a powerful odour. They are insoluble in water, but soluble in ether &c.

They have as a rule a strong attraction for oxygen, fuming and growing hot on exposure to the air, and often inflaming spontaneously. In some cases they appear to have even a greater affinity for oxygen than the primary bases. This is so according to Hofmann with all the secondary phosphines of the fatty series which he obtained. But apparently it is not the case with diphenyl phosphine. The products of this diphenyl phosphine. spontaneous oxidation do not appear to have been examined except in the case of dibenzyl phosphine, which yields dibenzyl phosphinic (phosphonic) acid and possibly the oxide also (C,H,),PO (Letts a. Blake, l.c.). But the products of their oxidation by nitric acid have been investigated, chiefly by Hofmann (B. 5, 104; 6, 303). These are in all cases phosphinic acids, R₂PHO₂, which are monobasic, and no doubt have the constitution R₂P(OH):O.

These acids are probably also produced when the chlorides R₂PCl are oxidised by nitric acid. Such is at least the case with (C_eH_b)₂PCl. Some of them can be distilled unchanged, e.g. dimethyl phosphinic acid; others, e.g. diphenyl and dibenzyl phosphinic acid, lose water, and give pyro-

Secondary phosphines combine with acids, the resulting salts being far more stable than those of the primary bases. Thus in most cases they are not decomposed by water, though some are (e.g. salts of diphenyl phosphine). paratively little is known regarding secondary phosphines, and very few of their compounds have been investigated.

Sulphur acts upon them, and in the case of diethyl phosphine, according to A. W. v. Hofmann a. Mahla (B. 25, 2436), triethyl phosphine sulphide and diethyl dithiophosphinic acid. PEt₂SSH are formed, together with a third substance which probably has the composition PSEt, S.S.S.PSEt. Bisulphide of carbon also acts upon them, but the products do not appear to have been investigated. They readily combine with alkyl iodides, giving hydriodides of tertiary phosphines.

General properties of tertiary phosphines. The following tertiary bases have been obtained:

Name	Condition	Boiling-point
Trimethyl phosphine	Liquid	40°-42°
Tri-ethyl ,,	2 1	1270
Tri-isopropyl "	3 ,,	?
Tri-isobutyl ,,	4 "	215°
Tri-isoamyl ,,	4 ",	about 300°
Tri-phenyl ,,	6 Solid	above 860°
Tri hanzyl	fia	
Ethyl-isopropyl-	"	"
isobutyl "	7 Liquid	about 190°
Methyl-diphenyl ,,	8 -	284°
Ethyl-diphenyl ,	9 "	293°
Di othel shonel	,,	220°
Di mathal athal	"	83°_85°
Di atherl motherl	2 "	110°-112°
Di-ethyl-propyl ,,	. "	146°-149°
Di sthelissonel	4 "	185°-187°
Di othel hongel	5 "	252°-255°
Fthel dihoneul	8 "	320°-380°
Dimethal m tolal 1	7 "	210°
This add and a dealers I	8 "	240°
Dimothel velel	9 "	230°
	"	260°
Diethyl xylyl ,, 2	" " 	200

- ** Hofmann a. Cahours, A. Ch. [8] 51, 85.

 * * Hofmann, B. 6, 292, 304.

 * Michaelis, B. 15, 801.

 * Michaelis a. Link, A. 107, 210.

 * Michaelis a. Link, A. 17, 210.

 * Michaelis a. Link, A. 181, 80.

 * Official Structure 1888, 714. Michaelis, A. 181, 845.

17 10 10 20 Czimatis, B. 15, 2014. The tertiary phosphines hitherto obtained are, with the exception of triphenyl and tribenzyl phosphine, liquids at ordinary temperatures, having a powerful odour. They are insoluble in water, but soluble in ether &c. As a rule, they oxidise rapidly in contact with the air, fuming and growing hot, and in some cases igniting spontaneously. The product of this oxidation is a tertiary phosphine oxide of the formula R.PO, and, no doubt, of the constitution R.P:O.

The final products of the oxidation of phosphuretted hydrogen, and of primary, secondary, and tertiary phosphines, are therefore respec-tively: H₂PO₄, RH₂PO₂, R₂HPO₂, R₃PO, the amount of oxygen absorbed by the phosphine decreasing in a regular manner as the series is ascended.

The oxides of tertiary phosphines are solid stances of remarkable stability. They can substances of remarkable stability. in the majority of cases be distilled, and even boiled with nitric acid, without change. By no means as yet discovered can they be reduced. Hydracids combine with them, and they give crystalline compounds with a number of metallic salts, such as the chlorides of platinum, zinc, mercury, iron, cobalt, &c., also in some cases with chloride of acetyl, bromine, and sulphur.

Tertiary phosphines also combine with the elements of the sulphur group, forming compounds analogous to the oxides.

The salts of tertiary phosphines are readily obtained by dissolving the bases in acids. They are stable, and are not, as a rule, decomposed by water. Their compounds with hydracids have been chiefly studied; those containing oxyscids have not been investigated (with very few exceptions). The haloid salts dissociate to a greater or less extent on heating. Their hydrochlorides combine with chloride of platinum to give chloroplatinates of normal composition.

Some of the tertiary phosphines combine with chloracetic acid to give hydrochlorides of phosphorised betaines. At present only two or three of these substances have been obtained—trimethyl phosphorus betaine by Meyer (B. 4, 734), the corresponding ethyl derivative by Hofmann (Pr. 11, 530), and in addition to these two the closely allied compound, tri-methyl phosphorus benzo-betaine hydrochloride, by Michaelis a. Czimatis (B. 15, 2018), (CH₃), PCl.C₂H₄.COOH.

The compounds of these phosphorised betaines are stable and well-defined substances. Letts (Tr. E. 30, pt. 1, 285) has investigated the reactions and decompositions of the ethylated body,

which are of some interest.

The hydrate and the salts of this betaine lose carbonic anhydride when heated, and give rise to the hydrate or salt of methyl-tri-ethyl-phos-

phonium

 $(C_2H_s)_sPX.CH_2.COOH = CO_2 + (C_2H_s)_sPX.CH_s$, a reaction which is entirely analogous to that occurring when the corresponding sulphur compounds (thetines) are heated: $(CH_s)_2SX.CH_2.COOH = CO_2 + (CH_s)_2SX.CH_s$. While it is perfectly different from that which the true (nitrogen) betaines experience, as they either dissociate into the original trialkyl-amine and the group $X.CH_2.COOH$ (or the products of its decomposition), or distil unchanged (Brühl, A.177, 214).

Treated with caustic potash, all the salts of tri-ethyl phosphorus betaine yield tri-ethyl phosphine oxide: (C₂H₂)₃PX.CH₂.COOH+2KHO

 $= (C_2H_3)_8PO + KX + CH_3 \cdot COOK + H_2O.$

Several of the tertiary phosphines combine directly and energetically with a molecule of bisulphide of carbon to give highly characteristic compounds, usually of a red colour, and possibly

having the constitution, R₃P<\(\frac{5}{C:S}\). So characteristic and so readily formed is this compound in the case of tri-ethyl phosphine, that its production may be employed as a test either for bisulphide of carbon or for the phosphine itself. As yet these (bisulphide) compounds have been obtained only with methyl, ethyl, and iso-propyl phosphine, and with those of the aromatic phosphines containing ethyl or methyl groups.

According to Czimatis (B. 15, 2016) these mixed phosphines combine very easily with bisulphide of carbon if they contain methyl, the readiness with which combination occurs diminishing, however, in proportion to the molecular weight of the aromatic radicle, while, if they contain ethyl, combination occurs only slowly and with difficulty. Hofmann (Tr. 1860, 431) has somewhat exhaustively studied the compound of tri-ethyl phosphine and the bisulphide, which forms with explosive violence. Among its properties are the following: It is insoluble in water, difficultly soluble in ether, but easily dissolves in hot alcohol, from which it separates on cooling in red needles like chromic anhydride. From an ethereal solution it is deposited by spontaneous evaporation in large deep-red monoclinic crystals exhibiting dichroism, which melt

at 95° and volatilise at 100°. It is soluble in strong hydrochloric acid, and if the solution is mixed with platinic chloride, a yellow amorphous compound is produced, $2(C_1H_3)_3PCS_2_3PtCl_4$. When heated with silver oxide or nitrate, it is decomposed as follows: $(C_2H_3)_3PCS_2 + 2Ag_2O = Ag_2S + Ag_2 + CO_2 + (C_2H_3)_3PS$, and moist air produces a similar change. But if is heated with water to 100°C., the following reaction occurs:—

 $\begin{array}{l} 4(C_{n}H_{n})_{n}PCS_{n} + 2H_{n}O \\ = 2(C_{n}H_{n})_{n}PS + (C_{n}H_{n})_{n}PO + (C_{n}H_{n})_{n}(CH_{n})POH + 3CS_{n}. \end{array}$

Heated with sulphuretted hydrogen, it suffers the following change: $3(C_2H_3)_3PCS_2 + H_2S = 2(C_2H_3)_3PS + (CH_2S)(C_2H_3)_3PCS_2 + CS_2$.

The action of halogens upon tertiary phosphines has not been very fully studied. Probably direct addition would occur in all cases. This has been proved to take place with tri-ethyl phosphine if the halogen is allowed to act very gradually upon it. The chloride (C₂H_a)₃PCl₂ thus obtained is crystalline, melting at 100° and volatilising readily, though its boiling-point is high. Similar compounds of bromine and iodine have been obtained.

Compounds of tri-methyl and tri-ethyl phosphine with mustard oils are formed easily, and give crystalline hydrochlorides. They, no doubt,

have the constitution, S:C:NR:PR3.

[Note.—Some of the aromatic tertiary phosphines, especially tri-phenyl phosphine, have properties which differ materially from those of other tertiary phosphines. Thus tri-phenyl other tertiary phosphines. Thus tri-phenyl phosphine is a crystalline solid having scarcely any odour, and it does not oxidise spontaneously. It is remarkably stable, and is not attacked by chlorine even when heated. The hydriodide and hydrochloride are formed when it is dissolved in the warm concentrated hydracids, and are crystalline, but on adding water they dissociate. By treating the phosphine with bromine and an alkali, or by oxidising it with hydrochloric acid and chlorate of potash, the hydrate $(C_eH_s)_sP(OH)_2$ is obtained as a crystalline solid. This when heated to 100° readily loses water, and is converted into the oxide, a substance which is not acted upon by bromine, oxygen, sulphur, &c. By dissolving the phosphine in fuming nitric acid a nitrate of the formula (C,H,),P(NOs)2 is obtained.]

Tertiary phosphines, apparently without exception, unite with alkyl iodides to form phos-

phonium salts.

General Properties of Quaternary Compounds (Phosphonium Salts). So many of these bodies have been obtained that a list appears inadvisable. It would include derivatives of the series C_nH_{2n-7} , and a large number of mixed phosphoniums containing various radicles, among which are vinyl, allyl, and ethylene.

The phosphonium salts are the most stable of all organic phosphorus compounds. None are decomposed by water, and most of them can be obtained readily in the crystalline state by

evaporating their solutions.

As a rule, they are soluble in water and in alcohol. They are readily prepared from their iodides, either by double decomposition with a silver salt, or by first obtaining their hydrates (by the action of moist oxide of silver), and

subsequently neutralising the solution with the

The hydrates R.POH are solid substances, having a powerful alkaline reaction and many properties similar to those of an alkali. Indeed, in the case of tetrethyl phosphonium hydrate, the only remarkable point of difference between it and caustic potash (so far as its reactions with metallic salts &c. are concerned) is that, when added to a zinc or aluminium salt, the zinc or aluminium hydrate, which is at first precipitated, is insoluble in an excess. Phosphonium hydrates are decomposed when heated, and in some cases, when their solutions are boiled or at the moment of production, into a tertiary phosphine oxide and a hydro-carbon,

$$R_4POH = R_3PO + R - H.$$

The action of heat upon the salts of the phosphoniums has been investigated in a number of cases, partly by Letts and N. Collie (Tr. E. 30, part 1, 213; P. M. August 1886), and partly by the latter chemist alone.

As regards the haloid salts, the chlorides decompose almost quantitatively into a hydrocarbon and a tertiary phosphine hydrochloride (Collie), furnishing an excellent method for retrograding from quaternary to tertiary

'When the phosphonium chloride contains several ethyl groups, then if more than one of the latter is present, ethylene is always formed, e.g.

(C₂H₄)₃(C₇H₇)PCl = (C₂H₄)₂(C₇H₇)P.HCl + C₂H₄. But when only one ethyl group is present, then, although ethylene is still formed, two decompositions occur, e.g. (1)

$$2(C_2H_5)(CH_3)_3PCl = 2(C_2H_5)(CH_3)_2P.HCl + C_2H_4$$
(2) $(C_2H_5)(CH_3)_3PCl = (CH_3)_3P.HCl + C_2H_4$.

If we compare the decomposition by heat of phosphonium chlorides with the decomposition of any of the compound ammonium salts, it must be with the hydroxides and not with the corresponding chlorides' (Collie, C. J. 1888, 636, 714), e.g.

$$\begin{array}{ll} (C_2H_3)_4PCl &= (C_2H_5)_5P.HCl + C_2H_4 \\ (C_2H_5)_4NOH &= (C_2H_5)_5N + C_2H_4 + H_2O \\ (C_2H_5)_4(C_7H_7)P(OH) &= (C_2H_5)_5PO + C_7H_5 \\ (C_2H_5)_6(CH_3)NCl &= (C_2H_5)_5N + CH_5Cl. \end{array}$$

The effect of heat on phosphonium salts derived from oxyacids is completely different. In the case of the ethyl series at all events, they suffer, as a rule, at least two, and occasionally three, different and distinct decompositions. In one of these the molecule splits up into three new groups, consisting respectively of carbonic anhydride, a (paraffin) hydrocarbon, and the tertiary phosphine. In the other, two hydrocarbons are formed—namely, an olefine and a paraffin—in addition to carbonic anhydride and the tertiary phosphine. Whilst in the third, a totally different change occurs, in which only two products are formed-namely, the oxide of the tertiary phosphine and a ketone,

(1)
$$\mathrm{Et_3P} < \frac{\mathrm{C_2H_3}}{\mathrm{OOC_1R}} = \mathrm{Et_3P} + \mathrm{CO_2} + \mathrm{C_2H_3R}.$$

(8)
$$\text{Et}_{a}\text{P} < \frac{\text{i}C_{2}H_{a}}{\text{O}: \text{OCH}} = \text{Et}_{a}\text{PO} + C_{2}H_{3}.\text{CO.R.}$$

It is possible, if not indeed probable, that the third reaction occurs subsequently to the first, and that it really depends upon the reducing action of the triethylphosphine upon the carbonic anhydride, at the high temperature at which the decomposition usually occurs, whereby carbonic oxide is liberated, which combines with the hydrocarbon radicle in statu nascendi, forming

$$\operatorname{Et_3P} + \operatorname{CO_2} + (\operatorname{Et}) + (\operatorname{R}) = \operatorname{Et_3PO} + \operatorname{EtCOR}$$

(or $\operatorname{Et_3P} + (\operatorname{OCO} - \operatorname{R}) + (\operatorname{Et}) = \operatorname{Et_3PO} + \operatorname{EtCOR}$).

If we merely consider the third kind of decomposition alone, it appears to be, to a certain extent, analogous to the decomposition which a sulphine compound suffers when heated, the difference depending on the greater attraction which phosphorus has for oxygen, compared with that of sulphur for the same element. In both cases a hydrocarbon group is detached from the molecule, and also the residue of the acid, but while with the sulphur compounds these two simply combine (forming a compound ether), and leave a hydrocarbon sulphide, in the case of the phosphonium salt the acid residue is reduced by the tertiary phosphine, and the group thus left combines with the hydrocarbon radicle, forming a ketone.

Thus-

$$Et_2S < Et_2S + EtOOCR$$

$$Et_{3}P < Et = Et_{3}PO + EtOCR.$$

A result of this kind is in perfect harmony with the views expressed by Crum Brown and Letts (Tr. E. 28, 371; Letts, Tr. E. 30, 285) regarding the analogies and differences existing between phosphorus and sulphur and their com-

When the phosphonium salts contain ethylene they suffer a different decomposition under the influence of heat; at least this has been ascertained to be the case with the bromide of bromo-ethylene-triethylphosphonium, and the bromide of hydroxy-ethylene-triethyl phosphonium, which decompose as follows (Hofmann) :-

 $\begin{array}{l} C_2H_4Br.P(C_2H_5)_{,B}r = HBr + (C_2H_3)(C_2H_5)_{_3}PBr \\ C_2H_4(OH)_{,P}(C_2H_5)_{_3}Br = H_2O + (C_2H_3)(C_2H_5)_{_3}PBr \end{array}$ and also in that of the hydrate of ethylene-hexethyl diphosphonium, which decomposes according to the equation: $C_2H_4(P(C_2H_3)_3OH)_9$ = $C_2H_4 + (C_2H_5)_3P + (C_2H_5)_3PO + H_2O$. Masson and Kirkland (C. J. 1889, 126) have

studied the action of bromine and chlorine on the salts of tetrethyl phosphonium, the results showing a very close similarity between the polyhaloid derivatives of tetrethyl phosphonium and those of trimethyl sulphine and of tetramethyl ammonium previously described by Dobbin and Masson (C. J. 1885, 56; 1886, 846). The tendency to form solid poly-haloid compounds is, however, more marked. For the table containing a list of the new substances, the methods of forming them, and their chief properties, the original memoir should be consulted.

Organic Phosphorus Compounds which cannot be placed in any of the above Groups. A phosphorised cacodyl (CH2),P2 (the methyl analogue of liquid phosphoretted hydrogen) was obtained by Thénard (C. R. 21, 144, 25, 829) by the action of chloride of methyl on phosphide of calcium, and is interesting not alone as being the sole representative (as yet prepared) of its class, but also as having been probably the first phosphine obtained. Thénard describes it as a colourless, highly refractive liquid, of an odour water, and boiling at about 250°. It inflames spontaneously in contact with air, but if oxidised slowly gives a crystalline acid (CH₂), P₂H₂O₄ = (CH₅)₂PHO₂ (dimethyl phosphinic acid ?), analogous to cacodylic acid. Treated with an excess of hydrochloric acid, it is converted into trimethyl phosphine, and a solid yellow substance (CH₃)₂P₄ (which is also formed in the original reaction), and which Thénard regarded as the methyl analogue of solid phosphide of hydrogen. Michaelis (B. 10, 807) obtained a substance,

Michaelis (B. 10, 807) obtained a substance, which he named diphosphenyl or phospho-benzane C_0H_3 .P:P.C. $_0H_3$ (corresponding to azobenzene), by the action of phenyl phosphorous chloride on monophenyl phosphine C_0H_3 .PC. $_2+C_0H_3$.P. $_2+2$ HCl. It is a pale-yellow powder, insoluble in water, alcohol, and ether, but readily soluble in hot benzene, and is slowly oxidised by the air to $(C_0H_3)_2$ P.20. Treated with chlorine, phenyl phosphorous chloride is regenerated. With nitric acid it is oxidised to phenyl phosphinous (benzene phosphinic) acid C_0H_3 PH.0. if the acid is dilute, but to phenyl phosphinic (benzene phosphonic) acid C_0H_3 PH.0. if the acid is strong. Treated with hydrochloric acid, it reacts so as to regenerate the substances from which it is formed.

Michaelis (B. 7, 499) also obtained a substance, which he called di-phospho-benzene hydrate C.H. P.P.OH, by the action of spontaneously inflammable phosphoretted hydrogen on phenyl phosphorous chloride. It is a yellow powder, soluble with ease in bisulphide of carbon, taking fire on exposure to air, and oxidised by nitric acid to phenyl phosphinic (benzene phosphonic) and phosphoric acids. In addition to the above, Michaelis (B. 11, 885) obtained a phenylated solid phosphide of hydrogen (C,H,)HP, by treating phenyl phosphorous chloride with a quantity of water insufficient for complete decomposition (for instance, by keeping it in a badlystoppered bottle). It is a dark-yellow amorphous body, having a faint odour of phenyl phosphine, soluble in hot bisulphide of carbon, but insoluble in water, alcohol, and ether. Treated with chlorine it reacts as follows: (C₆H₅)HP₄+6Cl₂= 3PCl₃ + (C₆H₅)PCl₂ + HCl. Nitric acid oxidises it to a mixture of phenyl phosphinic (benzene phosphonic) and phosphoric acids.

The action of phosphonium iodide on aldehydes has been studied by Girard (A. Ch. [7] 2, 50), while that of phosphoretted hydrogen and hydrochloric acid on the same bodies and on ketonic acids has been investigated by Messinger and Engels (B. 21, 328, 2919). Girard obtained products of addition containing four molecules of the aldehyde (valeric, propionic, salicylic, and benzoic) to one of phosphonium iodide.

Messinger and Engels obtained similar bodies by acting on the aldehydes with hydrochloric acid and phosphoretted hydrogen. The compounds thus formed are for the greater part solid, crystalline, and fairly stable. By treatment with

water they are decomposed, and the aqueous solution gives the reactions of hydrochloric acid and phosphoretted hydrogen. Their constitution is probably represented by the formula (R.OHOH), PCI. Chloride of tetra-hydroxyethylidene phosphine (C₂H₂O), PCI is decomposed by caustic potash into the free phosphine (C₂H₂O), PC₂H₄O and the hydrate (C₂H₃O), P(OH). Benzaldehyde and its mono-nitro derivative give compounds which differ from those obtained in the fatty series in that they contain no hydracid.

Messinger and Engels have summarised the results of their researches as follows: (1) Phosphoretted hydrogen does not act on an aldehyde alone, but is absorbed if at the same time a hydracid is present. The absorption occurs more completely if the aldehyde is largely diluted with ether. (2) The aldehydes of the fatty series combine with a molecule of phosphoretted hydrogen and a molecule of hydracid, while those of the aromatic series combine with phosphoretted hydrogen only, though in order that the compound shall be formed the presence of the hydracid is necessary. (With benzoic aldehyde the compound has the formula (C_cH_cCOH),PH₂.) (3) The phosphorised derivatives of the fatty series have an unpleasant smell, and are decomposed by water, while those of the aromatic series have no odour, and are nearly insoluble in water. All are soluble with difficulty in ether, and in some cases are insoluble.

By the substitution of a ketonic acid for an aldehyde in the above reaction, compounds are produced in certain cases. Thus lævulic acid gives an oil, and pyruvic acid a solid compound,

either $(CH_3,CO,CO)_3P$ or $(CH_3-C-C=O)_3P$,

which is a well-defined crystalline body, having neither basic nor acid properties, soluble in alkalis with decomposition, and also decomposed when heated with acids. It dissolves, however, without change in glacial acetic acid, and crystallises out on cooling. Boiled with water, it is decomposed into the substances from which it was originally produced. It forms crystalline compounds with aniline, phenyl hydrazine, and toluene diamine.

E. A. L.

PHOSPHINIC ACIDS. The name oxy-ethylphosphinic acid has been given to the acid CHMe(OH).PO(OH), which is got from aldehyde by successive treatment with PCl, and water. A more appropriate name for this acid is, however, oxy-ethane-phosphonic acid, the term phosphinic acid being reserved for acids R'P(OH), containing trivalent phosphorus.

PHOSPHINO-BENZENE C. H. PO. Phosphenylic anhydride. [100°]. Formed from phosphenylic acid and POOL (Michaelis a. Rothe, B. 25, 1747). White crystalline powder, v. sol. water, being reconverted into phosphenylic acid.

PHOSPHINO-ψ-CUMENE C₆H₂Me₂PO₂.
[216]. Formed from C₆H₂Me₂PO(OH)₂ and C₆H₂Me₂POCl₂ (Michaelis a. Rothe, B. 25, 1749).
Plates (from benzene), v. e. sol. chloroform. Converted by hot water into ψ-cumene phosphonia acid.

PHOSPHINO-TOLUENE C_eH₄Me.PO₂. Toluphosphonic anhydride. [101°]. Formed ene phosphonic anhydride. [101°]. Formed from C,H,PO(OH), and C,H,POCl₂ (Michaelis a. Rothe, B. 25, 1748). Converted by water into toluene phosphonic acid.

PHOSPHITES. Salts of the acid H, PO; v.

PHOSPHORUS, OXYACIDS OF, p. 151.

PHOSPHO-. Use of this prefix applied to inorganic compounds; for phospho-acids, phosphosalts, &c., v. the acids, salts, &c., to which phospho- is prefixed. Thus, phosphomolybdic acid and phosphomolybdates are described under

MOLYBDENUM, ACIDS OF, AND THEIR SALTS.
PHOSPHO-AMIDO-BENZENE SULPHONIC ACID PO₃H₂.NH.C₆H₄.SO₃H. The chloride C.H.NSPO.Cl. [158°] is formed from p-amidobenzene sulphonic acid and PCl, (Laar, J. pr. [2] 20, 250). It is converted by MeOH and alcohol into Me₃A" [114°] and Et₃A" [102°] re-

spectively.

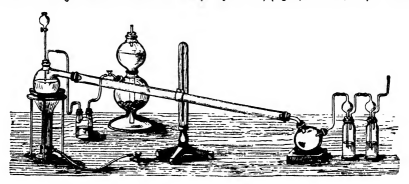
PHOSPHOBENZENE C₆H₅P:PC₆H₅. Formed by slowly adding C₆H₅PH₂ to C₆H₅PCl₂ in a current of H (Köhler a. Michaelis, B. 10, 812). Yellowish powder, insol. water, alcohol, and ether, sol. benzene. On exposure to air it is oxidised to (C₆H_bP)₂O. Dilute HNO₃ oxidises it to phosphenylic acid. Chlorine, diluted with CO₂, forms C₆H₅PCl₂. Conc. phenyl-phosphine and C₆H₅PO₂H₂. Conc. HClAq forms

PHOSPHONIUM COMPOUNDS. PH₃ combines with the halogen acids HX to form com-

20 atmos. pressure at 14°, or by cooling to -25° under ordinary pressure (Ogier, C. R. 89, 705). Skinner (Pr. 1887, 283) found the critical temp. to be 48° under 95 atmos.; and the maximum vapour-pressure line to be below those of HCl and PH, at all temperatures; from - 30° to 10° the line was normal, above 10° combination began; at temperatures near the critical the volume of liquid PH₄Cl formed was nearly half that of the liquid constituents.

Phosphonium hydroxide. (?)PH₄OH. When PH, is strongly compressed in presence of water the PH, liquefies and floats on the water. If the pressure is now suddenly decreased, white crystals are formed; these crystals are probably PHOH (no analyses have been made). The production and dissociation of the crystals occur at definite pressures and temperatures. No formation occurs above 28° at any pressure. The crystals can be distilled from one part of the tube to another by gently warming (Cailletet a. Bordet, C. R. 95, 58). When equal volumes PH₃ and CO₂ are compressed in presence of water, below 22°, the gases disappear and white crystals are formed, which C. a. B. (l.c.) take to be a mixture of PH₄OH and the hydrate CO₂-8H₂O discovered by Wroblewski (C. R. 94, 954).

PHOSPHONIUM IODIDE PH.I. H.F. [P,H4,I] = 29,500 (solid PH I from H, solid P, and solid I); $[PH^3,HI] = 24,100$ (solid PH_4I from gaseous PH₃ and HI) (Ogier, C. R. 89, 705). This com-



pounds PH.HX. The reactions of these compounds are most simply expressed by regarding them as salts of the radicle PH,; following the analogy of the ammonium compounds, the bodies PH,X are called phosphonium compounds. mol. weights of these compounds have not been determined.

PHOSPHONIU M BROMIDE PH,Br. [PH*,HBr]=23,000; [P,H*,Br]=44,100 (solid PH_Br from H, solid P, and liquid Br) (Ogier, C. R. 89, 705). Colourless cubes; melts at 30° . V.D. 27.7 corresponding with that required by PH, + HBr (Bineau, A. Ch. [2] 68, 431). Deliquesces in air; decomposed violently by water to PH, and HBrAq. Prepared by bringing together PH, and HBr (Serullas, S. 64, 238); by leading PH, into cold conc. HBrAq (Ogier, C. R. 89, 705); by heating conc. HBrAq with P to 100°-120° (Damoiseau, C. R. 91, 883).

PHOSPHONIUM CHLORIDE PH.Cl. Formed in colourless crystals, melting at c. 25°, by subjecting a mixture of equal volumes HCl and PH, to capacity, and to this solution, kept cool, 680 g.

pound was discovered by Labillardière and Gay-Lussac (A. Ch. [2] 6, 304), and more fully examined by Serullas (S. 64, 238) and H. Rose (P.

24, 151; 46, 636).

Formation.—1. By bringing together PH, and HI as gases or in conc. aqueous solutions .-2. By the action of gaseous PH, on I (Hofmann, A. 103, 355).—3. By the action of red Ponconc. HIAq at 160° (Oppenheim, Bl. [2] 1, 163).—4. Along with P_2I_4 , by the action of HI gas on P at ordinary temperatures (Damoiseau, C. R. 91, 883).-5. By the action of a little water on PI. (v. Preparation).—6. By the reaction of HI on H₂PO₂ and warming the product in CO₂ (Lissenko, B. 9, 1313).

Preparation.—P and I are combined in CS. solution, and the product is decomposed by a little water (Baeyer, A. 155, 269; Hofmann, B. 6, 286). Hofmann (l.c.) gives the following directions: 400 g. P are dissolved in the same weight of dry CS, in a retort of 1 litre

I are added by small portions. The whole of the CS, is then distilled off on a water-bath, an operation which takes 9 or 10 hours. The retort is then connected with a long wide condensing tube, and a two-necked globular receiver, from which a connecting tube passes to two condensing bottles, the first containing dilute HIAq, the second water (see fig.). Heat is then applied to the retort, gently at first, and 240 g. water are slowly added through the drop-funnel. PH,I and HI are formed; the HI collects in the condensing bottles, and the PH,I forms a crust, chiefly in the tube, and in smaller quantity in the globular receiver, which, to prevent stoppage, should be connected with the long tube by a wide tube. To prevent the liquid in the condensing-bottles from being forced back in consequence of the rapid absorption, a slow stream of CO₂ is passed through the apparatus during the whole operation. A well-conducted sublimation may be completed in 8 or 9 hours. When it is finished one end of the long condensing tube is closed with a cork, and the PH₄I is detached by means of a stout wire bent and sharpened at the Hofmann expresses the reaction thus: $18P + 9I + 21H_2O = 7PH_4I + 3H_4P_2O_7 + 2HI$. The PH,I may be sublimed in dry HI.

Properties.—Large, clear, lustrous, pyramidal crystals; boils at c. 80°; may be sublimed in HI.

Reactions.—1. With carbon compounds reacts as a very energetic reducing agent, and is much used in preparing phosphines &c. (v. Phosphines).—2. Decomposed rapidly by water or alkali solution, evolving non-inflammable PH₃ (v. Phosphorus trihydride, Preparation 2, p. 136). 3. With alcohol forms EtI and PH₃; if the reaction is conducted in a sealed tube PEt₃ and PEt₄I are produced.—4. With phosphorus trichloride forms P₂I₄, HCl, PH₃, and solid P₂H (de Wilde, B. 16, 217).—5. PH₄I is inflamed by contact with many cone. acids, e.g. HNO₃, HCl, HBr, HI.

PHOSPHONIUM SULPHATE. When PH₃ is passed into H₂SO₄ at ordinary temperatures much heat is produced, S separates, and SO₂ is formed. The reaction is much slower if the acid is cooled by ice and salt; if the H₂SO₄ is cooled to -20° or -25° (by rapid evaporation of CH₃Cl) a syrupy liquid is formed, from which a white, crystalline, very deliquescent solid separates; this solid seems to be phosphonium sulphate [?(PH₄)₂SO₄] (Besson, C. R. 109, 644). Exposed to air, H₂PO₄, H₂PO₃, and H₃PO₂ are formed with SO₂. S, and a little H₂S; dissolves in water with hissing noise and evolution of PH₃, but no reduction of the H₂SO₄.

PHOSPHORIC ACIDS. In this article are described the three acids, H₂PO₄, HPO₃, and H₄P₂O₇; the salts of these acids are described under Phosphates (p. 106); for the other oxyacids of P and their salts v. Phosphorus, oxyacids of, and their salts v. Phosphorus, oxyacids of, and their salts (p. 149). The oxide P₂O₃ is the anhydride of three acids: cold water added to P₂O₃ forms HPO₃, the long-continued action of moist air on HPO₃ produces H₄P₂O₇, and H₄PO₄ is formed by boiling for some time an aqueous solution of HPO₂ or H₄P₂O₇. The three acids may be formulated as P₂O₃.H₂O, P₂O₃.2H₄O, and P₂O₃.3H₂O (cf. Phosphates, p. 106).

ORTHOPHOSPHORIC ACID H₂PO₄. (Phosphoric acid. Tribasic phosphoric acid.) S.G. 1-88 (Schiff, A. 113, 183). Melts at 41.75°; liquid solidifies at 38°, temperature rising to 40·5° (Berthelot, Bl. [2] 29, 3). H.F. [P,O⁴,H³] = 302,600 (formation of solid acid); 300,080 (formation of molten acid); [P,O⁴,H³,Aq] = 305,290 (Th. 2, 225). Heat of fusion = -2520 (Th., l.c.). One mol. w. of the molten acid in grams occupies $52\cdot02$ c.c. (Th., l.c.). $\mu_{\rm A}1\cdot3584$, $\mu_{\rm B}1\cdot363$, $\mu_{\rm H}1\cdot3746$, for H₃PO₄Aq, S.G. 1·11 (= 29 p.c.) (Gladstone, C. J. [2] 8, 101, 147). Affinity in aqueous solution = c. 25 (HNO₄=100) (Ostwald). Formation.—1. By oxidising P, by long expo-

sure to moist air (Leeds, A. 200, 286; Wenzell, Ph. [3] 14, 24), or by HNO, HClO, HClO, SO, NO₂, &c.; phosphates are formed by treating P with many metallic oxides, and with alkali carbonates.—2. By oxidising H,PO,Aq or H,PO,Aq by ClAq, BrAq, or IAq, HClOAq, HNOs, or by Ag₂O or HgO; H₃PO₃Aq is oxidised to H₃PO₄Aq by standing in air.—3. By reaction of boiling water with P2O3, HPO3, or H4P2O7.-4. By burning PH, in moist air or O.-5. By reaction of water with POCl₃, or much water with PCl₃. 6. By oxidising, by HNO,Aq, the mixture of H,PO,Aq and H,PO,Aq obtained in making HI by the action of P, I, and H₂O (Pettenkofer, A. 138, 57).—7. By decomposing many phosphates, e.g. Ca₃P₂O₈, by H₂SO₄; Pb₃P₂O₈ decomposed by H₂S in presence of water gives H₃PO₄Aq. For production from bones v. DICTIONARY OF APPLIED CHEMISTRY.—8. By decomposing (NH₄)H₂PO₄ by warm conc. HClAq, separating NH₄Cl after cooling, and evaporating with HNO, till all NH, salts

and HCl are expelled (Joly, C. R. 102, 316).

Preparation.—One part P is mixed with 10-12 parts HNO₃Aq (free from H₂SO₄) S.G. 1·2 (the acid must not be more conc. than this) (Kranthausen, Ar. Ph. [3] 10, 410) in a large retort, the beak of which passes loosely into a receiver placed in cold water; about 3 to 6 g. I is added for 100 g. P; the retort is gently warmed until the P has completely dissolved. Should any liquid distil over, it is returned to the retort. Oxidation proceeds more regularly, and at a lower temperature, when a little I is present than when HNO3Aq alone is used (Horn, Ph. [3] 10, 468; Ziegler, D. P. J. 258, 239). If I is not added, and the acid is more conc. than S.G. 1.2, violent explosions may occur. Von Schrötter recommends the employment of amorphous P and acid more conc. than When the P is all dissolved, the con-S.G. 1·2. tents of the retort are boiled, and 5-7 parts of the liquid are distilled off (the distillate is HNO₈Aq, S.G. 1·1 to 1·14). During this part of the process copious evolution of NO2 often occurs, due to the oxidising action of HNO, on H,PO,. According to Reinsch (J. pr. 28, 385) little NO, but much N, is formed during the oxidation of P by HNO, at boiling temperature in absence of air, but much NO is produced in an open vessel at low temperatures. The liquid in the reto:t is evaporated in a basin until the fumes cease to turn blue litmus red. During this operation NO. is often freely given off (especially if little or none of this gas has been produced in the retort), as the oxidation of H₂PO, is not always completed until the liquid has been evaporated; should no gas be evolved it is advisable to add a fittle conc. HNO₂Aq to the liquid in the basin after concentration, and again evaporate till acid fumes cease to come off. The liquid in the basin is diluted with water, saturated with H_2S , kept in a warm place for 24 hours, and filtered from As_2S_3 (and Sb_2S_3), which may separate (ordinary P generally contains As). The filtrate is then evaporated, at a temperature not above 150° , until it becomes a thick syrup on cooling, and the cold liquid is placed over H_2SO_4 , when it slowly solidifies. By evaporating the syrup in vacuo over H_2SO_4 , crystalline tablets of H_2PO_4 are obtained; a fragment of these placed in O_4 are obtained; a fragment of these placed causes crystallisation (Huskisson, B. 17, 161).

Properties.—A thick semi-solid mass, S.G. 1-88 (Schiff, A. 113, 183); also obtained in crystalline tablets by evaporation of the syrup S.G. 1-88 in vacuo over H₂SO₄. According to Berthelot (Bl. [2] 29, 3) crystals of H₃PO₄ melt at 41·75°; the liquid solidifies at 38°, with a rise of temperature to 40·5°. H₃PO₄ is tribasic, forming salts M¹H₂PO₄, M¹₂HPO₄, and M¹₃PO₄ (v. Phosphates, p. 107). The affinity of H₂PO₄ in aqueous solution is about 25 (HNO₃ = 100) (Ostwald). Both crystalline and syrup-like H₂PO₄ deliquesce in air. The acid dissolves readily in water Hager (in his commentary to the Pharmacopæia Germanica) gives the adjoining table.

According to Sieber (Ph. [3] 9, 598), H₃PO₄Aq

containing 5 p.c. acid (or even less) stops putrefaction.

Reactions.-1. Heated to c. 213°, H,P2O, is formed; at a higher temperature HPO3 begins to be formed, and at full red heat only HPO, is produced.—2. Is not decomposed by electric current, according to Luckow (Fr. 1880.1); by electrolysis of H₃PO₄Aq, using C poles, Bartoli a. Papasogli (G. 11, 239, 468; 12, 113, 117, 125) obtained a substance containing C and P, which they called phosphomellogen. Molten H₂PO₄, on electrolysis, gives off H and O, and forms H.P. then HPOs, and then HsPOs, and inflammable P hydride (Janecek, C. C. 1888. 273).—3. It is not certain whether water forms hydrates with H.PO., when the acid is dissolved in water (v. Crompton, C. J. 53, 116).—4. By adding alkali solutions to H3POAq till neutral, salts M2HPO4 are formed; addition of considerable excess of alkali produces M₃PO₄. For connection between quantity of alkali added and electrical conductivity of the solution, v. D. Berthelot (C. R. 113, 851).-5. Reacts with phosphorus pentachloride at ordinary temperatures to form POCl, and HCl; with phosphorus oxychloride when hot to form HPO, and HCl, or H, P2O, and HCl, according to the relative quantities of H₃PO, and POCl₃; with phosphorus trichloride to form HPO₃, H₂PO₃, and HCl; and with metaphosphoric acid to form H₄P₂O₄, (Geuther, J. pr. [2] 8, 359). For reactions of salts of H₂PO₄ v. Phosphates, p. 107. For nitriles of H.PO. v. Phospham (p. 104), and PHOSPHORUS OXYNITRIDE (p. 144); for amide and imido amide v. Phosphamides (p. 105) and PHOSPHAMIDO-IMIDE (p. 105). For thio-amide v. PHOSPHORUS Reaction SULPHOCHLORIDE, (p. 148).

Pyrophosphobic Acid, H,P2O7.

Preparation.—1. An impure acid, containing some H₂PO₄ and HPO₅, is obtained by heating H₂PO₄ to 213° until a little dissolved in cold

Sp. gr.	Per cent. P.O.	Per cent. H.PO.	Sp. gr.	Per cent. P.O.	Per cent. H.PO.
1.809	68	93.67	1.325	85	48-21
1.800	67.5	92.99	1.319	34.5	47.52
1.792	67	92.30	1.314	34	46.84
1.783	66.5	91.61	1.308	83.5	46.15
1.775	66	90.92	1.303	83	45.46
1.766	65 5	90.23	1.298	32.5	44.77
1.758	65	89.54	1.292	32	44.08
1.750	64 5	88.85	1.287	31.5	43.39
1.741	64	88.16	1.281	31	42.70
1.733	63.5	87.48	1.276	30.5	42.01
1·725 1·717	63 62·5	86·79 86·10	1.271	30 29·5	41.33
1.709	62	85.41	1.265 1.260	29	40·64 39·95
1.701	61.5	84.72	1.255	28.5	39.26
1.693	61	84.03	1.249	28	38.57
1.685	60.5	83.34	1.244	27.5	37.88
1.677	60	82.65	1.239	27	37.19
1.669	59.5	81.97	1.233	26.5	36.50
1.661	59	81.28	1.228	26	35.82
1.653	58.5	80.59	1.223	25.5	35 ·13
1.645	58	79.90	1.218	25	34.44
1.637	57.5	79.21	1.213	24.5	33.75
1.629	57	78.52	1.208	24	33.06
1·621 1·613	66.5 56	77·83 77·14	1·203 1·198	23·5 23	32·37 31·68
1.605	55.5	76.45	1.193	22.5	30.99
1.597	55	75.77	1.188	22 3	30.31
1.589	54.5	75.08	1.183	21.5	29.62
1.581	54	74.39	1.178	21	28.93
1.574	53.5	73.70	1.174	20.5	28.24
1.566	53	73.01	1.169	20	27.55
1.559	52.5	72.32	1.164	19.5	26.86
1.551	52	71.63	1.159	19	26.17
1.543	51.5	70.94	1.155	18.5	25.48
1·536 1·528	51 50·5	70·26 69·57	1·150 1·145	18 17·5	24·80 24·11
1.521	50	68.88	1.140	17	23.42
1.513	49.5	68.19	1.135	16.5	22.73
1.505	49	67.50	1.130	16	22.04
1.498	48.5	66.81	1.126	15.5	21.35
1.491	48	66.12	1.122	15	20.66
1.484	47.5	65.43	1.118	14.5	19.97
1.476	47	64.75	1.113	14	19.28
1.469	46.5	64.06	1.109	13.5	18.60 17.91
1·462 1·455	46 45·5	63·37 62·68	1·104 1·100	$\begin{array}{c} 13 \\ 12.5 \end{array}$	17.22
1.448	45	61.99	1.096	12 0	16.53
1.441	44.5	61.30	1.091	11.5	15.84
1.435	44	60.61	1.087	11	15.15
1.428	43.5	59.92	1.083	10.5	14.46
1.422	43	59.23	1.079	10	13.77
1.415	42.5	58.55	1.074	9.5	13.09
1.409	42	57.86	1.070	9	12.40
1.402	41.5	57.17	1.066	8.5	11.71
1.396	41	56.48	1.062	8	11.02
1·389 1·383	40·5 40	55·79 55·10	1.058	7·5 7	10·33 9·64
1.377	39· 5	54.41	1.049	6.5	8.95
1.371	39	53.72	1.045	6	8.26
1.365	38.5	53.04	1.041	5.5	7.57
1.359	38	52.35	1.037	5	6.89
1.354	37.5	51.66	1.033	4.5	6.20
1.348	37	50.97	1.029	4	5.51
1.342	86.5	50.28	1.025	3.5	4.82
1.336	86	49.59	1.021	3 2·5	4.18
1.330	35· 5	48.90	1.017	2.9	8.44

water gives a white pp., without a trace of yellow Ag, PO,, with AgNO, Aq, after neutralisation by NH₂Aq.—2. Péligot (A. Ch. [2] 73, 286) obtained crystals of H₄P₂O, by allowing glacial HPO, to stand in moist air in a bottle for some years. Crystals of H₃PO₄ formed at the top of the bottle, a liquid S.G. 1.7 in the middle, and crystals of H₄P₂O₇ at the bottom.—3. An aqueous solution of H₄P₂O₇ is obtained by decomposing Pb₂P₂O₇ suspended in water by H₂S, filtering from PbS, and allowing the H2S to evaporate at the ordinary temperature (Geuther, J. pr. [2] 8, 359).—4. By heating together H,PO, and HPO, in the ratio HPO, H,PO, -5. By the reaction of H₃PO₄ and POCl₃ in the ratio 5H₃PO₄:POCl₃

Properties and Reactions. — A glass-like solid; Péligot (A. Ch. [2] 73, 286) obtained the acid in non-transparent crystals resembling loaf-sugar. Dissolves readily in water; dilute solutions remain unchanged at ordinary temperatures (Graham), when heated H₃PO₄Aq is formed. The solution, neutralised by NH₃Aq, gives a white pp. of Ag,P₂O₇ with AgNO₃Aq. H₄P₂O₇ is tetrabasic; for description of salts v. Phosphates, p. 107. When heated with v. PHOSPHATES, p. 107. When heated with PCl_s reacts according to relative masses used: (1) H₄P₂O₇ + 5PCl_s = 7POCl_s + 4HCl; (2) H₄P₂O₇ + PCl_s = 2HPO_s + PCCl_s + 2HCl (Geuther, J. pr. [2] 8, 359). PCl_s heated with H₄P₂O₇ forms HPO₃, H₃PO₅, and HCl; POCl_s produces HPO₃ and HCl (G., l.c.). For the amic acids desired from heated and the polysical from heated solid desired from heated solid from heated solid from heated solid desired from heated solid fro H₁P₂O₇, and other amic acids derived from hypothetical condensed pyrophosphoric acids, v. Рнозрнами асиз (р. 105).

METAPHOSPHORIC ACID HPO, (Graham, T.

1833. 253).

Preparation.—1. H.PO.Aq or H.P.O.Aq is evaporated to dryness and the residue heated to full redness until H2O ceases to be evolved.—2. PbP₂O₄ is suspended in water and decomposed by H₂S, the filtrate from PbS is evaporated to dryness and heated to full red heat .- 3. By heating together H₂PO₄ and POCl₂ in the ratio 2H₃PO₄:POCl₃, or H₄P₂O₇ and POCl₂ in the ratio 2H₄P₂O₇:POCl₃, or H₄P₂O₇ and PCl₃ in the ratio 3H₄P₂O₇:PCl₄ (Geuther, J. pr. [2] 8, 359).—4. By reacting on H₃PO₃Aq with Cl or Br.—5. A solution of P2Os in a little cold water contains HPOs.

Properties and Reactions .- According to Tammann (J. pr. [2] 45, 417), two varieties of HPO, exist; one is a soft, silky mass, formed by heating H₃PO, until one molecule of H₂O is removed; the other is obtained by heating H.PO. until it sets to a glassy mass on cooling. A transparent, colourless, glass-like solid. Ordinary containing considerable quantities of Na salts, and some H₂PO₄ or H₄P₂O₅. Very hygroscopic, soon deliquesces in air. Volatilised completely at bright-red heat (H. Rose, A. 76, 2, 13; 77, 319). Not dehydrated by heat nor by the action of dehydrating agents. Heated with PCl, forms POCl, and HCl, HPO, Aq is changed to H, PO, Aq, slowly at ordinary temperatures, more rapidly by boiling or by adding HClAq or HNO, Aq. (For rate of change v. Sabatier, C. R. 106, 63; 108, 788, 804.) HPO, is monobasic; several series of metaphosphates, some polymeric and some isomeric, exist (v. Phosphates, p. 106). M. M. P. M.

PHOSPHORIC ETHER v. TRI-ETHYL-PHOS

PHOSPHOROUS ACID H.PO. v. PHOSPHORUL,

OXYACIDS OF, p. 149. PHOSPHOROUS ETHER v. ETHYL PROS-PHITES.

PHOSPHORSELLIC ACID v. vol. iii. p. 782. PHOSPHORUS. P. At. w. 30.96. Mol. w. 123.84 and 61.92 (v. infra). The following data apply to ordinary crystalline P. Melts at 44.3° (Schrötter, P. 81, 299); at 44.2° (Desains, C.R. 23, 149). Boils at 290° (Pelletier, A. Ch. 4, 3); at 250° (Heinrich, Gm.-K. [6th ed.] 2, 102); at 287.3° at 762 mm., 165° at 120 mm. (Schrötter J. 1847-48). S.G. 1.8367 at 0°, 1.8232 at 20°. 1.8068 at 40° (Pisati a. de Franchis, B. 8, 70). -76 to 10 = 1035, 7 to 15 = 1054 (Regnauta, A. Ch. [8] 26, 286); 13° to 86° = 202 (Kopp, T. 155, 71); liquid = 2046 (Person, A. Ch. [8] 21, 295). S.V.S. c. 17; at b.p. c. 21 (Ramsaya. Young, C. J. 39, 50). S.V. of P in PCl, and PBr_s = 25'3. C.E. :0003674 + :000000211t from 50° to 280°; C.E. liquid P from 50° onwards = 0005167+ ·00000037 (t=50) (Graham-Otto, 1881. II. 2, 283; v. also Kopp, A. 93, 129; Pisati a. de Franchis, B. 8, 70; Leduc, C. R. 113, 259).

Refraction-equivalent $\left(\frac{\mu-1}{d}$. At. w.) 14.93;

spec. refractive power for H line $\alpha = 4816$ (Haagen, P. 131, 117). H.C. $[P^2, O^5] = 369,900$, $[P^2, O^5, Aq] = 405,500$, $[P^2, O^5, Aq] = 250,060$ (Th. 2, 408). Crystallises in dodecahedra and octahedra (Trantwein a. Buchner; Whewell, C. N. 39, 144; Story-Maskelyne; Mitscherlich, B. B. 1855. 409; Hermann, B. 6, 1415). For spectrum, v. B. A. 1884. 434.

The following data apply to red amorphous P. S.G. 2·106 at 17 (Schrötter, J. 3, 262), 2·19° Hittorf); in powder S.G. = 1.964 at 10° (S.), 2-6 (H.). S.H. 15° to 98° = 1698 (Regnault, P. 89, 496). S.V.S. c. 15. E.C. (Ag at 0° = 100) 00000123 at 20° (Matthiessen, P. 103, 428). H.C. [P', O'] = 862,820 (Troost a. Hautefeuille, C. R. 78, 748).

The following data apply to crystalline metallic P. S.G. 2:34 at 155°. S.V.S. c. 13:2

(Hittorf, P. 126, 193).

Historical.-In the early part of the seventeenth century a substance was prepared by heating heavy spar with combustible organic matter, e.g. egg-white and charcoal, and this substance glowed in the dark (Kopp, Geschichte der Chemie, iv. 42). From this time the name phosphorus was given to any substance which emitted light in the dark without being ignited $(\phi\omega\sigma\phi\delta\rho\rho\sigma)$ = light-bearer). The substance now known as phosphorus was prepared for the first time about 1670; Brand seems to have been the discoverer, but the discovery is sometimes attributed to Kunkel or to Boyle (for details v. Kopp's Geschichte der Chemie, iii. 327).

For a long time P was prepared in very small quantities; it was obtained by evaporating urine, mixing the residue with sand, and heating in a very hot fire. In 1787 charcoal was mixed with the sand and urine-residue. The discovery of phosphates in bones by Scheele in 1771, and in minerals by Gahn in 1780, led to the preparation of P on a comparatively large scale. discovery of P played an important part in the development of the phlogistic theory. Phosphorus was supposed to be a compound of phlogiston with the acid which was formed when P was burnt; this acid was composed of phlogiston and muriatic acid according to Stahl. and of vitriol and muriatic acid according to Hofmann. Marggraf in 1743 determined that P increased in weight when burnt, and that P was obtained by heating phosphoric acid with combustible matter. In 1777 and 1780 Lavoisier showed that P was a constituent of phosphoric acid (cf. Kopp, l.c.).

Occurrence.—P is not found uncombined. Phosphates of Al and Ca occur in large quantities in many rocks; phosphates of Fe, Pb, Mg, and NH, &c. are also widely distributed minerals. Coprolites (the fossilised excreformer land-animals) and guano ments of (the excrement of certain birds) consist very largely of Ca phosphate. Phosphates also occur in some kinds of coal (Carnot, C. R. 99, 154). Phosphates of Ca and Mg are found in the ashes of plants and the bones of animals; P occurs in combination with C, H, and N in the yolk of eggs, in blood, in semen, and in nerve and brain

matter.

Formation.—Ordinary phosphorus is formed: 1.—By heating CaPO, with powdered charcoal, 3Ca(PO_s)₂+10C $=4P+10CO+Ca_{2}(PO_{4})_{2};$ or with SiO, and charcoal, $2\text{Ca}(\text{PO}_3)_2 + 10\text{C} + 2\text{SiO}_2$ = $4P + 10\text{CO} + 2\text{CaSiO}_3$.—2. By reducing Pb₃(PO₄)₂ with charcoal (Donovan, P. M. [4] 2,

202; Fourcroy a. Vauquelin, Gmelin's Handb. [6th edit.] i. 2, 95).—3. By decomposing a strongly-heated mixture of Ca, (PO4), and C by

HCl (Cari-Montrand, C. R. 38, 864).

Red phosphorus is formed: 1. By the action of light, heat, or electricity on ordinary P, whether the P be solid or in solution, dry or moist, at ordinary temperature or at -14° (Schrötter, W. A. B. 1, 180; 8, 241; 9, 414; Corenwinder, A. Ch. [3] 30, 242; Lallemand, C. R. 70, 182; Hittorf, P. 126, 193; V. Meyer, B. 15, 297; Grove, C. J. 16, 269).—2. By heating common P with a little I. One part of I suffices to transform c. 100 parts of P into the red variety, by melting the P, mixing in the I, and heating again; the mass becomes red at 100°, at 120°-130° a red powder separates, at 200° a more or less explosive action occurs and the change is complete (Brodie, J. pr. 58, 171).-3. By adding a little I to a solution of common P in CS, evaporating to dryness, adding water, and washing with CS, (Corenwinder, A. Ch. [3] 30, 242; Lallemand, C. R. 70, 182; Brodie, J. pr. 58, 171). Rüdorff (P. 128, 463) supposed the red substance thus formed to be a hydride of P.

1. Crystalline ordinary phosphorus is formed: 1. By heating red P in a vacuous sealed tube to 447° (Hittorf, P. 126, 193).-2. By dissolving ordinary P in molten lead and crystallising therefrom (H., l.c.).

Preparation.-1. The details of the preparation of ordinary phosphorus from bone-ash are described in DICTIONARY OF APPLIED

CHEMISTRY, vol. iii. Ordinary P frequently contains notable quantities of As, derived from the H₂SO₄ used; this can be removed only by repeated treatment with dilute HNO, Aq, S.G. 1.1, a process which involves the loss of about $\frac{5}{6}$ of the P (Herz a. Bärwald, B. B. 32, 2, 133; cf. Dulk, ibid. 34, 1, 247; also Wackenroder, J. pr. 2, 340; and Liebig, A. 11, 260). Traces of red P may be removed by shaking in a closed vessel with conc. K₂Cr₂O₂Aq and H₂SO₄ (Wöhler, A. 45, 249). Lockyer (C. N. 40, 101) found that a specimen of dry P gave off considerable quanti-

ties of H when heated in vacuo.

2. Red or amorphous phosphorus.-A small flask of hard glass is fitted with a good cork carrying a short tube and an exit-tube which dips just beneath the surface of a little Hg; the entrance tube is fitted with a stopcock. Dry CO₂ is passed through the apparatus for some time; a dry stick of common P is then placed in the flask, and the stream of CO, is continued for a little, after which the stopcock is closed, and the flask is heated on a sand-tray or in an oil-bath, to 240°-280°, for 50-60 hours. After cooling, a layer of red P is found at the bottom of the flask, generally covered by a mix-ture of the two kinds of P. The ordinary P is melted by running in a little water at 50°-60°; the water and melted P are poured off; the residual red P is washed repeatedly with CS, (addition of CaCl, Aq, 38°-39° Beaume, causes the CS₂ solution to rise to the surface; Nicklés, C. R. 42, 646), or with boiling KOHAq S.G. 1.3, and then with water containing a little HNO₃; the red P is finally washed with water, and dried at c. 100° (Schrötter, W. A. B. 1, 130; 8, 241; 9, 414).

The change of common into red P occurs in ten minutes by heating the former to 300° in sealed tubes of thick glass, from which the air has been pumped out (Hittorf, P. 126, 193; V.

Meyer, B. 15, 297).

3. Crystalline metallic phosphorus. Hittorf (P. 126, 193) claimed to have formed a distinct variety of P by heating red P in sealed tubes of hard glass to c. 447° (in vapour of S); and also by dissolving common P in molten lead and crystallising. A wide tube of hard glass is closed at one end, filled to 1 with common P, and then filled with pieces of lead; the open end is narrowed, and the air is pumped out of the tube, which is then sealed. The tube is embedded in calcined MgO, placed in an iron tube, and heated above the melting-point of Pb for 8-10 hours. After cooling, the Pb is dissolved in HNO.Aq, S.G. 1.1, and violet-black crystals of P remain; these crystals have not, however, been obtained quite free from lead.

Properties. - P exists in three, and perhaps

in more than three, modifications.

1. Ordinary, octahedral, or phosphorus is a semi-transparent, colourless, crystalline solid, with a very distinct smell. The smell is very probably due to a mixture of ozone and P₂O₂, formed by the oxidation of the P (Schönbein, P. 65, 377; Thorpe a. Tutton, C. J. 57, 573). When melted and cooled quickly P is opaque; when cooled slowly it is nearly as clear as water. P must be kept under water; if the water is free from air the P remains transparent (Baudrimont, C. R. 61, 857). In ordinary water

P becomes covered with a whitish film, but this again becomes transparent at c. 50° (H. Rose, P. 27, 563; Baudrimont, l.c.). According to Marchand (J. pr. 20, 506) the white film contains from 4 to 7 p.c. water; Pelouze (A. Ch. [2] 50, 83) regarded this substance as a hydrate of P and Mulder (J. Ph. 23, 20) as a compound of oxide with hydride of P. When melted in large quantity and allowed to cool slowly, fairly large dodecahedral and octahedral crystals separate (Hautwein a. Buchner, Kastn. Arch. 10, 127, 504; Whewell, C. N. 39, 144; Hermann, B. 6, 1415). At ordinary temperatures P is as soft as wax; when cooled it becomes brittle. Exposed to light, P soon becomes yellow and then red on the surface. In a vacuum tube, or in a tube filled with an indifferent gas, P easily sublimes in colourless, transparent, very lustrous crystals (Hermann, B. 6, 1415; Mitscherlich, B. B. 1855. 409; Blondlot, C. R. 63, 397; Desains, C. R. 23, Melted P often remains liquid many degrees below its m.p.; this is shown especially when P is melted by warming with alcoholic KOH solution, or when a solution of P in CS₂ is gradually evaporated under water (H. Rose, P. 32, 469; Kallhofert, J. pr. 50, 1). In some cases P remains liquid at c. -15°. P may be obtained in finely-divided particles by melting it under water, and shaking carefully in a flask until the P solidifies; if pure P is melted under a solution of sugar, gum, dextrin, NH,Cl, (NH₄)₂CO₃, &c., and shaken until it solidifies, the P is obtained in a state of extremely fine division (Casaseca, J. Ph. 16, 202; Blondlot, J. 1865, 126; Schiff, A. 118, 88; also A. Suppl. 4, 37). P is slightly sol. water, alcohol, ether, ethereal and fatty oils, and hot conc. acetic acid (Vulpius, Ar. Ph. [3] 13, 38). Water shaken with P loses the smell and taste of P, and ceases to glow in the dark when it has been exposed to air for a time (Müller, B. 3, 84). P is very soluble in CS₂, S₂Cl₂, and liquid P₂S.

P volatilises in water-vapour, even at the

P volatilises in water-vapour, even at the ordinary temperature; when P is distilled in steam and the distillate is cooled rapidly, the P is obtained as a white snow-like mass (Remsen a. Kaiser, Am. 4, 459); it is also often obtained in this form by distilling crude HIAq containing

P (Hell, J. 1883. 312).

P oxidises very easily; slowly when exposed to low temperatures, and more rapidly as the ignition-temperature, 60°, is approached. When P is burnt in air or O it emits very bright yellow-white light, and much heat is produced. But if H is passed over slightly warmed P, or if a stream of the warm gas is charged with a very little P, the issuing gas glows with a very feeble greyish-white light, scarcely visible in daylight; the temperature of this flame is very low: if the H is ignited an emerald-green cone appears in the interior of the H flame (for spectrum of this green cone v. Christofle a. Beilstein, C. R. 56, 399; Salet, A. Ch. [4] 28, 56). The glowing of P in the dark was formerly supposed to be due to the evaporation of the P (Corne, J. Ph. [5] 6, 17; Marchand, J. pr. 50, 1). It was shown that P glows in an indifferent gas only when small quantities of O are present; also that P neither oxidises nor glows in pure O at the ordinary pressure, but only when the O is diluted with an indifferent gas or the pressure is diminished; and also that the glowing is stopped by the presence of H₂S, PH₃, SO₂, CS₂, Br, Cl, N₂O, NO₃, &c. (Fischer, J. pr. 55, 342; 39, 48; Schrötter, J. pr. 58, 158; Müller, B. 8, 84; Joubert, C. R. 78, 1858; Deschamps, C. R. 52, 355; Müller-Erzbach, B. 12, 2130; Chappuis, Bl. [2] 35, 419; cf. Thorpe, 'On the Glow of Phosphorus,' N. 41, 523). Much work has been done to determine whether ozone or H₂O₂, or both, is formed when P oxidises in moist air; the question is not yet finally settled (McLeod, C. J. 37, 118; Kingzett, C. J. 37, 792); Leeds (N. Y. Acad. of Sciences, 1, 363, and 3, 150) has given a full index to the memoirs on the subject. Schmid (J. pr. 98, 414) determined that ozone is not formed when P oxidises in dry O. P is extremely poisonous; the lethal dose for adults is from '2 to '5g. Burns with P are very dangerous; the parts should be rapidly and thoroughly washed with dilute bleaching powder solution.

The atomic weight of P has been determined, (1) by ppg. Ag and Au solutions by P and determining the ratio of Ag:P and Au:P (Berzelius, G. A. 53, 433; P. 8, 17); (2) by finding the Ag needed to ppt. Cl from PCl₂ (Pelouze, C. R. 20, 1047; Dumas, A. Ch. [3] 55, 174); (3) by oxidising red P in O to P₂O₅ (Schrötter, J. pr. 53, 435); (4) by analysing and determining V.D. of PCl₃, PF₅, &c.; (5) by measuring S.H. of P.

Determinations of the S.G. of P vapour made by Dumas, Mitscherlich, and Deville a. Troost at temperatures from 500° to 1000° gave numbers agreeing with the molecular wt. $P_4 = 124$. Biltz a. V. Meyer (B. 22, 725) obtained the values 52.5 at 1480° , 46.7 at 1680° , and 45.5 at 1700° for V.D. of P, showing that at a white heat the mol. formula is probably P_2 (V.D. = 31).

The mol. wt. of ordinary P in solution in C_9H_6 was determined by Paterno a. Nasini (B. 21, 2153) by the cryoscopic method; values were obtained between P_2 and P_4 . Hertz, using the same method and the same solvent, obtained the value $124 = P_4$ (Z. P. C. 6, 358). Beckmann also found $124 = P_4$ for P in solution in CS₂ by the method of raising of boiling-point of the solvent (Z. P. C. 5, 76).

The action of light, heat, or electricity on P changes it into allotropic red P. Light affects the change whether the P be in air or another gas, or in vacuo, whether water be present or not (Schrötter, W. A. B.1, 130; 8, 241; 9, 414), at temperatures as low as -14° , and whether the P be solid or dissolved in CS₂ (Corenwinder, A. Ch. [3] 30, 242; Lallemand, C. R. 70, 182). For an account of the action of tropical sunlight on P in CS₂ v. Pedler, C. J. 57, 599. The action of heat begins at c. 215° at the ordinary pressure; diminution of pressure causes a slackening of the rate of change, and the action stops at 393 mm. pressure, temp. being 215°. The change is complete by heating to 260° for c. 8 days; at 300°, in a sealed tube, it is finished in a few minutes (Schrötter, l.c.; Hittorf, P. 126, 193; V. Meyer, B. 15, 297; cf. Dissociation, vol. ii. p. 391). The change is accomplished by electricity by passing the current through a little P in a closed tube with Pt wires fused into the ends (Grove, C. J. 16, 269; Geissler, P. 152, 171). Hittorf (P. 126, 193) regards the action as due

to the heating of the P vapour by the current. Certain reagents also effect the change: e.g. heating with a very little I (Brodie, J. pr. 58, 171); or heating a solution of P in CS₂ with I (Corenwinder, A. Ch. [8] 30, 242; Lallemand, C. R. 70, 182), but Rüdorff (P. 128, 463) says the product is a hydride of P. In the change of 62 g. ordinary P into red P c. 7,100 g.-units of heat are produced; and there is a contraction of volume (v. Petersen, Z. P. C. 8, 601).

2. Amorphous or red phosphorus is a dull, dark carmine, odourless, tasteless powder, which becomes darker on heating, and black when boiled with KOHAq. It is brittle, and shows no trace of crystalline form. Amorphous P is heavier, and conducts electricity much better, than crystalline P, although its conductive power is very small compared with the metals (v. data at beginning of this article). Amorphous P does not melt when heated to redness in a sealed tube (Hittorf, l.c.); vapour is formed in the tube, and on cooling this solidifies as crystalline P, but the unvapourised portion remains amorphous. When distilled, amorphous P becomes crystalline; the action begins at c. 260° (Schrötter, l.c.). The process, like the reverse change of crystalline into amorphous P, is essentially one of dissociation (v. Dissociation, vol. ii. pp. 391-3). Amorphous P takes fire at c. 250°-260° (Hittorf, Schrötter, l.c.), but active combustion begins at c. 300°; it does not oxidise in dry air, but in presence of moisture oxidation proceeds slowly, with the smell of ozone and the formation of an acid liquid (Personne, C. R. 45, 113; Wilson, Ph. 17, 410; Pedler, C. J. 57, 599). Red P does not glow in the air; it is not poisonous (De Vrij, J. 1851. 313). This variety of P is insoluble in the menstrua which dissolve ordinary P; boiling turpentine, and a few liquids of high boiling-points, dissolve a little and change it into ordinary P.

3. Crystalline metallic phosphorus is described by Hittorf (P. 126, 193) as forming long, black, thin, rhombohedral crystals which appear red in transmitted light; S.G. 2:34 at 15:5°; less volatile than amorphous P; heated in a sealed tube drops of ordinary P appear at c. 358°. It is doubtful whether this is a distinct variety of P; it does not seem to have been obtained free from Pb (for preparation, v. ante). According to Troost and Hautefeuille (C. R. 78, 748), the S.G. and heat of combustion of red P change continuously with temperature; at 580° ruby-red crystals appear, which have the S.G. of Hittort's metallic P.

Thénard observed that P became black when heated and suddenly cooled (Gm.-K. [6th ed.] i. 2, 102); some experimenters claim this to be a special variety of P (v. Reichardt, Ar. Ph. [3] 9, 442; P. Thénard, C. R. 95, 409); according to others it contains foreign substances. Blondlot obtained it only when a trace of Hg was present (C. R. 70, 856; 78, 1130); Ritter found the presence of As necessary (C. R. 78, 192).

Napoli (C. R. 25, 369) thought that a special

Napoli (C. R. 25, 369) thought that a special yellow modification of P was produced by keeping freshly-melted ordinary P under water.

The atom of P is trivalent in most gaseous molecules, e.g. PCl₂, PH₃, PF₃, &c.; it is pentavalent in the gaseous molecule PF₂. P acts as a nonmetallic element; its oxides are acidic, although Vol. IV.

some of them combine with less acidic oxides to form salt-like compounds. P is closely related to As, Sb, and Bi, and also to N, V, Nb, Di, Er, and Ta; these elements form Group V. For a detailed consideration of the chemical relations of P v. Nitrogen group of elements, vol. iii. p. 571. The specific volume of P at its b.p. is 20-21 (Pisati a. de Franchis, B. 8, 70; Ramsay a. Young, C. J. 39, 50); the specific volume of P in combination is c. 25·3 (Thorpe, C. J. 37, 333).

Reactions and Combinations.—1. P combines with oxygen to form P4O, P2O3, and P2O5; the reaction begins at ordinary temperature with crystalline P, and at c. 250°-260° with amorphous P. (For more details v. Phosphorus, oxides of, p. 138.) P does not combine with pure O at the ordinary pressure, but only when an indifferent gas is present, or the pressure is diminished (for references v. Properties of P, p. 128). The combination of P in O is arrested in proportion to the dryness of the O (Baker, C. J. 47, 349; Pr. 46, 1). Amorphous P does not oxidise in dry air, but the oxidation proceeds slowly in moist air. Ordinary Poxidises readily in moist air; it is not yet finally settled whether or not ozone is produced (v. McLeod, C. J. 37, 118). In moist air H,PO, is formed; this may be a direct product of oxidation, or it may result from the reduction of H₃PO₄ formed by the direct exidation of the P (v. Schönbein a. Bohlig, J. 1863. 167; Blondlot, C. R. 66, 351; Stiassay, D. P. J. 199, 414).—2. P is easily changed to P2O5 by oxidising agents, e.g. HNO3, CrO₃, KNO₃, &c. A mixture of P and KClO₃ explodes violently when struck or rubbed; the explosion is less violent with amorphous P. Amorphous P is inflamed when rubbed with CrO₃, PbO₂, Pb₃O₄, HgO, K₂Cr₂O₇, or Ag₂O, and when heated with MnO2, CuO, or KNO30-3. P combines with sulphur, the halogens, and many non-metals when heated with these. The reactions occur at lower temperatures, and more rapidly, with ordinary than with amorphous P, yet amorphous P combines with Br and Cl at the ordinary temperature.-4. P decomposes solutions of salts of copper, silver, and some other metals, with separation of the metal, which often combines with the P .- 5. Boiling water is slightly decomposed by P in presence of air, with formation of PH₃, H₂PO₂Aq, and H₂PO₄Aq (Cross a. Higgin, C. J. 35, 254); at 250° the reaction is more rapid (Schrötter); with amorphous P the action is slow (C. a. H.).—6. Concentrated hydrochloric acid is decomposed by P at 140°, with formation of PH3 and H2PO3Aq (Oppenheim, Bl. [2] 1, 168).—7. Hydriodic acid reacts with P to form PH₄I and PI₂; hydrobromic acid reacts at 100°-120° to produce PH,Br (Damoiseau, C. R. 91, 883).—8. Chlorosulphonic acid [SO2-OH.Cl] reacts explosively with ordinary Pat 25°-30°, and more slowly and at a higher temperature with amorphous P, to form SO2, HCl, and H₂PO₄ (Heumann a. Köcklin, B. 15, 416).—9. Sulphuryl chloride [SO₂Cl₂] produces PCl, and SO₂; the reaction is violent, even with amorphous P (H. a. K., B. 15, 1736).—10. Adie (C. J. 59, 230) describes the action of sulphur trioxide on P as producing 3P₂O_{4.2SO₄.—11. With boiling solutions of alkalis P forms PH, and} alkali salts of H.PO. P left for a long time in

sontact with ammonia solution probably forms P hydride and NH, phosphite (v. Blondlot, C. R. 57, 1250; Commaille, C. R. 68, 263; Flückiger, J. 1863. 171). Red P is not acted on by NH₃Aq (F., l. c.).—12. Heated with the alkaline earths P forms phosphides and phosphates. -13. When P is strongly heated with anhydrous alkali carbonates C is separated. For an account of the technical applications of P v. DICTIONARY OF APPLIED CHEMISTRY; for description of the methods of testing for P and estimating it a manual of analysis must be consulted.

Phosphorus, acids of. The phosphoric acids are described under that heading (p. 124), the other acids under the heading Phosphorus, oxy-ACIDS OF, AND THEIR SALTS (p. 149). For phosphamic acids v. p. 105, and for thiophosphamic wids v. Thiophosphoryl chloride, Reaction No. 7, p. 148.

Phosphorus, bromides of. P and Br combine violently; two products of the reaction have been isolated, PBr_s and PBr_s. These for-

mulæ are probably molecular.

PHOSPHORUS TRIBROMIDE, PBr, (Phosphorous bromide). B.P. = 172.9° at 760.2 mm. (Thorpe, C. J. 37, 335). S.G. $\frac{6^{\circ}}{4^{\circ}}$ 2-92311 (T., *l.c.*). Does not solidify at $-13\cdot6^{\circ}$. A colourless liquid, with very penetrating odour; fumes in the air. PBr. is formed by the direct union of P and Br. If P is added to Br, the Br must be kept in a treezing mixture, and the P must be added in extremely small pieces until a colourless liquid is obtained; the process is dangerous. Lieben (A.146,314) recommends to pass CO_2 through slightly warmed Br, to lead the gas over small pieces of dry P, and to distil the liquid thus produced after it has stood for some days. Schorlemmer uses amorphous P. The P is placed in a flask connected with an inverted condenser, and having a dropping funnel passing through the cork. Br is added drop by drop from the funnel: the first few drops combine, with a production of flame, but after a little the Br may be added more freely; the PBr, is distilled from the excess of P (v. also Pierre, A. Ch. [3] 20, 5; Kekulé, A. 130, 16; Löwig, Gm.-K. i. 2, 834).

PBr, is not acted on by O; reacts with S to lorm PSBr_s (Gladstone, P. M. [3] 35, 345); Cl forms PCl, and Br; HgCl, forms PCl, and HgBr,; H₂S produces P₂S₃ and HBr. PBr₃ is decomposed slowly by cold water, rapidly at 25°, to HBrAq and HsPOsAq (Löwig, l.c.). P and I dis-

solve in PBr.

PHOSPHORUS PENTABROMIDE, PBr, (Phosphoric bromide. Perbromide of phosphorus). Produced by adding excess of Br to PBr, stirring well with a glass rod, and after a time removing the excess of Br by warming very gently in a stream of dry CO₂. Also formed by action of Br on PCl, in presence of a little I (Gladstone, P. M. 3] 53, 345). PBr, forms a citron-yellow solid; umes in the air; melts to a red liquid when gently warmed, which solidifies again to rhompoidal crystals, which are yellow if cooling has been rapid, and red if cooling has been slow. The ed crystals become yellow when rubbed (Baudrinont, Bl. 1861. 118); at a higher temperature below 100°) gives red vapours that condense to rellow needles. PBr, is separated at c. 100°

into PBr, and Br, which re-combine on cooling. (Ogier, C. R. 92, 83) gives the following heats of formation: [P,Br] = 63,000 from liquid Br;

=83,000 from gaseous Br; =62,300 from solid Br; in each case solid PBr, is formed.

Reactions.—1. Heated to 200° in oxygen, PBr, inflames with formation of P₂O₅ (Baumstark, A. 140, 75).—2. Neither hydrogen nor sulphur reacts in the cold with PBr. -3. Chlorine forms PCl, and Br; iodine produces PBr, and IBr_s.-4. In moist air, POBr_s is formed (q. v.).-5. A little water produces HBr and POBr₃; with much water HBr and H₃PO, are formed.—6. PBr, reacts with hydroxides to substitute OH by Br; the reaction is similar to, but not so energetic as, that of PCl_s.—7. Hydrogen sulphide forms PSBr₃ and HBr.—8. Phosphorus hydride forms at first PBr,, and then also P and HBr.— 9. Hydriodic acid has no action: dry boric acid forms some BBr, (Gustavson, B. 3, 426); dry oxalic acid produces POBr_s, HBr, CO, and CO₂. 10. Antimony sulphide produces SbBr, and PSBr, (Baudrimont, Bl. 1861. 118).—11. With many metals at high temperatures, PBr, produces phosphides and bromides.

Combination.—With ammonia to PBr_s.9NH_s. Prepared by passing dry NH_s into a solution of PBr, in CCl, (Besson, C. R. 111,

Phosphorus, bromochlorides of. Several compounds of P, Br, and Cl are formed by the reaction of PCl₂ with Br. PBr₂Cl₂, PBr₄Cl₃, or PBr_sCl_s is produced according to the relative quantities of PCl, and Br used; and from one or other of these the compounds PBr,Cl₂ and PBrCl₄ are obtained. According to Stern (C. J. 49, 815) PBr₂Cl₃ is formed, and on addition of more Br the compound PBr, Cl. is produced, and unites with Br to form PBr, Cl2; the whole of the Cl in PCl_s cannot be thus replaced by Br (Michaelis, B. 5, 9, 411; Wichelhaus, A. Suppl. 6, 277; Stern, C. J. 49, 815; Geuther, J. Z. 10, 128; Michaelis a. Geuther, J. Z. 6, 242; Prinvault, C. R. 74, 868). Michaelis regarded PBr Cl. and PBr,Cl, as compounds of PBr,Cl, with Br, and 3Br, respectively; Prinvault looked on PBr, Cl, as PBr, Cl, BrCl, and PBr, Cl, and PBr, Cl, and PBr, Cl, as PBr, 3BrCl and PBr, 2BrCl, respectively. Against Prinvault's view it was urged by Michaelis that if the compounds contain PBr, they must give POBr, and BrOl by reacting with SO, whereas the products are POCl, SBr, and Br; but Geuther showed that POBr, reacts with BrCl to give POCl, and Br. Geuther regarded the compounds as containing trivalent Br atoms.

PHOSPHORUS DIBROMOTRICHLORIDE, PBr2Cl3. Yellow-red crystals, melting at 35°; obtained by gradually adding 160 pts. Br to 187.5 pts. PCl. and keeping in a closed tube in a freezing mixture for some time (M.). Addition of 18 pts. water reacts thus (G. a. M.): 3PBr₂Cl₃+3H₂O $=2POCl_s + POBr_s + 3HCl + 3HBr.$ If SO₂ is passed into the contents of the tube, SBr. and

POCl, are formed (M.)

PHOSPHORUS TETRABROMOTRICHLORIDE PBr₄Cl₂. Large dark-red crystals with a blue sheen; formed by adding Br slowly to PCl, till two layers are produced, the lower of which is dark and oily, and then adding PBr₂Cl₃ until the lower layer solidifies (G.). Also produced by warming PBr₃ with PCl₃ to 40°: **4**PB \mathbf{r}_b +6PCl $_b$ =5PCl $_t$ +5PB \mathbf{r}_t Cl $_s$ (G.), when heated PB \mathbf{r}_t Cl $_s$ melts and separates into two layers, which unite again on cooling. A little H $_2$ O produces POCl $_s$, POB \mathbf{r}_s , HCl $_t$ HB \mathbf{r}_s , and B \mathbf{r}_s : much H $_2$ O forms H $_2$ PO $_s$, HCl $_t$ HB \mathbf{r}_s , and B \mathbf{r}_s ; SO $_2$ produces POCl $_s$, SB $_t$, and B \mathbf{r}_s .

PHOSPHORUS OCTOBROMOTRICHLORIDE PBr₈Cl₃. Brown needles, melting at 25°. Formed by adding 4½ pts. Br to 1 pt. PCl₃ (i.e. in the ratio 8Br: PCl₃), heating to 65°, and allowing the oily liquid to cool to c. 4°; much heat is produced in the reaction (P.). Geuther (l.c.) says the compound is also produced by adding BrCl to PBr₃.

in the ratio PBr₅: 3BrCl.

PHOSPHORUS HEPTABROMODICHLORIDE PBr,Cl₂. Prismatic crystals. Obtained by rapidly distilling the preceding compound at a little above 90° (P.); also by dropping PCl₃ into Br until two layers are formed, and allowing to stand. Soluble in PCl₃.

PHOSPHORUS MONOBROMOTETRACHLORIDE

PBrCl₄. According to Prinvault (l.c.) this compound is obtained, as yellow crystals, by heating to boiling a solution of the preceding compound in PCl₃, also by the reaction of PCl₃ with BrCl. P. says that PBr₈Cl₃ and PCl₃ react in accordance with the equation PBr₈Cl₃ + PCl₃ = PBr₇Cl₂ + PBrCl₄.

Phosphorus, bromofluoride of, PBr_xF₃. A liquid, which fumes in the air, and is violently decomposed by water; also decomposed easily by heat to PBr₃ and PF₃; formed by passing PF₃ into Br (Moissan, C. R. 100, 1348).

Phosphorus, bromonitride of, PBr₂N. This compound is said to be formed by heating the product of the action of NH₃ on PBr₅, in a sealed tube, to 250°-275°, with twice its weight of PBr₅ for twelve hours, and then heating the resulting substances under reduced pressure, when PBr₅ sublimes, and then at 200° a white crystalline sublimate of PBr₂N is formed; the compound is purified by resubliming in vacuo. The crystals melt at 188°-190°; sublime at 150° in vacuo; sol. ether, less sol. CS₂ or CHCl₃, insol. water (Besson, C. R. 114, 1479).

Phosphorus chlorides of. P and Cl com-

Phosphorus chlorides of. P and Cl combine directly and easily, with the production of light and much heat. Two compounds are thus produced, PCl₃ and PCl₃; these formulæ are molecular. According to Donny a. Mareska (C. R. 20, 717), and Dumas (A. Ch. [3] 55, 172), liquid Cl surrounded by solid CO₂ combines

with P.

Рноврновов тисновите PCl_3 (Phosphorous chloride). Mol. w. 137-07. Boils at 75-95° at 760 mm. (Thorpe, C. J. 37, 333; v. also Pierre, A. Ch. [3] 20, 5; Haagen, P. 131, 122). S.G. $\frac{\sigma}{c}$ 1-61275 (Thorpe, l.c.); V.D. 70-6 (Dumas, A. Ch. [3] 55, 172). S.H. of vapour 1347, equal weight of air = 1; '64, equal vol. of air = 1 (Regnault, J. 1863. 85). μ for H line a = 3222; $\frac{\mu-1}{d}$. Mol. w. = 44-3 (Haagen, P. 131, 122). H.F. [P,Cl]

= 44·3 (Haagen, P. 131, 122). H.F. [P,Cl] = 75,300 (Th. 2, 408; formation of liquid PCl₃). Regnault (J. 1863. 65) gave the vapour pressures of PCl₃ in mm. of Hg as follows:—37·98 at 0°, 62·68 at 10°, 100·55 at 20°, 155·65 at 30°, 233·78 at 40°, 341·39 at 50°, 485·63 at 60°, 674·23 at 70°.

Formation.—1. By direct union of P and Cl.—2. By reaction of P with HgCl₂, FeCl₂, CuCl₂,

ICl_s, S₂Cl₂, or SeCl₂ (Gm.-K. i. 2, 390).—3. By reaction of Cl or HgCl₂ with PBr₃ or PI_s (l.c.).—4. By decomposing PCl₃ by heat aided by H, P, PH₃, I, HI, S, or one of several metals.—5. By heating HPO₃ with NaCl (Gm.-K., l.c.).—6. By heating Na₂HPO₄ with NH, Cl (H. Rose).—7. By heating P with conc. HClAq (Oppenheim, Il. [2] 1, 163).—8. By passing POCl₄ through a tube filled with glowing charcoal (Riban, C. R. 95, 1160).

Preparation.—About 500-600 g. P, in pieces about 1 inch long, are dried quickly, twice, between filter-paper, and placed in a dry retort of c. 1 litre capacity, through the neck of which a stream of dry CO2 is kept passing. The beak of the retort is now passed into a well-cooled receiver, and the tubulus is connected with an apparatus which is evolving dry Cl. The tube through which the Cl passes into the retort is fixed somewhat loosely in the cork, so that it may be moved up or down easily. The retort is placed in warm water until the P is melted; after the action has fairly begun, the heat produced suffices to keep the P molten. A large basin with cold water is kept beneath the retort in case the latter should break. The reaction is regulated by moving the tube from the Cl apparatus near to, or away from, the surface of the P; if this tube is too near the P, the heat of the reaction vaporises much of the P, which then condenses in a reddish layer in the upper part of the retort; if the delivery-tube is too far from the P, solid PCl, forms and deposits on the upper part of the retort in yellow crystals. As the reaction proceeds, pieces of red P appear here and there in the retort. Towards the end much red P is formed; this must be heated carefully with a lamp, a process in which the retort is apt to be cracked. The distillate is digested with a little P for a few days, to decompose any PCl_s, and then redistilled from a retort heated on a water-bath (Donny a. Mareska, C. R. 20, 717; Dumas, A. Ch. [3] 55, 172). Thorpe (C. J. 37, 333) recommends to heat red P in dry Cl; he says that the reaction is very regular, and that large quantities of PCl, may be prepared thus with great ease and rapidity; the red P should be washed with water, then with alcohol, then with ether, and dried at a gentle heat.

Properties.—A clear, colourless, highly refractive, liquid, which can be vaporised without decomposition in absence of moisture; fumes in air with decomposition; does not redden dry blue litmus paper; vapour has an acrid smell and excites tears. Does not solidify at -115° (Natterer, P. 62, 133). Does not conduct electricity. Dissolves P; also I (Gladstone, P. M. [3] 35, 345). Mixes with C.H., Et.O. CHCl., and CS2. In a gas-flame or alcohol-flame, PCl., burns brightly. Sinks in water and then begins to decompose to H., PO, Aq and HClAq; if a little warm water is used, amorphous P separates. Slowly decomposes in the dark to PCl., and P (Casselmann, A. 83, 247). Combines with Cl to form PCl. Not acted on by O at ordinary temperatures; POCl. is slowly formed at b.p. of PCl. POCl. is formed by action of ozone and various O compounds which readily part with O (v. Reactions).

Testing PCl, for POCl, PCl, has little ac-

ĸ 2

tion on zinc-dust at ordinary temperatures; POCl, produces P2Os and Zn phosphide, which is decomposed by H2O, giving inflammable P hydride. To apply this reaction, a few drops of the liquid are added to a little zinc-dust in a test tube, and then a few drops of water; if POCl, is present, small bubbles of gas are given off, and take fire

in the air (Deniges, Bl. [3] 2, 787).

Reactions.-1. Water produces H,PO, and HOl; the reaction proceeds somewhat slowly. If a little warm water is added to PCl3, amorphous P separates as the product of a secondary reaction —3 H₃PO₃Aq + 2PCl₃ = 3HPO₃Aq + 6HClAq + 2P (Michaelis, B. 8, 504; but cf. Geuther, J. Z. [2] 3, 2nd Suppl. 116).—2. Oxygen does not react at ordinary temperatures, but POCl₃ is slowly formed at b.p. of PCl₃ (Michaelis, J. Z. 6, 93; Remsen, Am. S. [3] 11, 365).—3. Ozone rapidly forms POCl₃ (Remsen, l.c.).—4. Potassium chlorate produces POCl, -5. Sulphur trioxide reacts violently, forming POCl₃ and SO₂ (Michaelis, J. Z. 6, 239; 7, 110; Armstrong, J. pr. [2] 1, 255; cf. Rose, P. 44, 304).—6. Selenion dioxide produces POCl₃ and Se at 110°; if the SeO₂ is in excess, SeCl₂ and P₂O₃ are also formed; SO₂ has no action at 140° (M., l.c.).—7. Heated in a closed tube at 130° with arsenious oxide, forms As, AsCl₃, and P₂O₅ (Michaelis, l.c.).—8. Thionyl chloride forms PCl₅, POCl₃, and PSCl₃. Chromyl dichloride produces POCl3, P2O5, PCl5, and CrCl3. 9. Nitrogen trioxide or pentoxide reacts with strongly cooled PCl₃ to form NOCl, P₂O₅, POCl₃, and P₂O₅Ol₄ (Geuther a. Michaelis, B. 4, 766).— 10. Many metallic oxides withdraw Ol from PCl. forming metallic chlorides; PbO forms Pb and Pb(PO₃)₂ besides PbCl₂.—11. Many metallic sulphides when heated with PCl₃ react to form P_2S_3 and metallic chlorides; e.g. K_2S , BaS, CaS, Sb_2S_3 , PbS, &c. (v. Gm.-K. i. 2, 320).—12. Sulphides, PbCl phur reacts at c. 130° with production of PSCl, (Henry, B. 2, 638).—13. Antimony forms SbCl, and P (Gm.-K., l.c.).—14. Red-hot iron filings produce Fe phosphide and FeCl₂ (Gm.-K.).— 15. Zinc, at 100°, forms ZnCl₂ and P (Most, B. 18, 2029).—16. Platinum and silver are not acted on by PCl_s (Gm.-K.).—17. Sulphuric acid reacts slowly in the cold, rapidly when hot, to form SO₂, HCl, P₂O₅, and SO₂.OH.Cl (Michaelis, J. Z. 6, 239).—18. Phosphorous acid produces P₂O₅ (Nacquet's Chem. German ed. 1868. p. 218). 19. Hydriodic acid reacts at the ordinary temperature, forming PI, and HCl (Hautefeuille, Bl. [2] 7, 198, 200, 203).—20. PSCl, and PCl, are formed by heating for some hours at 160° with sulphur chloride. -21. Iodine pentabromide is said to form ICl, and PBr,.—22. Phosphorus hydride produces HCl and solid P,H (Besson, C. R. 111, 972). SbH, is said to be without action (Mahn, Z.[2]5,729).—23. Heated with hydrogen sulphide, HCl and P2S2 are formed (Baudrimont, A. Ch. [4] 2, 5).-24. Ammonia seems to form various compounds which have not been fully examined (v. Phospham, p. 104).— 25. Passed over heated ammonium chloride, phospham (q. v.) is formed.—26. With many carbon compounds containing hydroxyl, PCl. reacts to substitute Cl for OH.

Combinations.—1. With chlorine, to form PCl₃ (q. v.). — 2. With bromine to form several bromochlorides (q. v.).—3. With iodine

to form PI₂Cl₃ (v. Phosphorus iodochloride, p. 138).—4. With titanium chloride to form PCl₂.TiCl₄ (Bertrand, M. S. [3] 10, 1331); also with a few other metallic chlorides, e.g. AuCl.— 5. With dry ammonia to form PCl_s.5NH_s (Rose; Besson, C. R. 111, 972).

PHOSPHORUS PENTACHLORIDE PCl₅ (Phosphorus perchloride). Mol. w. 207.81. Melts at 148° under pressure; boils at 160°-165°, with partial decomposition to PCl, and Cl. V.D. varies with temperature; c. 103 in an atmosphere of PCl₃ (v. Properties). H.F. [P,Cl³] = 104,990 (Th. 2, 408).

Formation.—1. By combining PCl₃ with Cl,

or by the action of excess of Cl on P.-2. By reacting with PCl₃ on PH₃, S₂Cl₂, or SOCl₂.—3. By the action of SbCl₅ on PH₃ (Mahn, Z. [2] 5, 729).

Preparation.—PCl, may be prepared by the action of Cl on P, but the PCl, is apt to retain pieces of unchanged P, the presence of which may lead to violent explosions; it is, therefore, better to prepare PCl, from PCl, PCl, is placed in a strong flask with a wide neck; the flask is surrounded by a freezing mixture, and dry Cl is passed in by a wide tube which reaches almost to, but does not touch, the surface of the liquid. As the contents of the flask get semi-solid they must be thoroughly broken up by a glass rod. The whole of the PCl, is not changed to PCl, until Cl has passed for a considerable time; the change is not complete until, after closing the flask, shaking thoroughly, and allowing to stand for some time, the presence of greenish-yellow gas in the flask is quite apparent. Excess cf Cl is finally removed by the long-continued passage of a slow stream of dry CO₂ (Muller, Z. 1882. 295; Gräbe, Ar. Sc. [3] 5, 477).

Properties. — A white, or slightly yellow, crystalline solid with a strong odour; crystallises in rhombic plates; fumes in moist air, with decomposition; sublimes without melting under 100°, but melts at c. 148° under considerable pressure; boils at c. 160°-165°, with partial dissociation to PCl, and Cl, which re-combine on cooling. PCl, does not conduct electricity (Gm.-K.i. 2, 390). Decomposed by O at c. 300°.

Burns in a candle-flame.

Vapour density.-The S.G. of the vapour obtained by heating PCl, is less than that calculated for PCl_s, and it decreases as temperature increases until it is equal to half the calculated value. Wanklyn a. Robinson (C. R. 56, 547) showed that the vapour contained free Cl, by diffusing into CO2, and that PCl3 was present in the residue. Deville noticed that the vapour showed the colour of Cl, and that the intensity of colour increased as temperature rose (A. 141, 147). The V.D. was determined by Cahours at different temperatures, with the following results (A. 141, 42; v. also Gibbs, Am. S. [3] 18, 277,**371)** :--

Temp.	8.G. $(air = 1)$	P.c. Dissociation
182°	5.08	41.7
190	4.99	44.3
200	4.85	48.5
230	4.80	67.4
250	4.00	80.0
274	8.84	87.5
288	8.67	96.2
289	8.69	95.7
800	8.65	97.3

The temperature whereat 50 p.c. is dissociated is c. 202°. Wurtz (B. 3, 572) showed that, making certain assumptions, the S.G. of PCl, vapour formed in an atmosphere of PCl, is c. 7.4-6.8, which agrees with that calculated for

PCl₅, viz. 7.2.

Reactions .- 1. Heat dissociates PCI, into PCl, and Cl, which recombine on cooling (v. ante).--2. PCl, is decomposed by water to HClAq and POCl₃, and finally to H₃PO₄Aq with some HPO₃Aq (*Gm.-K.* i. 2, 390). Thomsen gives [PCl⁵,Aq] = 123,440 (*Th.* 2, 322).—3. Boric acid and antimonic hydrate react similarly to water (Gerhardt, A. 87, 66, 290; Schiff, A. 102, 111; 106, 116). Boric oxide when heated with PCl, forms B2O3.P2O5 and BCl3 (Gustavson, B.3, 426).-4. Hydroxides generally, including carbon compounds which contain the group OH, exchange OH for Cl, with production of POCl, and chloride of the radicle before in combination with OH .-5. Compounds containing the group SH generally react similarly to OH compounds; e.g. SH.H gives ClH and PSCl₃.—6. Oxides generally exchange O for Cl2; carbon compounds containing the group CO generally also react in this way. -7. Oxygen reacts with vapour of PCl, at c. 300°, producing POCl₃, P₂O₅, and Cl (Gm.-K. i. 2, 390; Wanklyn a. Robinson, C. R. 56, 547).— 8. PCl, vapour mixed with hydrogen and passed through a red-hot tube produces HCl, PCl, and P (Baudrimont, Gm.-K. i. 2, 394).—9. Heated with phosphorus PCl₃ is produced.—10. The compound PS2Cl, is formed by distilling PCl, with 8 parts of sulphur (Gladstone, C. J. 3, 5).—
11. Selenion produces Se.Cl. and PCl.—
12. Iodine forms PCl., and ICl which combines with part of the PCl, to produce PCl,.ICl (Gm.-K., l.c.).—13. PCl, reacts with many metals to form metallic chlorides and PCl3; sometimes P and metallic phosphides are formed (Baudrimont, J. pr. 87, 300; 88, 78; Casselmann, A. 98, 213). Among the metals which react with PCl, are Al, As, Cd, Au, Fe, Pt, K, Na, Sn, and Zn. The metallic chloride formed often combines with the undecomposed PCl₃; this occurs, e.g., with AlCl₃, FeCl₃, HgCl₂, PtCl₄, and SnCl₄ (Wöhler, B. 13, 875; cf. Goldschmidt, C. C. 1881, 489).—14. Phosphorus hydride produces PCl_s and HCl; with excess of PH_s, HCl, and P are formed.—15. Antimony hydride forms PCl., SbCl2, and HCl; silicon hydride acts similarly but more slowly (Mahn, Z. [2] 5, 729; ammonia forms phospham (q.v., p. 104) and nitrogen phosphochloride (vol. iii. p. 570); also phosphamidoimide (q. v., p. 105; cf. also Besson, C. R. 114, 1264).—16. Sulphur dioxide produces SOCl₂ and POCl₃ (Schiff, A. 102, 111; 106, 116; Persoz a. Bloch, C. R. 28, 86; Kremers, A. 70, 297).-17. Sulphur trioxide forms S2Cl2O, and POĆl₃ (Schiff, *l.c.*; Michaelis, J. Z. 6, 235, 240, 292; Williamson, Pr. 7, 11).—18. Selenion dioxide when distilled with PCl, forms POCl, SeCl₂, and P₂O₅, SeOCl₂ being formed as an intermediate product (Michaelis, Z. [2] 6, 465). 19. Phosphoric oxide produces POCl_s.—20. Boric oxide reacts slowly when heated to c. 140° with PCl_a; BOCl_a is probably formed, and on heating more strongly BCl, and B,O,P,O, remain (Gustavson, B. 3, 426).—21. Nitrogen dioxide produces NOCl and POCl, (Gm.-K. i. 2, 390).—22. Silicon dioxide forms POCl., and SiCl, which

combines with the excess of PCl, (Weber, P. 107, 375).-23. Vapour of PCl, reacts with many metallic oxides, when these are strongly heated, to form POCl, and metallic chlorides which frequently combine with undecomposed PCl. The following react in this way—Al₂O₃, As₂O₃, and As₂O₄ (Hurtzig a Geuther, A. 111, 159; Michaelis, J. Z. 6, 239), [no reaction with Sb₂O₈; Schiff, A. 102, 111], CdO, Cr₂O₃, Fe₂O₃, MnO₂, MoO₃ (Schiff, l.c.), SnO, TiO₂ (Weber, P. 107, 375; Tüttscheff, A. 141, 111), WO₃ (Gerhardt a. Chiozza, C. R. 36, 1050; Schiff, l.c.).—24. Phosphorus sulphide forms PSCl, (Weber, P. 107, 375; Gm.-K. i. 2, 390).-25. Carbon disulphide does not react at 100° (Hofmann, A. 115, 264); at 200° PSCl₃ is produced (Carius a. Fries, A. 112, 193), and CCl. (Rathke, Z. [2] 6, 57). — 26. Many metallic sulphides when heated are decomposed by vapour of PCl, with formation of PSCl, S2Cl, and metallic chloride; the following react in this way—sulphide of As, Ba, Bi, Cd, Ca, Pb, Na, Sn, Zn (Weber, l.c.; Baudrimont, l.c.). -27. Antimony selenide and lead selenide form SeCl₄, which forms a double compound with the excess of PCl₅.—28. Hydriodic acid produces PCl, HCl, and I (Wurtz, Gm.-K., l.c.); hydrobromic acid is without action (Gladstone, C. J. 3, 5); syrupy phosphoric acid solution reacts slowly, the vapours of this acid do not act (Gerhardt, A. 87, 66, 290; Schiff, A. 102, 111; 106, 116; Persoz a. Bloch, C. R. 28, 86).-29. Conc. nitric acid reacts energetically, forming POCl, and HCl.-30. Sulphuric acid forms SO2.OH.Cl, HCl, and HPO3, and S2Cl2O3 as the result of a secondary reaction (Michaelis, J. Z. 6, 235, 240, 292; Williamson, Pr. 7, 11; Baumstark, A. 140, 75).-31. Many metallic salts react with PCl,; KNO, gives NOCl and POCl, (Nacquet, Bl. 1860); sulphites form SOCl₂ (Buchanan, B. 3, 485); AgF produces AgCl and probably a fluoride of P (Pfaundler, W. A. B. 46, 258); KClO, gives POCl, KSCy forms PSCl, S2Cl, and PCl₃; Na₂HPO₁ gives H₃PO₄ and HCl; haloid salts of the alkalis, and also KCy and K₄FeCy₆, are not acted on by PCl₅.

Combinations.—1. With iodine chloride to form PCl₃.ICl (Gm.-K. i. 2, 390).—2. With selenion tetrachloride to form SeCl . 2PCl .. -3. With many metallic chlorides (cf. Reactions, Nos. 13 and 23). These compounds are solids, which sublime with partial decomposition when heated, and are decomposed by water (Baudrimont, J. pr. 87, 300; 88, 78; Weber, ibid. 77, 65; Casselmann, A. 83, 258).—4. With ammonia to form PCl₅.8NH₂; produced by slowly passing dry NH₃ into a solution of PCl₅ in CCl₄ (Besson,

C. R. 111, 972; 114, 1264).

Phosphorus, chlorobromides of, v. Phosphorus BROMOCHLORIDES, p. 130.

Phosphorus, chlorofluoride of, PCl,F2. w. 175.07. V.D. 78. Poulenc (C. R. 113, 75). Preparation.—Two stoppered flasks of c. 500

c.c. capacity are connected by a bent tube, which passes through the stoppers; one flask is filled with dry Cl, and the other with dry PF; that containing PF, is connected with a bulb full of Hg, which can be caused to flow into the flask and so force the PF, into the flask full of Cl. The PF, is slowly driven into the other flask, and the PCl.F, thus formed is kept in contact with Hg for some days (without being shaken with the Hg, else some PF, may be decomposed) to remove traces of Cl.

Properties — A colourless gas, with very irritating odour; incombustible; liquefied at —8° at ordinary pressure; absorbed and decomposed by water; heated to c. 250° gives PF₅ and PCl₅, also decomposed in the same way by electric sparks.

Reactions.—1. Heat produces PF₈ and PCl₅; the action proceeds at c. 250°.—2. Electric sparks also produce PF₈ and PCl₅.—5. Heated with sulphur to c. 115°, PSF₈ and S₂Cl₂ are formed.—4. Heated with phosphorus to c. 120° PF₃ and PCl₈ are formed.—5. Several metals when heated to c. 180° produce metallic chloride and PF₃, e.g. Al, Fe, Pb, Mg, Ni, and Sn; Hg exerts a slight action at ordinary temperatures, and acts rapidly at 180°; Na seems to absorb PCl₃F₂ entirely.—6. Water reacts rapidly; if the products are POF₃ and HCl; if the gas is passed into water, H₃PO₄Aq, HClAq, and HFAq are formed.—7. Absorbed by alcohol, with formation of a compound not yet examined.—8. Amnonia is absorbed at ordinary temperature, forming a white solid, which is probably PF₃(NH₂)₂.

Phosphorus, chloro-iodide of, v. Phosphorus

iodochloride, p. 138.

Phosphorus, chloronitride of, v. Nitrogen Phosphochloride, vol. iii. p. 570. Besson (C. R. 114, 1479) recommends to prepare PCl₂N by forming a layer of PCl₅ on the sides of a large flask, by acting on PCl₃ with Cl, then to heat the PCl₅ with NH₃ till it is superficially saturated, and then to heat under reduced pressure.

Phosphorus, compounds of, with hydrogen and oxygen. Those compounds of P, H, and O, which are acids, are described in the articles Phosphoric ACIDS (p. 124), and PHOSPHORUS, OXYACIDS OF, AND THEIR SALTS (p. 149); but two compounds which have been described by Gautier will find a place here. Gautier (C. R. 76, 49) says that the compound P.HO is formed by heating crystallised H₃PO₃ with 5 or 6 times its weight of PCl₃ to 79°, washing with water, and drying at -10° ; at between 80° and 100° the products of the reaction are H₃PO₄, HCl, and amorphous P. This compound is described as an amorphous yellow powder; unchanged in air when dry, but oxidised slowly when moist; insoluble in most menstrua; burns when heated in air to c. 260°; burns explosively when heated with CuO; unchanged at $240^{\circ}-250^{\circ}$ in CO_2 , but evolves PH₃ at 265°, and at 350°-360° ordinary P distils off; not acted on by dilute acids; oxidised by ordinary HNO, Aq; conc. H2SO, evolves SO, at c. 200° ; decomposed by H_{2}° at c. 170° to H_{2}° PO₂Aq, H_{2}° PO₃Aq, and PH_{3} ; reacts with dilute alkali solutions; NH_{3} combines to form a very unstable brown substance. Gautier (C. R. 76, 178) described another compound P₆H₃O; obtained by adding PI2, little by little, to H2O at 80°-90°, allowing the yellow solution to deposit a flocculent pp., washing this with warm water, and drying in vacuo. If water is added slowly to PI₂ the products are HIAq, H₃PO₃Aq, and H₂PO₂Aq. P₃H₃O is described as a pure yellow solid; amorphous, or perhaps showing traces of crystalline form; tasteless and odourless; insoluble in all menstrua; oxidises in air when moist, when dry oxidises slowly at 100°; oxidised violently by HNO₂Aq; evolves SO₂ from H₂SO₄; gives off PH₂ when heated to 135° in CO₂, and ordinary P at 350°; decomposed by dilute alkali solutions to PH₃, H₁H₂KPO₂, and HK₂PO₄; combines with NH₃ to form a brown very unstable substance.

Phosphorus, cyanide of, v. Cyanogen phosphide, vol. ii. p. 358.

Phosphorus, fluorides of. Two fluorides of P are known, PF_s and PF_s; these formulæ are molecular. The compounds are gases at ordinary temperatures.

PHOSPHORUS TRIFLUORIDE PF₃. (Phosphorous fluoride.) Mol. w. 87.96. V.D. 43.7 (Moissan, A. Ch. [6] 6, 433).

Formation.—1. By the reaction of PbF₂ with Cu phosphide (M., l.c.).—2. By adding AsF₃ to PF₃ (M., l.c.).—3. By the reaction of PbF₂ with PCl₃ or POCl₃ (Güntz, C. R. 103, 58).—4. By adding PBr₃ to gently heated ZnF₂ (M., A. Ch. [6] 19, 286).

Preparation.—1. AsF₃ is dropped into perfectly dry PCl₃; the gas is shaken with a little water, and dried over H₂SO₄ (M., C. R. 100, 272).

2. A mixture of well-dried Cu₃P₂ and PbF₂ free from SiO₂ is heated in a brass tube to dull redness, the gas is passed by a leaden tube through two very small bottles (2 or 3 c.c. capacity) containing H₂SO₄, then through pumice soaked in H₂SO₄, and is then collected over Hg.—3. PBr₃ is gradually added to warm ZnF₂, the gas is washed by passing through water, then dried by passing over pumice soaked in H₂SO₄, and collected over Hg (M., A. Ch. [6] 19, 286).

Properties.—A colourless gas, condensed to a colourless liquid at -10° and 40 atmos, pressure. The gas is not liquefied at 24° under a pressure of 180 atmos, but liquefaction occurs when the pressure is suddenly reduced to 50 atmos. (Moissan, l.c.). PF₃ does not fume in the air; it is very slowly decomposed by water, with formation of H_3PO_3Aq and HFAq; burns when mixed with O; decomposed by heat; also by

electric sparks.

Reactions and Combinations.-1. Decomposed by heat; in a glass apparatus P and SiF. are formed; in contact with Pt black a gas is formed, probably containing F (Moissan, C. R. 102, 763).—2. Electric sparks slowly decompose PF_s, forming PF_s and P (Moissan, C. R. 102, 763).—3. Burns when mixed with pure oxygen and ignited; a mixture of 1 vol. PF, and ½ vol. O explodes violently when sparked, with formation of POF₃ (M., C. R. 102, 1245).—4. Very slowly decomposed by water, with formation of H₃PO₃Aq and HFAq; more rapidly decomposed by steam at 100°.—5. According to Berthelot (A. Ch. [6] 6, 358) potash solution produces a fluophosphorous acid probably analogous to fluoboric or fluosilicic acid; Moissan (C. R. 99, 655) says that a fluoride and a phosphite are formed.—6. Absorbed rapidly by bromine with formation of PBr₂F₈ (v. Phosphorus bromofluoride, p. 131), also by chlorine, with formation of PCl₂F₃ (v. Phosphorus chlorofluoride, p. 133).—7. Decomposed rapidly by solutions of chromic acid or potassium permanganate.—8. Decomposed by hot sodium, copper, boron, and silicon (M., C. R. 99, 655).—9. Combines with ammonia to form white, flocculent compound, which is decomposed

by water (M., l.c.).--10. Combines with fluorine to form PF (Moissen, RI [3] 5, 454)

to form PF, (Moissan, Bl. [3] 5, 454).

Determination of composition.—The compound was analysed by Moissan (C. R. 100, 272) by heating a known volume in a glass vessel, measuring the volume of SiF, formed, dissolving the P deposited in HNO, Aq, and estimating as Mg-NH, phosphate.

PROSPHORUS PENTAFLUORIDE PF. (Phosphoric fluoride). Mol. w. 125.96. V.D. 63.2 to 65.1 (Thorpe, A. 182, 201; Moissan, C. R. 102, 763;

103, 1257).

Formation.—1. By passing PF₈ into Br at —15°, PBr₂F₃ is formed, and when heated this gives PF₅ and PBr₅ (Moissan, C. R. 101, 1490).
2. By the reaction of AsF₃ with PCl₅.—3. By direct combination of PF₃ with F (M. Bl. [3] 5, 454).

Preparation. — AsF₃ is added gradually to PCl, surrounded by a freezing mixture; the gas which comes off is passed through PCl, contained in a tube which is kept cold, and is collected over

Hg (Thorpe, A. 182, 201).

Properties and Reactions. - A colourless, strongly-smelling gas, which fumes in air and rapidly attacks the membranes of the mouth and the bronchial tubes. Condensed at 15° and 46 atmos. to colourless liquid which does not act on glass (Moissan, C. R. 101, 1490); on partially releasing pressure the liquid solidifies, but soon again becomes liquid (M., l.c.). Decomposed by powerful electric sparks (150-200 mm. long) to PF_s and F (M., C. R. 103, 1257), but not by ordinary sparking (Thorpe, l.c.). Incombustible and does not support combustion. Not acted on by heating with phosphorus to dull redness, nor with sulphur vapour at 440°, nor iodine at 500° (M., l.c.). In presence of trace of water acts on glass, forming SiF, and POF, Passed over gently-heated spongy platinum is partially decomposed to PF, and F; when the Pt is heated to dull redness a compound of Pt with P and F, probably PF₃.PtF₂, is formed (M., Bl. [3] 5, 454).

Phosphorus, fluobromide of; v. Phosphorus

BROMOFLUORIDE, p. 131.

Phosphorus, fluochloride of; v. Phosphorus

CHLOROFLUORIDE, p. 133.

Phosphorus, haloid compounds of. P combines readily with the halogens, with production of much heat, to form compounds PX₈ and PX₅; besides these, the compound PI₂ is also known. Several compounds of the type PX₅, and at least one of the type PX₅ (viz. PF₆), have been gasified, so that the formulæ are molecular. The haloid compounds of P are decomposed by water, the fluorides only very slowly, with formation of oxyacids of P and halogen acids; these compounds also combine with many other haloid compounds, both of metals and non-metals, to form double salts.

Phosphorus, hydrides of. P and H do not combine directly; but hydrides of P are produced by evolving H in contact with phosphites or hypophosphites. Three hydrides of P have been isolated; PH₃, PH₂ (or P₂H₄), and P₂H (or P₄H₂). Janssen (Repert. Chim. app. 3, 393) obtained a crystalline, explosive substance by the reaction of milk of lime with P, to which he gave the formula P₂H₃. At the ordinary temperature and pressure the first hydride is gaseous, the second liquid, and the third solid. The formula PH₂ is molecular; Croullebois (C. R. 78, 496)

said that the V.D. of the liquid compound shows the mol. w. to be P₂H₄, but no details are given; the mol. w. of the solid compound is unknown. The liquid hydride takes fire in contact with air; the gas inflames in air at c. 150°, and the solid at c. 200°; the processes for preparing gaseous PH₄ generally also produce more or less liquid P₂H₄, the resultant gas is therefore spontaneously inflammable. PH₅ resembles NH₃ in its reactions, but it is much less decidedly alkaline; phosphonium compounds, e.g. PH₄I, similar to ammonium compounds, are known.

PHOSPHORUS TRIHYDRIDE PH, (Phosphoretted hydrogen. Phosphine). Mol. w. 33 96. Liquefies - 90°; solidifies - 133 5°; melts at -132 5°; boils c. -85° (Olszewski, M. 7, 371). S.G. (air = 1) 1 185. V.D. 17 2. H.F. [P,H] = 36, 600 (Ogier, C. R. 87, 210). S. 112 (Dy-

browski, J. 1866. 735).

Formation. - The gas obtained by the following methods is never pure PH,; it contains more or less P2H4, and generally also H; it is usually spontaneously inflammable. evolving H in contact with phosphites or hypophosphites in solution; phosphates are not reduced (Gm.-K. i. 2, 138; Fresenius, Fr. 6, 203; Herapath, Ph. 7, 57). P and H do not combine directly (Fourcroy a. Vauquelin, A. Ch. 21, 202; Dusart, C. R. 43, 1126).—2. By heating solutions of phosphites or hypophosphites (Gm.-K., l.c.).-3. By the action of boiling alkali solutions on P (H. Nose, P. 6, 199; 8, 191; 14, 183; 24, 109, 295; 32, 467; 46, 633).—4. By heating P with syrupy H3PO4Aq, or with HClAq, HBrAq, or HIAq (Oppenheim, Bl. [2] 1, 163).—5. By heating P with water to 200° (Oppenheim, l.c.).—6. By long-continued heating P with blood to 35°-41° (Dybrowski, J. 1866, 735).—7. By the action of the electric current on moist molten P (Groves, C. J. 16, 268).—8. By heating P with H₂O to c. 200°.—9. By decomposing Zn phosphide by dilute acids, or boiling alkali solution (Schwarz, D. P. J. 191, 896). According to Lüpke (C. C. 1890. ii. 642), the gas obtained by acting on phosphide of Mg, Sn, or Zn with dilute acid is not spontaneously inflammable. — 10. By treating Cu phosphide with KCN and a little 80 p.c. alcohol (Gm.-K. i. 2, 138).—11. By decomposing phosphides of the alkali or alkaline earth metals by water or dilute acids.

Preparation .- 1. A small flask carrying an exit tube is nearly filled with a conc. solution of KOH in 80 p.c. alcohol; small pieces of P are dropped in, and the flask is heated; the gas is passed through a CaCl₂ tube to absorb alcohol vapour, and collected over water. The gas thus prepared does not contain more than c. 45 p.c. PH, (Hofmann, B.4,200); it is generally free from the spontaneously inflammable hydride. The gas prepared by the action of KOHAq on P contains some P₂H₄ and much H; it is spontaneously inflammable in air. The simplest way of preparing this gas is to place a few small pieces of P in a little flask which carries a cork with two tubes, one fitted with a stopcock and passing just through the cork, and the other bent like an ordinary gas-delivery tube and dipping under water in a basin; the flask is nearly filled with conc. KOHAq; the tube with the stopcock is attached to the gas-supply, the cork is fitted loosely into the flask, and a stream of coal gas is passed

through the apparatus for a few minutes, after which the cork is fitted tightly into the flask, the stopcock is closed, and the flask is heated. During this process the end of the delivery-tube is kept under the surface of the water in the basin. After a few minutes gas is evolved, and each bubble takes fire as it comes into contact with the air after passing upwards through the water in the basin. If the whole of the air is not driven out of the apparatus before the flask is heated, serious explosions may occur.—2. The best way of preparing pure PH3 is to decompose PH4I by KOHAq. A small flask is fitted with a caoutchouc cork, which carries a tube furnished with a bulb and stopcock and an exit-tube; pieces of PH.I about the size of peas are placed in the flask along with several small pieces of glassrod, KOHAq (c. 1 pt. KOH in 2 pts. H₂O) is allowed to drop from the bulb-tube very slowly on to the PH,I in the flask. Pure PH, is evolved without heating; if the KOHAq is added too rapidly the gas may contain a little P2H4 and be spontaneously inflammable (Hofmann, B.4,200; Rammelsberg, B.6,88). About 1 litre PH₄ is obtained from $7-7\frac{1}{2}g$. PH₄I (for preparation of PH,I v. Phosphonium compounds, p. 123).

Properties.—PH3 is a colourless gas, with a very disagreeable smell like that of onions mixed with decaying fish; very poisonous; slightly soluble in water (v. beginning of this article). The solution glows in the dark, and deposits amorphous P when exposed to light and air; on boiling, PH3 is given off. PH3 is slightly soluble in alcohol, ether, ethereal oils, and blood; it is completely absorbed by solution of bleaching powder, also by Cu2Cl2 in HClAq (v. Combinations, No. 4). PH3 combines with the halogen acids to form salts similar to those of NH, (v. Phosphonium compounds, p. 123); the H of PH₃ is replaceable by organic radicles (v. Phosphines, PH₃ reduces solutions of salts of heavy PH₃ free from P₂H₄ takes fire in air at p. 116). c. 150°. The gas prepared as mentioned under Formation takes fire immediately on contact with air; it loses this spontaneous inflammability by exposure to sunlight, by passage through a U-tube cooled to -10° , by mixing with small quantities of Al_2O_3 , wood charcoal, various gases, &c. (v. Graham, P.M.5, 401). These processes remove P2H4, to the presence of which the ready inflammability of the gas is due. Addition to PH₃ of $\frac{1}{500}$ of its weight of P₂H₄ causes the gas to become inflammable in air; the presence of $\frac{1}{1000}$ to $\frac{1}{10000}$ of NO₂ produces the same effect (Graham, l.c.; cf. Landolt, A. 116, 193). The gas prepared by the action of KOHAq on PH_{*}I (v. Preparation, No. 2) is described by Hofmann as taking fire when gently warmed, and being sometimes inflamed by the friction of the stopper of the bottle in which the gas may be kept. According to Rammelsberg (B. 6, 88), the gas prepared in this way is sometimes spontaneously inflammable. The gas prepared in this way takes fire when passed into conc. AgNO, Aq (Poleck a. Thümmel, B. 16, 2442).

Reactions.—1. PH, is decomposed by heat; when passed through a red-hot tube, lustrous, amorphous P is deposited (Merz a. Weith, B. 18, 718).—2. Decomposed by electric sparks to

P and H; the process takes place with regularity, 20 c.c. are decomposed in 5-6 mins. with production of P and 30 c.c. H (for description of an apparatus for lecture purposes v. Hofmann, B. 4, 204).—3. PH, burns in air at c. 150° with production of P2O, and H2O; if a cold substance is brought into the flame, yellow-red specks of amorphous P are deposited.—4. No reaction occurs when pure PH3, free from P2H4, is mixed with oxygen, but the mixture explodes if the pressure is suddenly lowered; explosion is said to occur after some hours at the ordinary pressure (Labillardière; v. Gm.-K. i. 2, 144).— 5. Bubbles of PH, burn explosively to PCl, and HCl if passed into a vessel filled with chlorine; if the Cl is diluted with CO2, the action is slower, and some P separates.—6. Bromine and iodine react similarly to Cl; if PH₃ is heated gently with iodine, some PH₄I is formed.—7. Heated with sulphur, H₂S and P sulphide are formed (Jones, C. J. [2] 14, 648).—8. Salts of the alkali metals and of the alkaline earth metals decompose PH_s, producing phosphites and hypophosphites (Winkler, P. 111, 443).—9. Solutions of salts of heavy metals generally decompose PH_s, with production of metallic phosphides, which sometimes combine with the excess of the metallic salts (v. Kulisch, A. 231, 327). AgNO3Aq forms a yellow compound (? Ag_sP.3ÁgNÖ_s), and then black Ag,P which is quickly reduced to Ag (Poleck a. Thümmel, B. 16, 2442); HgCl.Aq, according to Ashan (Chem. Zeitung, 10, 82, 102), forms a yellow compound 3Hg₃P₂,7HgCl₂, a red compound 4Hg₃P₂.5HgCl₂, and a brown compound Hg₃P₂.HgCl₂; PtCl₄ is said to form Pt.PH₂ (Gavazzi, B. 16, 2279); chlorides of Cr, Co, Cu, Au, Fe, and Ni are reduced, with formation of phosphides of the metals, or of metal and P .-10. Potassium permanganate solution is reduced by PH₂ to Mn₂O₃, K₂HPO₄ being also formed (Gavazzi, B. 16, 2279).—11. PH₂ is decomposed by many oxides, acids, and salts, e.g. by H₂S, SO₂, NO, N₂O, PCl₃, HNO₃ (no reaction at -25°; Besson, C. R. 109, 644), H₂SO₄ (v. Besson, l.c.), SbCl., &c., also by certain metals, e.g. Sb, Cu, Fe, K, and Zn (v. Gm.-K. i. 2, 138).—12. With haloid compounds of arsenic PH, reacts to form haloid acid and phosphide of As (Besson, C. R. 110, 1258).-13. With an ethereal solution of bismuth bromide forms a black lustrous body, probably PBrH(BiBr₂)₂ (Cavazzi a. Tivoli, G. 21, ii. 306).

Combinations.—1. With the halogen acids to form compounds of the type PH₄X (X=Cl, Br, or I); combination with HI and HBr occurs at the ordinary temperature and pressure, with HCl combination occurs at -30° to -35° under the ordinary pressure, or at 14° under a pressure of 20 atmos. (Ogier, Bl. [2] 32, 483; v. also Skinner, Pr. 42, 283). At very low temperatures PH₃ seems to combine with H₂SO₄ (v. Phosphonium sulphate, p. 124). The compounds PH₄X are described under Phosphonium compounds, p. 123.—2. When PH₄ is strongly compressed in presence of water, the gas liquefies and floats on the water. If the pressure is now suddenly released, a white crystalline solid is formed, but this decomposes again when the pressure is reduced below a certain amount (Cailletet a. Bordet, C. R. 95, 58; v. Phosphonium compounds, p. 123).—3. White compounds, easily

decomposed on removing pressure, are said to be formed by compressing PH, with carbon dioxide, and with carbon disulphide, in presence of water (C. a. B., l.c.).—4. A solution of cuprous chloride in HClAq absorbs PH3 rapidly; a white crystalline mass of Cu₂Cl₂·2PH₃ is formed, but this liquefies as more PH₃ is passed in (? with formation of Cu₂Cl₂·4PH₃); the crystals Cu₂Cl₂.2PH₃ are decomposed by warming, or by addition of water, to PH3, HCl, and Cu3P; the liquid compound loses PH, in a stream of an indifferent gas, with production of the crystalline compound (v. Riban, Bl. [2] 31, 385). This behaviour of PH₃ towards Cu₂Cl₂ gives a method for preparing pure PH₃, and also for estimating PH, in a mixture of gases.

PHOSPHORUS DIHYDRIDE PH₂ or P₂H₄ (Liquid phosphoretted hydrogen). Mol. w. said to be 65.92, corresponding with V.D. c. 33, but no details given (Croullebois, C. R. 78, 496). Gattermann a. Hausknecht (B. 23, 1174) failed to determine V.D. as the compound always decomposed. A spontaneously inflammable gas containing P and H was obtained by Gengembre in 1783 (Crell's Ann. 1789. 450). The fact that this gas lost its inflammability by standing over air containing water was explained by Dumas (A. Ch. [2] 31, 113) by supposing that the gas was a mixture of two hydrides of P, one only of which was spontaneously inflammable; to one of these Dumas gave the formula PH2 and to the other the formula PH_s. In 1832 H. Rose (P. 6, 199; 8, 191) thought that the two gases had the same composition, but were isomeric one with the other. After Graham (P. M. 5, 401) had the other. After Graham (P. M. 5, 401) had examined the conditions under which spontaneous inflammability was removed from or bestowed on the gas, Le Verrier (A. Ch. [2] 60, 174) came to the conclusion that the gas consisted chiefly of PH, but that it contained also a very inflammable compound PH₂, which was decomposed by light into PH₃ and a solid hydride PH. În 1845 P. Thénard made a fuller study of the subject (A. Ch. [3] 14, 5); he showed that spontaneously inflammable phosphoretted hydrogen, when passed through a tube cooled to under 10°, deposited a liquid having the composition PH2, which liquid was extremely inflammable, and that this liquid decomposed in light into gaseous PH, and a solid to which he gave the formula P.H. In 1874 Croullebois said that the V.D. of the liquid hydride corresponds with the formula P₂H₄, but no details were given (C. R. 78, 496); in 1890 Gattermann a. Hausknecht attempted to determine V.D., but found that the compound decomposed (B. 23, 1174).

Preparation.—Impure Ca phosphide is prepared by heating dry CaO to bright redness in a crucible, and adding dried P little by little, the lid being replaced after each addition of P. The crucible should be arranged so that the P is brought under the strongly-heated CaO; this may be done by placing a few pieces of P in a crucible, filling up with dry CaO, covering tightly (but leaving one small hole), placing the crucible in a furnace arranged so that the lower part of the crucible projects downwards through the bottom of the furnace, heating the upper part of the crucible to bright redness, and then warming the lower part so as to vapourise the P. Small quantities are prepared by placing a dry piece of tube filled with Hg: 100 parts liquid phos-

P in the closed end of a tube of hard glass, partly filling the tube with dry CaO, arranged so that there is a little space between the CaO and the P, placing the tube in a combustion furnace, heating the CaO strongly, and then warming the P. The CaO is transformed into a mixture of Ca phosphide and phosphate; the phosphide is nearly black; the unchanged CaO is picked out when the action is over, and the phosphide is at once placed in a stoppered bottle. (For modification of process \boldsymbol{v} . Gattermann a. Hausknecht, B. 23, 1174.) A three-necked bottle of about 2 litres capacity is filled three-quarters with water; one of the necks carries the tube from a H apparatus, which tube dips considerably under the water, the middle neck carries a piece of tubing about 15 mm. internal diameter, dipping 2 or 3 cm. under the water, and the third neck carries an exit tube in communication with a condenser; between the condenser and the three-necked bottle is placed a test tube, to collect part of the water vapour coming over. The condenser consists of a tube about 100 mm. long and 30 mm. diameter, narrowed so that the lower end forms a tube about 40 mm. long by 12 mm. wide; this condenser is closed by a cork through which pass the entrance tube from the threenecked bottle and an exit tube, which is bent downwards and dips under the surface of water; these two tubes are cut off obliquely just below the cork. The condenser is placed in a deep vessel full of cold water which contains sufficient ice to keep the temperature at 0°. H is passed through the apparatus until all the air is driven out, the three-necked bottle is placed in a waterbath until the contents are warmed to 60°, and Ca phosphide, in pieces the size of peas, is dropped down the wide tube in portions about 2 g. at a time and at a rate such that 50 g. are added in 15 to 20 minutes. Gaseous P hydride passes off and bubbles up through the water, and the liquid hydride collects in the condenser, about 1-2 c.c. being obtained for 50 g. Ca phosphide used (Hofmann, B. 7, 531; Gattermann a. Hausknecht, B. 23, 1174; the many precautions to be taken in working with this compound are described by G. a. H.).

Properties and Reactions. - A colourless, highly refractive liquid; insol. in water; burns in air, with bright flame to P₂O₅ and H₂O; does not solidify at -10°; boils 57°-58° at 735 mm. (G. a. H., l.c.). S. G. 1.007 to 1.016 (G. a. H.). Decomposed by light into PH, and solid P2H; the liquid may he kept for a short time in sealed tubes, but decomposition soon begins, and the tubes explode violently (G. a. H.). The same decomposition is effected by conc. HClAq, air mixed with CO₂, &c. Passage of the gas prepared as described above through a little conc. HClAq removes every trace of PH₂, and hence causes the gas which issues to be non-inflammable. If CO2 is passed through the U-tube containing liquid PH2 a nearly invisible greenish light appears at the end of the tube; this flame is not hot enough to ignite a candle (v. Hofmann, B. 7, 531; Thénard, A. Ch. [8] 14. 5; Bonet a. Bonfil, J. pr. 55, 247; Gattermann a. Hausknecht, B. 23, 1174). The composition of PH₂ was determined by Thénard by decomposing a known quantity by direct sunlight in a graduated

phide gave 61.8 parts gaseous PH, and 38.2 parts solid P_2H ; i.e. 1 molecule P_2H was produced for 3 mols. PH₃, hence the equation FPH₂=P₂H+3PH₃. This equation was confirmed by G. a. H., who made also direct estimation of H by burning with PbCrO₄ in a current of CO_2 (B. 23, 1174).

DIPHOSPHORUS HYDRIDE P2H or P4H2. Mol. w. unknown. H. F. $[P^2,H] = 17,700$ (Ogier, C. R.

89, 707).

Formation -1. By decomposing phosphide of Ca or K (Magnus, P. 17, 527) by HClAq.— 2. By the action of light, HClAq, Cl (Le Verrier, A. Ch. [2] 60, 174), and various other reagents, on PH2.—3. By the decomposition of PI2 by water (Rüdorff, P. 128, 473). Hittorf (P. 126, 193) obtained only amorphous P by this process.

Preparation. — A two-necked bottle arranged with an exit tube dipping under water, and a wide entrance tube, down which is passed the delivery tube of a H-apparatus; conc. HClAq is placed in the bottle, H is passed through until all air is driven out, the H apparatus is removed, and small pieces of freshly-prepared Ca phosphide (for preparation v. Preparation of phosphorus dihydride, p. 137) are dropped slowly into the bottle. The P2H which forms in the bottle is rapidly washed with cold water, and dried in vacuo (P. Thénard, A. Ch. [3] 14, 5).

Properties. - A yellow flocculent powder, which becomes orange-yellow in light; tasteless and odourless. When dry may be heated to c. 200° before taking fire; ignited by a blow of a hammer.

Reactions.—1. Ignited by heating in air to c. 200°; also by a blow of a hammer.—2. Slowly decomposed in *moist air*, especially if in sunlight, to P₂O₅ and H.—3. Distillation in hydrogen produces PH, (Thénard, l.c.).-4. Chlorine forms PCl, and HCl.-5. Dissolves in dilute nitric acid; oxidised with ignition by conc. HNO, Aq. -6. Alcoholic solution of potash produces PH, and H; addition of water to the reddish solution ppts. yellow flocks of? amorphous P or a lower oxide of P.—7. Explodes when mixed with potassium chlorate, silver oxide, mercuric oxide, or cupric oxide, and struck or warmed. -8. Ppts. metallic phosphides from solutions of salts of many heavy metals (Le Verrier, l.c.).

Determination of composition. - The quantity of P in the solid hydride was determined by Thénard by heating with a weighed quantity of PbO, and weighing the mixture of PbO and Pb phosphate thus produced; the H was determined by decomposing by red-hot Cu, and collecting the H set free. Rüdorff decomposed the hydride by heat, and collected and analysed the mixture of PH, and H thus formed; he also oxidised the compound by HNO₈Aq, and estimated P as Pb

phosphate.

Phosphorus, iodides of. P and I combine directly, even at -24°; two iodides, PI, and PI, have been isolated, and the existence of a

third, PI, is probable.

PHOSPHORUS DI-IODIDE PI2 or P.I4. Mol. w. not known. H.F. [P,I2] = 9,880 from solid P and I; 20,680 from solid P and gaseous I (Ogier, C. R. 92, 83).

Formation .- 1. By adding I to P Combination occurs even at -24° , with production of much heat, and ignition of the excess of P if air is admitted; amorphous P is produced be-

sides PI₂ (v. Wurtz, A. Ch. [3] 42, 129).—2. By the reaction of I with PH₃; also of I in acetic acid on POl₃ (Hofmann, A. 103, 305; Ritter, A. 95, 210).

Preparation.—One part by weight of P is dissolved in CS₂, and S₂ parts of I are added little by little. When the colour of the dark brownish-red liquid thus produced has changed to orange it is cooled to 0° for some hours, when the vessel becomes filled with crystals; the crystals are freed from CS2 by heating in a water-bath, while a stream of dry air is passed over them (Corenwinder, A. Ch. [3] 30, 242). The crystals may also be obtained by evaporating the solution in CS, in a stream of CO, (Berthelot a. Luca, C. R. 39, 748). Corenwinder says that from 2 to 3 g. P, and 60 to 75 c.c. CS2, are convenient quantities to use.

Properties and Reactions.-Large orangecoloured prisms, melting at 110° (Corenwinder, l.c.). Heated in O gives P₂O₅ and I (Berthelot, C. R. 86, 628, 787, 859, 920; 87, 575, 667). Decomposed by H₂O to H₃PO₃Aq, PH₃, HIAq, and a yellow flocculent solid supposed to be amorphous P (Hittorf, P. 126, 193), regarded by Rüdorff as solid P4H2 (P. 128, 473), and said by Gautier (C. R. 76, 49, 173) to be P.H.O.

PHOSPHORUS TRI-IODIDE PI, (Phosphorous iodide). Mol. w. 410-55. H.F. [P,I*] = 10,900 (Ogier, C. R. 92, 83). This compound is obtained in large red prisms by dissolving one part of P in CS2, adding a solution of 121 parts of I in CS2, concentrating much out of contact with air, and then placing in a freezing mixture. As the crystals are very sol. CS2, they should be at once freed from mother-liquor, and dried in a stream of dry air at c. 50°. The crystals melt at 55°, and boil at a higher temperature, with evolution of I. PI, is very deliquescent; moist air decomposes it at once to H,PO,Aq and HIAq (Corenwinder, A. Ch. [3] 30, 242).

PHOSPHORUS PENTA-IODIDE PI, (Phosphoric iodide). Isolation doubtful. Hampton (C. N. 42, 180) obtained a dark-crimson, very deliquescent solid, giving numbers on analysis agreeing with 80 p.c. $PI_5 + 20$ p.c. PI_8 , by dissolving P in a little CS₂ in a stream of pure dry N, adding rather more than enough I to form PI_s, distilling off CS₂ (in a N stream) at 45° under reduced pressure, and then warming to 50°; at 55°, under the reduced pressure, I began to

sublime.

Phosphorus, iodo-chloride of, PI2Cls. Produced by adding much I to a little PCl, allowing to stand in moist air for some days, drying the crystals in a stream of air, dissolving in CS₂, and crystallising: large, red, six-sided crystals, very hygroscopic; decomposed by water to HsPOsAq, HClAq, and HIAq (Moot, B. 13, 2029).

Phosphorus, nitride of. No compound of

P and N has been isolated with certainty. substance supposed to be a nitride by Rose (P. 24, 308; 28, 529) and Wöhler a. Liebig (A. 11, 139) was found to contain H. Briegleb a. Geuther (A. 123, 236) think that a compound P₃N₅ is perhaps produced by the reaction of PCl₅ on hot Mg₅N₂ in an atmosphere of N.

Phosphorus, oxides of. P and O combine very readily, with production of much heat; four compounds are produced according to the conditions: P₄O, P₂O₂, P₂O₄, and P₂O₅; the V.D. of the second of these shows that its molformula as a gas is P_4O_6 . The oxides P_2O_2 and P_2O_5 are anhydrides; the former reacts with water to produce the acid H_3PO_2 , and the latter to produce three acids, HPO_2 , H_2PO_4 , and $H_4P_2O_7$; the oxide P_2O_4 is not the anhydride of a corresponding acid—with water it produces the two acids H_3PO_3 and H_3PO_4 .

The only oxide formed when P glows in air at ordinary temperature is P₂O₅, but P₂O₅ is formed by drawing air over P without the latter glowing (Thorpe a. Tutton, C. J. 57, 573); at 50°-60° small quantities of P₂O₅ are produced, and this oxide is produced in larger quantities when the P is actually ignited; at moderately high temperatures and with a limited supply of air the oxide P₄O is formed in addition to P₂O₅ and P₂O₅. If the products of the slow burning of P in dry air are heated together in CO₂, the oxide P₂O₄ is formed, along with some P₄O. The products of burning P in a limited supply of dry air generally contain small quantities of P; this may be removed by sucking the products through a tube heated by steam (Thorpe a. Tutton, C. J. 49, 833).

PHOSPHORUS SUBOXIDE P.O. Mol. w. uncertain. Some doubt still remains concerning the composition of the lowest oxide of P; but the evidence is in favour of the existence of a definite oxide P₄O. Le Verrier (A. 27, 167) obtained a yellowish solid by exposing P in PCl₃ or in ether to air, warming the solid thus formed with water, and drying over H₂SO₄. A similar product was obtained by the incomplete combustion of P in air, or by the action on P of such oxidisers as iodic or periodic acid in presence of water (v. Gm.-K. i. 2, 107). This substance was generally looked on as an oxide of P; Le Verrier gave it the formula P₄O; Schrötter (W. A. B. 8, 246) thought it was only a mixture of ordinary and amorphous P. Reinitzer a. Goldschmidt (B. 13, 845) obtained the oxide P₄O by heating POCl₃ with P to 200°-250°, also by heating POCl₃ with Zn, Mg, or Al at 100°. Thorpe a. Tutton (C. J. 49, 833) found that P₄O is one of the products of burning P in a limited supply of air at moderately high temperatures, and that it is formed when the products of the slow burning of P in dry air are heated to c. 300° in CO₂. The experiments of R. a. G. indicate the existence of two varieties of P₄O; one of these reacts with hot alkali solutions, evolving PH3, and reduces salts of Au and Ag and mercurous salts, and the other is not acted on by alkalis, and does not reduce salts of Au, Ag, or Hg.

Preparation and Properties of P₄O which reduces salts of Ag, Au, and Hg.—1. Thin plates of pure Zn are heated with POCl₃ in a sealed tube in a water-bath at 100° for some hours, the liquid, with suspended solid matter, is poured off, the solid is allowed to settle and the liquid is decanted off; the solid is heated as before for some time with POCl₃ (to remove any Zn present), the liquid is poured off, the solid is washed with CHCl₃ to remove POCl₃, then with HClAq, and finally with water; the solid is then dried for some days in vacuo (Reinitzer a. Goldschmidt, B. 13, 849); the other products of the reaction are ZnCl₂ and Zn2PO₃. As thus prepared P₄O is an orange-red powder. It very obstinately retains water, the sample made by R. a. G. con-

tained c. 8 p.c. water, but the P and O were in the ratio P₄O. This oxide in moist air evolves PHs; it reacts with boiling alkali solutions forming PH, and alkali phosphite and phosphate; solutions of salts of Au and Ag, and mercurous salts, are reduced with ppn. of the metals; heated in H, P is given off, and P₂O₅ remains.—2. POl₃ is placed in an open flask, of c. 1 litre capacity, containing a layer about 2 cm. thick of P in small pieces; the PCl_s just covers the P; after c. 2 days the PCl_s is poured off, the pieces of P and the adhering yellow film are loosened from the flask, and allowed to fall slowly into cold water (if water is added in quantity, heat is produced, and the products of the slow oxidation of P are decomposed). After a little, the water is filtered from suspended P, and the clear yellow filtrate is heated to 80° when a very finely divided yellow solid settles down; this solid is washed with warm water in a filter, removed from the filter while moist, and placed in a basin, over H.SO, in vacuo. Le Verrier's P₄O was tasteless and odourless, did not change in dry air or O, but in moist air it gave off PH₃; it was decomposed somewhat above 360°, giving off P and leaving P_2O_5 ; oxidised by Cl to PCl, and P2O5; heated with conc. H₂SO₄, gave SO₂; unacted on by HClAq; oxidised and dissolved by HNO_sAq ; exploded with $KClO_s$; readily combined with H_2O to form $P_4O.H_2O$. The oxide obtained by Thorpe a. Tutton (C. J. 49, 833) was formed by slowly burning P in dry air, transferring the products to a tube filled with CO2, exhausting by a Sprengel pump, and heating to 290°; as the oxide is described as orange-red, it was probably the form of P₄O which reduces salts of Au, Ag, and Hg.

Preparation and Properties of P₄O which does not reduce salts of Ag, Au, and Hg.—POCl, is heated with an excess of P in a sealed tube to 200°-250°; the sides of the tube become covered with a scarlet-coloured solid, which is washed with CS₂ (to remove P) and dried in vacuo (R. a. G., l.c.; the other products are PCl₃, P₂O₃Cl₄ and a little P₂O₃). As thus prepared, P₄O is a scarlet-red solid; S.G. 138; it is not acted on by water or alkali solutions, and it does not reduce salts of Ag, Au, or Hg. This form of P₄O seems to have been obtained by Pelouze (A. 3, 52) by burning P in O; and by Le Verrier (A. 27, 175) by burning P in a thin layer on a porcelain plate, washing with water to remove oxyacids, and boiling with PCl₂ to remove P.

PHOSPHOROUS OXIDE P₄O₆. (Phosphorus trioxide. Phosphorous anhydride.) Mol. w. 219·6; same in solution in C₆H₆ (Thorpe a. Tutton, C. J. 57, 545). Melts at 22·5°; boils at 173·1° (T. a. T.). S.G. $\frac{31^{\circ}}{16}$ liquid P₄O₆ 1·9481; solid P₄O₆ at $\frac{21^{\circ}}{6^{\circ}}$ c. 2·13·5. S.G. at b.p. = 1·6897 (T. a. T.). V.D. 111·9. S.V. 130·2 (T. a. T.). $\mu_{\rm A} = 1.5311$; Mol. w. $\times \mu_{\rm B} = 60.5$ (T. a. T.). S.G.

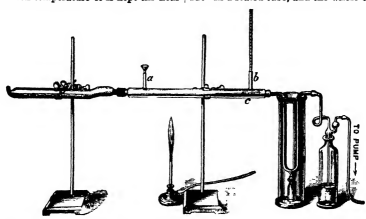
Mol. w.
$$\frac{\mu_H - \mu_A}{S.G.} = 4.17$$
. M.M. 9.962.

Preparation.—A piece of combustion-tubing about 40 mm. bore is drawn into the shape shown in the figure on following page, and is fitted by a

good cork into a brass tube, which is surrounded | by a wider tube of brass; water is introduced at a between the brass tubes, and a thermometer is placed at b. A loose-plug of glass-wool is placed in the inner brass tube at c. The U-shaped condenser is of rather narrow bore, the limbs are about 300 mm. long, a small bottle is attached to the vertical piece at the bottom of the condenser, which is itself connected with the inner brass tube by a good cork; the condenser and small bottle are surrounded by ice-and salt. The bottle between the condenser and the pump contains oil of vitriol. Two sticks of P are dried between filter-paper, cut into pieces c. 25 mm. long, and placed in the combustion tube, which is then narrowed at the open end and fitted into the brass tube; the pump is set in action, and the P is ignited by holding a small flame beneath the open end of the combustion-tube. Air is sucked over the P as rapidly as possible; if the combustion is too local, much P2O5 is formed; considerable quantities of red P4O are formed near the burning P. When the P has burnt for c. 15 mins. the water in the brass tube is heated to c. 50°, at which temperature it is kept till near

132°. Mol. w. as gas, and also in solution in C.H. (determined by cryoscopic method), corresponds with P.O. Soluble without change in C₆H₆, CS₂, CHCl₃, and ether. Heated to c. 200° in sealed tube begins to decompose to P2O4 and P. Decomposed by light, with separation of red P. T. a. T. (C. J. 59, 1019) several times obtained clear, transparent, perfectly-formed crystals of P₄O₈ by slow spontaneous sublimation in vacuo, which remained unchanged for many days, although exposed to light. Spec. volume at b.p. = 130.2. (For details of thermal expansion and S.G. at b.p. v. T. a. T., C. J. 57, 545.) Reacts slowly with cold water, forming H₃PO₃Aq; with boiling water reaction is violent, PH, is evolved, and red P, or perhaps P,O, and H,PO,Aq are formed. Oxidises to P2O5 by exposure to air or O; if temperature is raised the P,O, ignites and may explode. P.O. has a well-marked physiological action; the action of P may be due to P, Oe, formed by the air oxidising the P (T.a. T., l.c. p. 573).

Reactions.—1. Heat decomposes P_4O_4 , forming P_2O_4 and P (? $P_4O)$; the action begins at c. 210° in a sealed tube, and the whole of the P_4O_4



the end of the experiment, when it is raised to 60°. P_4O_6 begins to collect in the condenser about 30 mins. after starting the combustion; if the glass-wool is packed properly (it should not be too tight) no P_2O_5 passes into the condenser, and if the temperature of the water does not exceed 60° only very small quantities of P pass over. The process is stopped when c. 5ths of the P is burnt, else the P_4O_6 may be oxidised. The condenser is removed, the P_4O_6 is melted by the heat of the hand, and run down into the little bottle, from which it may be transferred to another bottle which has been filled with dry CO_2 ; this bottle should have a very tightly-fitting stopper, and should be kept in the dark in an atmosphere of CO_2 (T. a. T., l.c.).

Properties.—A white snow-like solid, with an alliaceous odour resembling that of P; when melted and cooled slowly crystallises in long thin prisms, probably monoclinic; melts at 22.5° to a clear, colourless, very mobile liquid, which solidifies again at 21°; the liquid may be cooled in a narrow tube several degrees below m.p. without solidifying; boils at 173.1° in CO₂ or N. Can be gasified in Hofmann's V.D. apparatus at

is decomposed at 440° (T. a. T., p. 552). -2. Rapidly acted on by light, becoming yellow and then dark-red (T. a. T., p. 553). The action of light separates red P; the amount obtained after several months' exposure does not exceed 1 p.c. (T. a. T., C. J. 59, 1019). By exposing P.O. to light for some months in a sealed tube filled with dry CO2, then melting, filtering through glass-wool, and repeating this process several times, T. a. T. eventually obtained P4O6 which remained perfectly clear and colourless after twelve months' exposure (l.c., p. 1023).—3. Oxidises to P2O5 in air or oxygen; when quite free from P, the oxide is not spontaneously inflammable. Under reduced pressure in O the P.O. glows; on raising temperature to c. 70° at ordinary pressure the glow gives place to flame. Combination occurs between vapour of P.O. and O; ozone is not formed. When P,O, is thrown into O heated to 50°-60° ignition occurs with an intensely brilliant flame (T. a. T., C. J. 57, 569).-4. Glows continuously, and oxidises to P₂O₅, when a stream of ozonised oxygen is passed over it.—5. Ignites in chlorine, burning with a greenish flame; when surrounded by ice and

exposed to slow stream of Cl a liquid is formed, which on distillation gives POCl₂, while PO₂Cl (?) remains (T. a. T., C. J. 57, 572).-6. Reacts violently with liquid bromine; using Br vapour at ordinary temperature, T. a. T. (C. J. 59, 1020) obtained PBr, and P2O5, and on then heating POBr, and PO2Br (?) were formed.-7. Iodine reacts slowly; by heating with I and CS, under pressure, P_2O_5 and P_2I_4 are produced (T. a. T., *l.c.*, p. 1021). 8. Heated with sulphur, in CO₂ or N, two layers of liquid are formed; at 160° there is violent reaction, and solid P₄O₆S₄ is produced (v. Phosphorus, sulphoxide of, p. 149; T. a. T., l.c., p. 1022).-9. Seems to form a Se compound analogous to P₄O₆S₄ by heating with selenion (T. a. T., i.c., p. 1026).—10. Dissolves very slowly in water, forming H_3PO_3Aq (T. a. T., C. J. 57, 567); $P_4O_6+6H_2O+Aq=4H_3PO_3Aq$. The action of hot water is very energetic; red P, or red P,O, is ppd., inflammable hydride of P is evolved, and H₂PO₄Aq is formed; if the quantity of P₄O₆ exceeds 2 g. the action of hot water is violently explosive.—11. Dry hydrogen chloride is rapidly absorbed; PCl₃ is formed along with H₂PO₃, H₃PO₄, and yellow P (T. a. T., C. J. 59, 1022).-12. Conc. sulphuric acid reacts violently, forming H₃PO₄ and SO₂; if 1 g. or more P₄O₆ is used, the mass ignites (T. a. T., l.c. p. 1026).—13. P.O. is rapidly oxidised to P.O. by sulphur trioxide, which is reduced to SO₂; no compound could be obtained (T. a. T., l.c. p. 1026). According to Adie (C. J. 59, 230), if a little water is present, P4O6 and SO8 form an unstable compound H₃PO₄.3SO₃.—14. Nitrogen tetroxide seems to react like SO₃, forming P₂O₅ and N2O2 or NO (T. a. T., l.c. p. 1028).—15. Phosphorus pentachloride produces PCl3 and POCl3; phosphorus trichloride reacts at c. 180° in a sealed tube, forming a mixture of P₂O₃, PCl₃, and red P (T. a. T., l.c. p. 1028).—16. Sulphur chloride reacts violently, producing POCl, PSCl, SO, and S (T. a. T., l.c. p. 1026).—17. Ammonia reacts with some violence, probably producing OH.P. $(NH_2)_2$ (v. Phosphamides, p. 105).—18. Cold dilute caustic soda, or caustic potash solution, slowly dissolves P₄O₆, forming a solution of Na or K phosphite; cold conc. or hot dilute solution of soda or potash produces red P (or P.O?) and alkali phosphate, and evolves inflammable hydride of P.—19. Ignites in contact with absolute alcohol; by allowing the alcohol to drop slowly on to cooled P₄O₆ the acid P(OEt)₂OH is formed (T. a. T., C. J. 57, 569). P₄O₆ seems not to react with H, PH₃, CO, CO₂, SO₂, N, NO, CN, or C₂H₄ (T. a. T., C. J. 59, 1029).

PHOSPHORUS TETROXIDE P2O4 (Phosphorosophosphoric oxide. Hypophosphoric oxide). Mol. w. not determined; corresponding sulphide has mol. w. P₂S₆. Hautefeuille a. Perrey (C. R. 99, 33) noticed that a crystalline sublimate is obtained by heating the products of the combustion of P; Thorpe a. Tutton (C. J. 49, 833) proved that this sublimate is a definite oxide P₂O₄. Concerning the combustion of P in air v.

pp. 128 and 139.

Preparation .- P is burnt slowly in a stream of air dried by H₂SO₄ and P₂O₅; the products are collected in a glass tube, surrounded by an outer tube filled with steam, and are then quickly transferred to a tube filled with dry CO₂, the tube is drawn out, exhausted by a Sprengel pump

(care being taken to prevent entrance of moisture), and sealed; at c. 290° the white mass becomes orange-coloured, and then red, and a white crystalline sublimate of P2O4 is formed, leaving P_2O_5 and P_4O in the lower part of the tube. T. a. T. (p. 838) think that the P_2O_4 is derived from the P_2O_3 , thus $7P_2O_3 = 5P_2O_4 + P_4O$; that P₂O₄ is formed by heating P₂O₅ out of contact with O is shown by T. a. T. (C. J. 57, 552).

Properties and Reactions.—Colourless crystals, probably orthorhombic; do not melt at 100°. Volatilises at c. 180°. Very deliquescent, forming H₃PO₃Aq and H₃PO₄Aq; from this reaction T. a. T. conclude that P₂O₄ is not the anhydride of $H_4P_2O_8$, as an aqueous solution of this acid does not behave like a mixture of H₃PO₈ and H₃PO₄, and can be boiled without change. Solution of P2O4 in water reduces AgNO, Aq to Ag, and HgCl, Aq to HgCl; KMnO, Aq is very slowly decolourised; 'magnesia mixture' gives an immediate pp., and after standing some time the filtrate gives a large pp., with NH, molybdate after heating with HNO,Aq.

PHOSPHORIC OXIDE P2O5 (Phosphoric an-Phosphorus pentoxide). hudride. Formula probably molecular, from analogy of P2S5. S.G. 2:387 (Brisson, Pesanteur specifique des Corps [Paris, 1787]). H.F. [P²,0³] = 369,900; [P²O¸,Aq] = 35,600 (Th. 2, 409). Formation.—1. This oxide is formed by burn-

ing P in a large excess of air; concerning the combustion of P in air v. pp. 128 and 139.— 2. It is also formed by burning P in such gaseous O compounds as NO, NO₂, ClO₂.—8. By distilling

P₂O₃Cl₄.

Preparation.—A large glass balloon with a wide opening and two side necks is thoroughly dried; through a cork in the wide opening passes a piece of wide tubing, which reaches to about the centre of the balloon, and from the lower end of this tube a small porcelain basin is suspended by Pt wires; one of the side necks is connected with U-tubes containing pumice soaked in H₂SO₄, and the other side neck is connected with a wide-mouthed perfectly dry bottle, which is again in connection with a water-pump, a bottle with H2SO4 being placed between the pump and the wide-mouthed bottle. A small piece of well-dried P is placed in the little basin, and is ignited by passing a hot wire down the glass tube, the mouth of which is then closed by a good cork; a rapid stream of air is sucked through the apparatus; the P is burnt to P2O3, part of which collects on the bottom of the balloon and part passes into the bottle attached to the side neck. When the P is burnt another small piece is dropped down the glass tube into the little basin, and the process is continued. When sufficient P2O5 has collected in the bottle this is closed by a well-fitting stopper (Delalande, A. Ch. [2] 76, 117). For other forms of apparatus v. Marchand, J. pr. 16, 373; Mohr, Gm.-K. i. 2, 120. Grabowski (A. 136, 119) has described a vessel of tinplate for preparing large quantities of P2O5.

Shenstone says that P2O5, prepared by the ordinary process, has generally reducing powers, and is not, therefore, trustworthy as a drying agent. S. recommends to heat for some days at c. 300° in a very slow stream of dry O; then to

heat at 300° in a more liberal, but still limited, supply of dry O until the reddish colour produced by the first heating has disappeared; then to heat at 300° in a more rapid stream of dry O so long as sublimation occurs; and finally to sublime, fractionally, in a current of dry O, from retorts of hard glass into receivers of the same material. The first portions of the sublimate should be rejected. The whole of these processes may be done in glass vessels. The retorts used in the final sublimation may be heated in a combustion furnace (private communication).

Hautefeuille a. Perrey (C. R. 99, 33) say that P2O5 can be obtained in three forms: crystalline, amorphous and powdery, and glass-like. They say that when P is burnt in a glass tube in dry air, the crystalline form of P₂O₅ is deposited on the colder part of the tube, the amorphous powdery form on the hotter part, and the vitreous form on that part of the tube which is heated to redness. These authors assert that crystalline P₂O₃ is obtained free from the other forms by distilling the products of the burning of P; that heating the crystals in vapour of S produces the powdery amorphous form; that the vitreous variety is formed by heating either of the others to low redness; and that this vitreous form yields crystalline P₂O₅ when sublimed at a red heat. Considering the facts brought to light by the work of Thorpe a. Tutton on the combustion of P (C. J. 49, 833; 57, 545; v. beginning of this article, p. 139), the existence of various modifications of P2O3 must be looked on as very doubtful. T. a. T. (C. J 49, 838) think it very likely that the crystalline form of P2Os obtained by H. a. P. was P₂O₄; and it is probable that the differences in the properties of the powdery and vitreous P2O5 were due to admixture with P2O3. No analytical data are given by H. a. P. Shenstone (priv. comm.) says that P2Os purified by his method is crystalline, but becomes amorphous when suddenly heated.

Properties .- A snow-white, amorphous, inodorous, very deliquescent, solid. Melts and sublimes below red heat (Lautermann, A. 113, 240); according to Davy (A. Ch. [2] 10, 218) volatile only at white heat. Absorbs water very rapidly; hence is used as a very efficacious drying agent. P_2O_5 is the anhydride of three phosphoric acids: HPO2, H2PO4, and H4P2O7. Dry P2O3 does not change the colour of dry litmus paper. Commercial P₂O₃ often contains traces of P, which cause it to redden in light, and also traces of As2O, derived from the impure P used; it generally has more or less marked reducing powers.

Reactions. — 1. Water produces HPO; ; H,P,2O, is formed from HPO, by the longcontinued action of moist air; and H.PO. is produced by continued heating of HPO2Aq.-2. P₂O₅ removes H₂O from many compounds which contain water or the elements of water; e.g. it produces anhydrides when heated with several acids, H₂SO₄ gives SO₃, HNO₃ gives N₂O₅, and hydrocarbons are often formed by heating P₂O₅ with compounds of C, H, and O.—3. Heated with several metals, e.g. Fe, K, Na, Zn, metallic oxide, phosphide, and phosphate are formed .-4. Salts of acids the anhydrides of which are volatile are decomposed by heating with P₂O₅, with formation of phosphates and volatilisation

of the anhydrides (Odling, P. M. [4] 18, 168).-5. POCl₂ is formed by heating P₂O₅ with sodium chloride, or with phosphorus pentachloride. - 6. Heated with phosphoryl chloride, P2O3Cl, and P,O15Cl5 (which may be a mixture) are formed (Huntly, C. J. 59, 202).—7. Ammonia produces PONH₂(OH)₂ and P₂O₃(NH₂)₂(OH)₂ (v. Pноs-PHAMIC ACIDS, p. 105).—8. Heated with boron chloride to 200° for 2-8 days, P₂O₅·B₂O₅ and POCl₃.BCl₃ are formed (Gustavson, B. 4, 976). P2O5.B2O2 is also formed by heating P2O5 with boric acid.

Combinations.—1. With water (v. supra, Reactions, No. 1).—2. With sulphur trioxide to form P₂O₃.3SO₃, decomposed at 30° (Weber, B. 20, 86).—3. With silica and water to form P₂O₃.SiO₂.4H₂O; formed by heating conc. H₃PO₄Aq with side (Hautefeuille a. Margottet, C. R. 104, 156). Similar compounds with oxides of titanium, zirconium, and tin are described by H. a. M. (C. R. 102, 1017).

Phosphorus, oxyacids of. The three phosphoric acids are described in the article Phos-PHORIC ACIDS, p. 124, and the salts of those acids in the article Phosphates, p. 106; the other oxyacids of P and their salts are described under PHOSPHORUS, OXYACIDS OF, AND THEIR SALTS, p.

Phosphorus, oxybromide of, POBr_s (Phosphoryl tribromide). Mol. w. 286·2. Melts at 55° (Baudrimont, A. Ch. [4] 2, 58); at 45°-46° (Ritter, A. 95, 210). Boils at 195° (Ritter, l.c.); at 193° (Gladstone, P. M. [3] 35, 345). V.D. 145·5. S.G. 2·822 (Ritter, J. 8, 301). H.F. $[P,O,Br^3] = 108,000$ from liquid Br; 120,000 from gaseous Br (Ogier, C. R. 92, 83).

Formation.—1. By the action of moist air or a very little H₂O on PBr₆ (Gladstone, P. M. [3] 35, 345).-2. By the action of oxalic or acetic acid on PBr_s (Baudrimont, A. Ch. [4] 2, 58; Ritter, A. 95, 210).—3. By passing O into boiling PBr₃; explosions generally occur (Demole, Bl. [2] 84, 201).

Preparation.—To 137.5 parts PCl, are added 18 parts H₂O, and then 160 parts Br are added drop by drop, HBr and HCl are evolved, and POCl, and POBr, remain (part of the PCl, is decomposed to H,PO, and HCl, and the H,PO₃, Br, and remaining PCl₃ react, (?) thus 2PCl₃+H₂PO₃+3Br₂=2POCl₃+POBr₃+3HBr); the liquid is distilled, the portion boiling above 150° being kept separate and surrounded by a freezing mixture till it solidifies (Geuther a. Michaelis, J. Z. 6, 242).

Properties and Reactions.—Large colourless plates, which melt and boil in dry air without decomposition; soluble in conc. H₂SO₄, reppd. by H₂O; soluble in CHCl₃, CS₂, ether, and turpentine. Decomposed rapidly by water to HBrAq and H, PO, Aq; bromine chloride forms POBrCl, or POCl₃, and Br, according to the quantity used (Geuther, J. Z. 10, 130); hydrogen sulphide probably forms PSBr₃; chlorine forms POCl₃ and Br; bromine seems to form an additive compound which separates into POBr, and Br on heating (Gladstone, l.c.); tin and antimony remove Br (Baudrimont, l.c.).

Phosphorus, oxybromochlorides of. Two of these compounds are known, but it is doubtful whether the second is a definite compound or a

mixture of POCl, and POBr,.

PHOSPHORUS OXYBROMODICHLORIDE POBrCl2. (Phosphoryl bromodichloride). Mol. w. 197 4. Melts at 11°; boils at 137.6 (Thorpe, C. J. 37, 343). S.G. & 2.12065 (Thorpe, l.c.). A colourless liquid, which solidifies, when cooled below 0° to large colourless tablets which melt at 11°; when repeatedly distilled, or more quickly when heated in a closed tube to c. 185°, decomposed to POCl₃ and POBr₅ (Chambon, J. Z. 10, 92); decomposed by H₂O to H₃PO₄Aq, HClAq, and HBrAq; with a little alcohol forms POCl₂(OEt) and HBr. Prepared by the reaction of POBr, with BrCl in the ratio POBr₃: BrCl (Geuther, J. Z. 10, 130); by the action of PBr, on $P_2O_3Cl_4$ (G. a. Michaelis, J. Z. 7, 103). By reacting on P.OEt.Cl, with Br (Menschutkin, A. 139, 343); 80 g. Br are allowed to drop slowly into 74 g. P.OEt.Cl₂ kept well cooled and constantly shaken; the reaction is very violent; the POBrCl, is separated from the C.H.Br formed by fractionation (Thorpe, C. J. 37, 343). The P.OEt.Cl₂ is prepared by slowly dropping absolute alcohol into well-cooled PCl₃ in the ratio C₂H₆O: PCl₃, allowing to stand for some time, and fractionating; the boiling-point is 117°-118°.

PHOSPHORUS OXYDIBROMOCHLORIDE POBr₂Cl (Phosphoryl dibromochloride). By the reaction of POBr₃ and BrCl, in the ratio POBr₃: 2BrCl, Geuther (J. Z. 10, 130) obtained a liquid boiling between 150° and 160°, and decomposing very easily by heat into POCl₃ and POBr₃; analyses agreed with the formula POBr₂Cl, but the liquid was possibly a mixture of POCl₄ and POBr₃

 $(3POBr_2Cl = POCl_3 + 2POBr_3).$

Phosphorus, oxychlorides of. Two oxychlorides of P have been isolated, POCl₂ and P₂O₃Cl₄. A third substance obtained by Gustavson (B. 4, 853) by heating together P₃O₃ and POCl₃, and said by him to be PO₂Cl, is declared by Michaelis (G. O. [1881] 2, 396) not to be a definite compound, and this is confirmed by Huntly (C. J. 59, 202).

PHOSPHORYL CHLORIDE POCI₂. Mol. w. 153·3. Boils at 107·2° (Thorpe, C. J. 37, 337). Melts at -1·5° (Geuther a. Michaelis, B. 4, 769). S.G. ⁹⁰ 1·71163 (Thorpe, L.c.). V.D. at 150° = 77·2 (Cahours, A. Ch. [3] 20, 369). For expansion by heat v. Thorpe (L.c.). S.V. 101·4 (T., L.c.). H.F. [P,O,Cl³] = 145,960; [PCl³,0] = 70,660 (Th.

2, 325).

Formation.—1. By heating PCl₃ in air or O, or with ozone, SO₃, SO₂Cl₂, and various other oxidisers.—2. By strongly heating PCl₃ in O, or with SO₃, SO₂, B₂O₃, &c., or by decomposing by a little H₂O.—3. By distilling NaCl mixed with P₂O₅ (Kolbe a. Lautemann, A. 113, 240).—4. By passing CO and Cl over Ca₃2PO₄ mixed with C and heated to redness (Riban, C. R. 95, 110; Oglialoro, G. 13, 328).—5. By heating PCl₅ with P₂O₅.—6. By the reaction of Cl with P₂O₃ (Thorpe a. Tutton, C. J. 57, 572).

Preparation.—An intimate mixture of 1 part P_2O_8 with 2.9 parts PCl_8 is distilled, and the product is again distilled, that boiling at $106^{\circ}-108^{\circ}$ being collected separately; the equation $P_2O_8+3PCl_8=5POCl_3$ is realised.

Properties.—A colourless, highly refractive, strongly smelling liquid, which fumes in the air; the fumes strongly affect the eyes and respiratory organs; solidifies at -10°, when touched

with a glass rod, to long, colourless tablets, which melt again at -1.5° (Geuther a. Michaelis, B. 4, 769). POCl₃ does not conduct electricity even when heated (Buff, A. 110, 257).

Reactions.—1. Water reacts rapidly, forming HClAq and H₃PO₃Aq [POCl³, Aq] = 72,190 (Th. 2, 325).—2. Alcohol produces HCl and PO.OEt.Cl2 3. Sulphur trioxide at 160° forms P₂O₃Cl₄ and P₂O₅ (Michaelis, G. O. 2, 393).—4. Sulphuric acid produces SO₂.OH.Cl, HCl, and HPO₃ (Michaelis, l.c.).—5. Most organic acids react similarly to H₂SO₄, producing acid chlorides; e.g. C₂H₃O.OH forms C₂H₃O.Cl; salts of organic acids react similarly, forming acid chloride and metaphosphate.—6. Sulphates form SO,Cl, and orthophosphates.—7. Sulphites are partially changed to SO₂, mixed with metallic chloride and phosphate (Divers a. Shimidzu, C. J. 47, 208).— 8. Nitrates produce chlorides and P₂O₅ &c.; nitrites give chlorides and phosphates and NO. (D. a. S., l.c.).—9. POCl_s separates I from potassium iodide (Schiff, A. 102, 181).-10. Chromyl chloride forms Cl, P₂O₅, CrCl₃, and Cr₂O₅ (Casselmann, A. 91, 241; 98, 213).—11. Phosphorus pentasulphide at 150° produces PSCl₃ and P₂O₃. 12. Phosphorus pentoxide reacts at c. 200°, forming P2O3Cl4 and a substance P,O1,Cl5, which may prove to be a mixture (Huntly, C. J. 59, 202).—
13. Phosphorus reacts at 200° to form P.O. PCl., P₂O₃Cl₄, and a little P₂O₅ (Reinitzer a. Gold schmidt, B. 13, 845).—14. POCl₃ is reduced by heating in closed tubes with many finely divided metals, e.g. Ag, Hg, Fe, Cu, As (Pb is without action); the products are generally PCl, and P₂O₅, and oxide, chloride, or phosphate of the metal; sometimes P₂O₃Cl₄ is said to be formed (v. R. a. G., l.c.); Mg, Al, and Zn are said to form red P₄O (R. a. G., l.c.). Zinc-dust with a little POCl, forms Zn phosphide, which evolves inflammable P hydride on addition of a little water (Deniges, Bl. [3] 2, 787). — 15. With ammonia forms various phosphamic acids (q. v., p. 105), and also phosphamides (q. v., p. 105).— 16. Boric oxide heated with POCl, in a sealed tube to 150°-170° for 8 to 10 hours forms

P₂O₃.B₂O₃ and POCl₃.BCl₃ (Gustavson, B. 4, 975).

Combinations.—1. With ammonia to form a compound not yet examined.—2. With several metallic chlorides to form double compounds, e.g. with AlCl₃, MgCl₂, and SnCl₂ (Casselmann, A. 91, 241; 98, 213). Some of these compounds may be vapourised in absence of moisture; they are decomposed by water.—3. With boric chloride to form POCl₃.BCl₃, by passing BCl₃ into POCl₄; also by heating POCl₃ with B₂O₃. This substance is decomposed by heat; with water it gives H₃PO₄Aq, H₃BO₂Aq, and HClAq (Gustavson,

B. 4, 975).

Constitution of POCl₂.—Two isomeric forms of POCl₃ are possible: Cl₂POCl and Cl₃PO; but the compounds obtained by different methods are identical. Wichelhaus (A. Suppl. 6, 257) thought that POCl₂ must have the constitution Cl₂POCl, because of its formation from Cl₂POEt, in which the O atom is in direct union with the P atom (Cl₂POEt+Cl₂=Cl₂POCl+EtCl); but Geuther a. Michaelis (J. Z. 6, 242) pointed out that this formation of POCl₂ may be interpreted also in favour of the constitution Cl₃PO by supposing that Cl first replaces OEt, forming PCl₂ and that the PCl₃ and EtOCl then react:

(1) $Cl_2POEt + Cl_2 = Cl_3P + EtOCl_4$ (2) $Cl_{\mathfrak{g}}P + EtOCl_{\mathfrak{g}} = Cl_{\mathfrak{g}}PO + EtCl_{\mathfrak{g}}$ \mathbf{T} he experiments of Reinitzer a. Goldschmidt (B. 13, 845) on the action of metals on POCl, show that in most cases O is withdrawn, leaving PCl3; and that in a few cases the 3 Cl atoms are withdrawn with formation of P₂O₅. These results do not indicate any differences between the functions of the Cl atoms, and hence they tend to favour the formula Cl₃PO. Thorpe (C. J. 37, 387) arrived at the formula Cl2POCl from consideration of the specific volumes of POCl, and PCl, Masson a. Ramsay (C. J. 39, 50), also from considering the specific volumes of P and P compounds, deduced the formula OPCl. Although but one POCl₃ has been obtained, the isomeric compounds (C₆H₅)₂POC₆H₅ and (C₆H₅)₅PO have been isolated (La Coste, B. 18, 2118).

PYROPHOSPHORYL CHLORIDE P.O.Cl. phorus trioxytetrachloride). Mol. w. not known. Boils 210°-215°, with partial decomposition.

S.G. 1.58 at 7°.

Preparation.-1. 100 g. PCl, are placed in a cylinder surrounded by ice and salt, and the vapour from $20\,\mathrm{g}$. liquid NO_2 is passed into the cylinder by a tube which reaches just to the surface of the PCl₃. The action proceeds at once; the products are P₂O₃, POCl₃, P₂O₃Cl₄, and NOCl. When all the NO₂ has been distilled into the PCl₃, the cylinder is placed in warm water to remove NOCl, and its contents are then fractionated; PCl, distils over first, then a considerable quantity of POCl, between 105° and 110°, and then P₂O₃Cl, from 200° upwards. It is best to prepare a considerable quantity of the impure product before fractionating; 350g. PCl₂ yielded 40 g. P₂O₂Cl₄ (Geuther a. Michaelis, B. 4, 766).—2. The vapour obtained by heating 30 g. starch with 180 g. HNO, Aq is led into 100 g. PCls; the product is fractionated, and that boiling under 100° is again treated as before; finally, the whole is fractionated. About 20 g. P2O3Cl4 are obtained from 200 g. PCl2 (G. a. M., l.c.).

Properties.—A colourless liquid, which fumes in the air; the vapour attacks cork. Has not been solidified by cooling. Boils, with partial decomposition into P2Os and POCls, at 210°-

214°.

Reactions.—1. Heat produces partial decomposition into POCl₃ and P₂O₅.—2. Decomposed by water to H₂PO₄Aq and HClAq.—3. Alcohol produces PO.OEt.Cl₂, PO.OEt.(OH)₂, and HCl. 4. Phosphorus pentachloride produces POCls. 5. Phosphorus pentabromide forms POBr. and POCl₂Br (G. a. M., l.c.).

METAPHOSPHORYL CHLORIDE PO2Cl. Gustavson (B. 4, 853) said that a compound of this composition is formed by heating P_2O_5 with POCl,; but Michaelis (G. O. 2, 396) says that the substance is not a true compound, and this is confirmed by Huntly (C. J. 59, 242).

Phosphorus, oxychlorobromides of, v. Phos-

PHORUS, OXYBROMOCHLORIDES OF, p. 142.

Phosphorus, oxyfluoride of, POF₃ (Phosphorus, oxyfluoride of, POF₃ (Phosphorus, Oxyfluoride) w 103.92. V.D. 52 phoryl fluoride). (Thorpe, C. J. 55, 759; Moissan, Bl. [3] 4, 260). A gas which fumes in air and is rapidly absorbed by water. Formed by the reaction of P_2O_3 with fluorides (Schulze, J. pr. [2] 21, 438); also by exploding a mixture of 2 vols. PF₃ and 1 vol. O

by passing electric sparks (Moissan, C. R. 102, 1245); by gently heating an intimate mixture of 2 pts. finely-powdered cryolite with 3 pts. P.O. in a brass tube, and collecting over Hg as soon as the gas which comes off is wholly absorbed by NaOHAq (Thorpe, l.c.). Moissan (l.c.) prepares POF, by heating ZnF, with POCl, in a brass tube, fitted with a leaden delivery tube; he allows the gas to pass through a brass tube cooled to -20°, and then over ZnF₂ to remove traces of POCI.

PÖF₃ liquefies at 16° under a pressure of 15 atmos., or at the ordinary pressure by cooling to -50° (M., C. R. 102, 1245). When compressed under 50 atmos. and the pressure then suddenly released, it solidifies to a snow-like mass. When quite dry it is without action on glass or Hg; heated in a glass tube SiF, and an alkaline phos-

phate are formed.

Phosphorus, oxyiodide of, PsI6O8. (Am. 3, 280) obtained a red crystalline compound, to which he gave this composition, as a by-product in preparing C_2H_3I . M.P. c. 140° ; sublimable, with partial decomposition; e. sol. water, alcohol, and ether.

Phosphorus, oxynitride of, PON (Phosphoryl nitride. Phosphoryl nitrile. Formerly called Phosphomonamide). Mol. w. uncertain. A white powder, which melts at red heat to black glass-like mass. Obtained by heating PO(NH₂), or PONHNH2, the former of which is formed by the reaction of POCl₃ with NH₃, and the latter by the reaction of PCl₅ and NH₂ (Schiff, A. 101, 300; cf. Phosphamides, p. 105). Gives K₃PO₄ and NH₃ by action of molten KOH (v. also Gladstone, C. J. [2] 7, 18).

Phosphorus, selenides of. P and Se combine directly to form four compounds: P.Se, P₂Se, P₂Se₃, and P₂Se₅. P₂Se, P₂Se₃, and P₂Se₅ combine with the selenides of the more positive metals to form compounds R2Se.P2Se, (R2Se)2.P2Se3, and (R2Se)2.P2Se5. The formulæ

are not necessarily molecular.

PHOSPHORUS SUBSELENIDE PaSe (Phosphorus hemiselenide). Prepared by melting together Se and ordinary P in the ratio Se: 4P(1:1.57) in an atmosphere free from O, and separating from amorphous P by distillation in absence of O. A dark yellow, oily, fetid-smelling liquid, which solidifies at -12°; e. sol. in CS, insol. alcohol and ether; takes fire in air; decomposed by water containing air, with formation of H, PO, Aq and Se compounds; not acted on by cold alkali solu-tions, but with boiling alkali solutions gives alkali phosphate and selenite, metallic selenide and P hydride; in solutions of metallic salts becomes covered with a crust of metallic phosphide

and selenide (Hahn, J. pr. 93, 430). DIPHOSPHORUS SELENIDE P₂Se. DIPHOSPHORUS SELENIDE P_2 Se. (Phosphorus protoselenide or monoselenide). Prepared by melting together, in absence of O, P, and Se in ratio 2P: Se(1:1.28). A red solid; unchanged in dry air, in moist air gives off H₂Se; insoluble alcohol and ether; Pis withdrawn by CS₂; boiling alkali solutions produce H.Se and a red substance containing Se and P. By heating, in absence of O, with equivalent weights of metallic selenides, compounds R12Se.P2Se and R11Se.P2Se are formed; R=Ba, Cu, Fe, Mn, K, Ag, Na. The alkali compounds are decomposed by water; the compounds containing heavy metals deconpose only at high temperatures, but they generally take fire when rubbed in a mortar; they are slowly decomposed by hot alkali solutions (Hahn, l.c.). From their empirical composition, R1PSe, these compounds might be called seleno-meta-

hypophosphites.

Phosphorous selenide P.Se. (Phosphorus sesquiselenide or trisclenide). Prepared by heating together P and Se in ratio 2P:3Se(1:3.82) in absence of O. A dark ruby-red solid; sublimes to yellow vapours when heated out of O; burns when ignited in air. Oxidised slowly in moist air; evolves H2Se when boiled with water. Insoluble alcohol, ether, and CS2; easily soluble in potash, less soluble in alkali carbonate solutions. Forms compounds with metallic selenides (R¹₂Se)₂.P₂Se₃, and 2R¹¹Se.P₂Se₃ (Hahn, l.c.); R = Ba, Cu, Pb, Mn, K, Ag, Na. From their empirical composition, R¹₄P₂Se₃, these compounds might be called selenopyrophosphites.

PHOSPHORIC SELENIDE P.Se. (Phosphorus pentaselenide). Prepared by heating together amorphous P and Se in ratio 2P:5Se (1:6:37) in a stream of CO2. A dark-red, nearly black, glasslike solid; insoluble in CS₂; crystallises from CCl, in black needles. In moist air, or water, decomposes slowly to H2Se and H2POAq; in conc. KOHAq quickly forms K₂Se and K₃PO₄Aq; alcohol forms H₂Se and Et₃PO₂Se₂ and also Et₂HPO₂Se₂. Very unstable compounds probably of the forms (R^{I}_{2} Se)₂, P_{2} Se₃ and $2R^{II}$ Se. P_{2} Se₃ are obtained by heating P_{2} Se₃ with metallic selenides; R = Ba, Cu, Pb, Mn, K, Ag, Na. The empirical composition of these compounds would lead to the name selenopyrophosphates (Bogen, A. 124, 57; Rathke, A. 152, 200). Rathke (l.c.) supposed that the K salt of a selenophosphoric acid is formed by the reaction of P2Se, with KOHAq.

Phosphorus, selenion acids of, salts of; v. DIPHOSPHORUS SELENIDE, PHOSPHOROUS SELENIDE,

and Phosphoric selenide (supra).

Phosphorus, sulphides of. P and S combine, when heated together, with production of much heat and light; the action is often violently explosive. Four definite compounds seem to exist; the simplest formulæ are P4S2, P2S3, PS2, and

P₂S₃.

Two compounds containing less S than P₄S₃, viz. P2S and P4S, are often described, and substances said to be compounds of P and S in the ratio P:3S and P:6S have been mentioned by

Dupré and Berzelius.

The substance to which the formula P₄S was given was obtained by heating P and S in the proper proportions (Berzelius, A. 46, 129, 255), or by digesting P with an alcoholic solution of K.S. (Böttger, J. pr. 12, 857), or according to Wicke (A. 86, 115) by contact of P and S at ordinary temperatures; it was described as a colourless, oily liquid. A solid red form of P.S was said to be formed by heating liquid P₂S with Na₂CO₃ (Berzelius, l.c.). The formula P2S was given to a substance of which two forms were described: a yellow liquid obtained by carefully heating together P and S in the proper ratio, and a dark-red powder obtained by heating the liquid form with dry MnS in H (Berzelius, l.c.). Various double compounds of P₂S with metallic sulphides, R¹₂S.P₂S and RIS.P.S (R = Cu, Fe, Mn, Hg, Ag) were described by Berzelius. The composition of these bodies Vol. IV.

would lead to their being called thiometahypophosphites. Isambert (C. R. 96, 1771) has shown that when a solution of P in a large excess of S is distilled in vacuo the whole of the P passes off and leaves S. Schulze (J. pr. [2] 22, 113) found that P separated at 8° from a solution of S in P in the ratio P.S, and that S separated when the elements were present in the ratio P2S. Schulze also found that a stream of CO2 removed P from both solutions at 150°; that heating to this temperature generally produced explosion; that the so-called liquid P₂S was soluble in CS, and that most of the S was removed by shaking this solution with alcohol, ether, or CHCl_s. Further, Schulze (B. 16, 2066) observed that solutions of S in P, in the ratio P₄S and P₂S, did not solidify homogeneously, but that P separated from the first, and S from the second, solution. From these facts Isambert and Schulze concluded that the so-called P4S and P2S are merely solutions of S in P. Lemoine (C. R. 96, 1630) maintained that these substances were true compounds. According to Mai (A. 265, 192), when 67 parts P and 96 parts S are melted together, and the product is distilled at 10-11 mm. pressure, a distillate is obtained which slowly solidifies, and digestion of this solid with CS₂ under pressure gives a solution from which crystals of P4S, separate, and the mother-liquor deposits what seems to be a mixture of P₂S₃ and P₄S₅. The melting-points of mixtures of P and S are much below those of either constituent (Pelletier, A. Ch. [2] 4, 1; Faraday, A. Ch. [2] 7, 71; Dupré, A. Ch. [2] 73, 435). The experiments of Faraday (l.c.) and Ramme (B. 12, 940, 1350) show that the crystals obtained by dissolving S in liquid P sulphides-to which Berzelius gave the composition PS, and Dupré the composition PS,—are merely S with more or less adhering P.

Tetraphosphorus trisulphide P4S2 (Sesquisulphide of phosphorus. Hypophosphorus sulphide). Mol. w. 219·8. Melts at 166° (Ramme B. 12, 1350). Boils at 380° (Isambert, C. R. 96, 1499). S.G. 2·0 at 11°. V.D. 114·8 (Isambert, Lc.; Ramme, Lc.). H.F. [P¹,S³] = 36,800 (I., Lc.).

Formation.—1. By heating together P and S in the ratio 4P:3S .- 2. By heating P.S. with or-

dinary P to 320° (Ramme, l.c.).

Preparation.—A mixture of 1 part S with 1.3 parts amorphous P is placed in a flask with a long wide neck connected with a wide tube dipping beneath the surface of Hg; the mixture is heated towards 260°, when the reaction begins, and proceeds with production of much heat; the product is dissolved in CS_2 and crystallised (Lemoine, Bl. [2] 1, 407). Ramme (B. 12, 1350) heats the red P and S together in a sealed tube at 260° for eight hours. If ordinary P is used, combination occurs at 130°, with violent explosion. The explosion may be prevented by melting the S and P in the water-bath, and adding twice as much sand from which air has been removed by CO₂ (Isambert, C. R. 96, 1499). Ramme (B. 12, 1851) obtained P₄S₂ by heating P.S. with ordinary P in a sealed tube, filled with CO₂, to 320°.

Properties .- A yellow crystalline solid, which melts at 167° to a slightly coloured liquid. Boils 230°-240° at 11 mm. pressure, with partial decomposition to red P (Mai, A. 265, 192). Crystallises from CS_2 , PCl_3 , or $PSCl_3$ in rhombic prisms (Lemoine, Bl. [2] 1, 407); sublimes under 800°, probably in regular crystals (Isambert, C. R. 96, 1499). Very soluble CS₂, PSCl₂, and PCla; dissolved and decomposed by alcohol (Lemoine, l.c.). Burns at c. 100° to P₂O₄ and SO₂;

scarcely changed in ordinary air.

Reactions.-1. Burns when heated in air to e. 100°, forming P₂O₅ and SO₂.—2. Not acted on by cold water; hot water slowly produces H2S and H₂PO₂Aq.—3. Dissolves in potash solution, forming K phosphite and sulphide, and evolving H and P hydride (Lemoine, l.c.).—4. Hydrated lead oxide reacts at 200° with separation of PbS (L., l.c.).—5. Soluble in potassium sulphide solution, probably with formation of a double compound (L., l.c.).—6. Chlorine, when moist, slowly produces H₂SO₄ and H₂PO₄ (L., l.c.; I., l.c.).—7. Not acted on by cold sulphuric or hydrochloric acid; nitric acid and aqua regia oxidise P4S, slowly and regularly on warming.

PHOSPHOBOUS SULPHIDE P.S. (Trisulphide of phosphorus. Tetraphosphorus hexasulphide. Thiophosphorous anhydride). Mol. w. 315.72. Melts at c. 200°, and boils at c. 490°. V.D. 147.5, to 173.6 at lower temperatures (Isambert,

C. R. 102, 1386).

Formation.—1. By heating red P and S in the ratio 2P:3S in CO. Combination occurs without explosion, but so much heat is produced that a part of the product sublimes rapidly (Kekulé, A. 90, 310; Michaelis, A. 164, 22). 2. By the reaction of PCl, with H2S (Serullas, P. 17, 101). Ramme (B. 12, 940) failed to obtain P4S, by heating a solution of P and S in

CS, in a sealed tube.

Preparation.—A mixture is made of 1 part red P with 1.55 parts S; a small portion of this mixture is heated in a long-necked flask in a stream of CO2 until combination occurs; the flame is removed, and the rest of the mixture is introduced into the flask in small portions at a time; the heat produced by the combination of one quantity suffices to cause combination of the next quantity. The PAS is sublimed into the neck of the flask; or it may be dissolved in CS₂ and crystallised (Michaelis, A. 164, 22).

Properties .- A greyish-yellow, crystalline, solid (Kekulé, A. 90, 310; Isambert, C. R. 102, 1386). Tasteless, inodorous; does not fume in air; melts readily (c. 200°), and sublimes more easily than S. Soluble in CS₂. The V.D. shows that the molecular formula is P₄S₆, corresponding with the oxide P4Os. P4Ss acts as a thio-anhydride, forming compounds with the sulphides of several metals (v. Reactions, No. 7). As P₄S₆ is acted on by moist air, it must

be kept in sealed tubes.

Reactions.—1. Decomposes rapidly in moist air (? to H₂S and H₃PO₃).—2. Decomposed by water to H2S and H3POAq.-3. Burns when heated in air.—4. Dissolves in potash, soda, or ammonia solution, and is reppd. in yellow flocks by addition of acid.—5. Decomposes carbon compounds containing OH, forming corresponding SH compounds and H,PO,.-6. With todine in CS₂ forms PI₂ (Isambert, C. R. 96, 1771).—7. According to Berzelius, P₂S₂ combines with many metallic sulphides forming compounds 2R¹₂S.P₂S₂ and 2R²S.P₂S₂ the compounds 2R¹₂S.P₂S₃ and 2R²S.P₂S₃ the compounds 2R¹₂S.P₂S₃ and 2R²S.P₂S₃ the compounds 2R²₂S.P₂S₃ and 2R²S.P₂S₃ the compounds 2R²S.P₂S₃ 2R²S.

sition of which would lead to their being called thiopyrophosphites; R = Cu, Fe, Hg, Ag.—8. By dissolving in caustic soda solution cooled to 0°. Lemoine (C. R. 93, 489; 98, 45) obtained the compound P₂OS₂Na₂O.5H₂O and the compound P₂OS₂3Na₂O.4H₂O; and by using ammonium sulphide he obtained P₂OS₂2(NH₁)₂O.3H₂O and P2O2S.2(NH4)2O.6H2O. These substances evolved H2S by the action of HClAq; heated to 200°-240° they gave off H2S, but retained S; they may perhaps be called thio-oxypyrophosphites.

TRIPHOSPHORUS HEXASULPHIDE P₈S₆ (Phosphorus disulphide [PS₂]. Phosphorus tetrasulphide [P₄S₄]). ? Mol. w. 284·8. Melts at 296°–298° (Ramme, B. 12, 940); at 248°–249° (Seiler, Dissertation, Göttingen, 1876). Boils at 835°–240° 340° at pressure of 10-11 mm. (Mai, A. 265, 192). V.D. Ramme (B. 12, 1350) says that V. D. agrees with formula P.S., but no numbers are given.

Formation.—1. By heating a solution of P and S in CS₂ in sealed tubes to 210°; according to Ramme (B. 12, 940), P₂S₆ is produced with P and S in the following ratios:—P;S, P:2S,

2P:S, 3P:S, 4P:S, 6P:S (v. also Dervin, Bl. [2]

41, 433).-2. By heating together P and Sin the ratio P: 2S (Seiler, I.c.).

Preparation.—P and S are dissolved in CS, in the ratio P:2S, the solution is heated in a sealed tube to 210° for 8-10 hours, after cooling the liquid is poured off, and the crystals are recrystallised repeatedly from CS, in a sealed tube

(Ramme, l.c.).

Properties and Reactions. - Long, clear yellow, needle-shaped crystals. Ramme (l.c.) says that V.D. corresponds with formula P.S. but he gives no data; the corresponding oxide is generally written P₂O₄, but its molecular weight has not yet been determined. Heated with water in a sealed tube to 150° for some hours, P.S. gave H.S., H.PO.Aq, and H.PO.Aq, and an orange-yellow solid insoluble in CS, and not melting at \$10° (Ramme, B. 12, 1850).

PHOSPHORIC SULPHIDE P₂S₅ (Phosphorus pentasulphide or persulphide. Thiophosphoric anhydride). Mol. w. 221·8. Melts at 274°-276° (V. a. C. Meyer, B. 12, 610). Boils at 518°, pressure = 730 mm. (Goldschmidt, B. 15, 803); at 520° (Isambert, C. R. 102, 1386); at 530° (Hittort B. 126, 106). at 332° 240° pages 530° (Hittorf, P. 126, 196); at 332°-340°, pressure being 11 mm. (Mai, A. 265, 192). V.D. 110.7, at 580° in N (V. a. C. Meyer, B. 12, 610).

Formation.—1. By heating a solution of ordinary P and S, in the proper ratio, in CS₂ to 210° for 8-10 hours, and crystallising from CS₂ (Ramme, B. 12, 940).—2. By melting together red P and S, in ratio 2P:5S, in CO₂ (Kekulé, A. 90, 810); with ordinary P violent explosion

Preparation.—A mixture of 40 parts S and 18.5 parts red P is melted together, the mass is powdered, placed in a retort filled with CO2, and distilled to c. $\frac{1}{8}$ to $\frac{1}{8}$ (to remove P, S, and more volatile P sulphides); the residue is again powdered and distilled in CO2; the distillate is pure

Soluble in CS2; P2S3 seems to react as a thio-

anhydride (v. Reactions, No. 7).

Reactions.-1. Decomposes in moist air or by water, to H₂S and H₂PÕ₄; burns when heated in air to P₂O₄ and SO₂.—2. With phosphorus pentachloride forms PSCl₃ (Weber, J. 1859. 80). 3. With carbon tetrachloride at 200° produces CS₂ and PSCl₃ (Rathke, Z. 1870. 57).—4. With ferric chloride gives FeS, and PSCl, (Glatzel, B. 23, 37).—5. With sulphonyl chloride, PSCl, SO, and S are formed (Prinz, J. Z. 13, Supplbd. 1, 90) .- 6. With many carbon compounds containing O and also with many inorganic oxides, P2S, form S compounds, exchanging S for O (Kekulé, A. 90, 310; Carius, A. 106, 331; 112, 180; v. also Prinz, l.c.).—7. According to Berzelius (A. 46, 129, 255), P_2 S, combines directly with several metallic sulphides, forming compounds 2R12S.P2S, and 2R11S.P2S, (R=Cu, Fe, Mn, Hg, Zn): these compounds may be called thiopyrophosphates.
Thio-oxyorthophosphates. Salts of the hypo-

thetical acids H_aPSO_a (monothio-oxyorthophosphoric acid) and $H_aPS_aO_a$ (dithio-oxyorthophosphoric acid) phoric acid) have been obtained by the action of alkali solution on P₂S₅ or on PSCl₂. Salts of trithio-oxyphosphoric acid (H₂PS₂O) have not

been isolated.

Monothio-oxyphosphates. The Na salt, Na,PSO,.12H2O was obtained by Wurtz by warming PSCl, with fairly conc. NaOHAq (A. Ch. [3] 20, 472), and by Thorpe a. Rodger by the action of NaOHAq on PSF, (C. J. 55, 318). Kubierschi (J. pr. [2] 31, 93) obtained the salt by adding powdered P₂S₅ to NaOHAq, in ratio P2S.:6NaOH, cooling, adding absolute alcoholwhich ppd. a mixture of Na,PSO, and Na,PS,O, (leaving Na polysulphides in solution)—dissolving the pp. in water, warming to 90° till H,S ceased to come off, evaporating, and crystallising. Thin six-sided prisms, melting at 60°. The Mg salt (with 20H₂O), and the Mg-NH₄ salt (with 9H₂O) were also obtained.

Dithio-oxyphosphates. The Na salt, Na₃PS₂O_{4.11H₂O was obtained (K., l.c.) by dis-} solving in water the pp. obtained as above by alcohol, warming to 50°-55°, ppg. by alcohol, and crystallising from water at 40°. Colourless, lustrous, needles, melting at 45°-46°. The NH salt (with 2H2O), Ba salt (with 8H2O), and Mg-NH, salt (with 6H2O) were also obtained in

crystalline form.

For reactions of these thio-oxyorthophosphates v. Kubierschi, l.c. (cf. Michaelis, B. 5, 5). Phosphorus, sulphobromides of. Two comounds, corresponding with the two oxychlorides,

have been prepared.

THIOPHOSPHORYL BROMIDE PSBr. (Sulphophosphoryl bromide. Phosphorus sulphobromide). Formula probably molecular, from analogy of POCI,

Formation.—1. By reaction of H₂S with PBr₅ (Gladstone, P. M. 35, 345; Baudrimont, A. Ch. [4] 2, 58).—2. By distilling PBr, with S, or Br with P₂S₃.—3. By the action of Br on a solution of P and S in CS2.

Preparation.—Equal parts of P and S are dissolved in CS₂, the solution is well cooled, and 8 parts Br are allowed to drop in slowly from a funnel with a stopcock; the CS2 is removed by warming in a water-bath, and the liquid is then and H.PO.Aq. Prepared by adding 80 parts Br

distilled quickly over a large flame. Part of the PSBr, solidifies in the neck of the retort, and part passes over and remains liquid in the receiver; the contents of the receiver are shaken with fresh quantities of water until the oily liquid solidifies to a yellow crystalline mass which is PSBr₃.H₂O; this hydrate is dissolved in CS2, water is removed by contact with CaCl2, CS₂ is distilled off in a stream of CO₂, and PSBr, remains as a yellow oil, which solidifies on touching it with a glass rod. The solid may be recrystallised from PBr, (Michaelis, A. 164, 36).

Properties. - Yellow octahedral crystals, with an aromatic but unpleasant odour; vapour attacks the eyes. Melts at 38°, when molten remains for a long time without solidifying unless touched by a solid (Michaelis, B. 4, 777). Easily soluble in ether, CS₂, PBr₃, or PCl₃. Cannot be distilled without decomposition.

Reactions.—1. Decomposed by heat to S and a yellow liquid, PSBr, PBr, which is resolved by repeated distillation into S and PBr, and by repeated washing with water yields PSBr, Ho. (Michaelis, A. 164, 36).—2. Decomposed by water, slowly when cold, rather more rapidly when boiling, forming H₂S, S, H₂PO₄Aq and H₂PO₄Aq (v. Michaelis, B. 5, 4).—3. With alcohol forms PS(OEt)₃.—4. Decomposed by ammorphic solution slowly when cold ammonia solution, slowly when cold, more rapidly when hot, giving NH, phosphite and phosphate, H.S. S. and NH, sulphide (M., L.c.). Reacts with phosphorus pentachloride to form PSCl, and PBr, (M., l.c.).
 Combinations. — 1. With water to form

PSBr. H2O (v. Preparation). A yellow crystalline solid; melts at 35°, being resolved into H₂O and PSBr.; S.G. 2.7937 at 18°; gradually decomposes in air, giving off HBr (Michaelis, A. 164, 36).—2. With phosphorus bromide to form PSBr. PBr.; obtained by heating PSBr. (v.

Reactions, No. 1).

Pyrophoryl Browide P2S2Br4 (Phosphorus trisulphotetrabromide). A light-yellow oil, with aromatic and pungent odour; S.G. 2.2621 at 17°; fumes in air, with separation of S. Decomposed by distillation into S, P,S, and PSBr., PBr., i decomposed by water, giving S, PSBr., H₂S, P, and a substance probably P₂S₃(OH)₄; alkalis act similarly to, but more violently than, water, but neither S nor PSBr, is formed; with absolute alcohol yields P₂S₃(OEt)₂Br, P₂S₃(OEt)₄, P₂S₃(OEt)₂(SEt)₂; bromine reacts, when heated, forming PBr, and PSBr. (Michaelis, A. 164, 22). Prepared by moistening 100 g. finely powdered P.S. with CS. and adding 200 g. Br diluted with an equal volume of CS2, shaking thoroughly, distilling off CS2 in water-bath not above 80°, removing last traces of CS, in stream of CO, dissolving residue in six times its volume of dry ether, and removing ether by a stream of CO₂ (M., l.c.).

Phosphorus, sulphobromochloride of, PSBrCl, (Thiophosphoryl bromochloride). probably molecular, from analogy of POBrCl2 (Michaelis, B. 5, 6). A slightly yellow liquid, with aromatic odour; boils c. 150°, but boilingpoint soon rises and S separates. Decomposed slowly by water, more completely when heated in a sealed tube to 150° for some hours, giving S, H₂S, HClAq, HBrAq, H₂PO₂Aq,

drop by drop to 163 parts PCl₂.SEt [obtained by action of EtSH on PCl,; for details v. Michaelis, l.c.]; much heat is produced; the liquid is distilled, the distillate from 150°-180° is shaken with water so long as any action takes place; water is removed by a separating funnel; the liquid is dried by CaCl, and HCl and HBr are removed by warming gently (M., l.c.).

Phosphorus, sulphochlorides of. But one compound, PSCl_s, is known with certainty. is doubtful whether the substance obtained by Gladstone (C. J. 3, 5) by the reaction of S with PCl, is a definite compound PS2Cl, or a double

compound PSCl₂.SCl₂.

THIOPHOSPHORYL CHLORIDE PSCI, (Phosphorus sulphochloride). Mol. w. 169.05. Boils at 125·12° (Thorpe, C. J. 37, 341). S.G. 40 1·6682 (T., l.c.). V.D. 85 at 160°-300° (Cahours, A. Ch. [3] 20, 369; Chevrier, C. R. 68, 1174). For thermal expansion v. Thorpe (l.c.). S.V. 116·1.

Formation.—1. By the reaction of H₂S with PCl_s (Serullas, A. Ch. [2] 42, 25).—2. By heating S with PCl_s to 130° (Henry, B. 2, 638).—3. By heating P with S2Cl2 (Wöhler a. Hiller, A. 98, 274).—4. By heating together P₂S₅ and PCl₅ (Weber a. Thorpe, Z. 1871, 467).—5. By the reaction of PCl₃ with SO₂, SO₂Cl₂, or S₂Cl₂; and by the reaction of PCl₅ with CS₂, or with Sb₂S₅ (Michaelis; Cairns, A. 112, 190; 119, 291; Rathke, Z. 1870. 57; Baudrimont, J. pr. 87. 301).

Preparation.—An intimate mixture of P2S, and PCl_s , in the ratio $P_2S_s: 3PCl_s (=1:2.82)$, is heated in a sealed tube to c. 120°; after a short time the liquid which has formed is distilled. Should the product have a yellowish colour, it is shaken with a little water, dried by CaCl2, and distilled. The equation $P_2S_5 + 3PCl_5 = 5PSCl_2$ is realised (Thorpe, C. J. 37, 341).

Properties.—A colourless, highly refractive liquid; with an aromatic, somewhat pungent, and very peculiar odour, which becomes very marked on warming. Slowly decomposed by water.

Reactions. -1. Very slowly acted on by water, with formation of H₂PO₄Aq, HClAq, and H₂S. When distilled in steam the greater part of the PSCl₂ passes over unchanged.—2. Heated with alcohol under pressure, PS(OEt), is formed.—8. Passed through a hot tube, with excess of hydrogen sulphide, P₂S₅ and HCl are formed (Baudrimont, J. pr. 87, 301).—4. Decomposed by chlorine with formation of S₂Cl₂ and PCl₅. by chartne with formation of S_2O_2 and FO_3 .—

5. Silver nitrate reacts rapidly, producing $A_{G_3}PO_4$, $A_{G_2}O_5$, NOCl, and NO_2 (the SO_2 and NO_2 then reacting to form $S_2O_5(NO_2)_2$; Thorpe a. Dyson, C. J. 41, 297).—6. Alkali solutions form salts of H_3PSO_3 and $H_2PS_2O_2$ (v. Thiooxyonthophothemates, p. 147).—7. The prolonged action of ammonia—continued until 60 p.c. of the PSCl, has reacted with the NH, -produces thiophosphamide PS(NH2):(?); this compound remains as an amorphous, yellow-white solid on washing the product of the action of NH₂ on PSCl₂; S.G. 1.7 at 13°; decomposed at 200°, gives (?)PS(ONH₄), with hot water (Chevrier, C. R. 66, 748; Schiff, A. 101, 292). According to Gladstone a. Holmes (C. J. 18, 5), thiophosphamic acid PS(NH₂)(OH)₂ and thiophosphodiamic acid PS(NH₂)₂OH, are produced by the action of ammonia on PSCI, the former by using fairly dilute NH, Aq, and the latter by using NH, gas-which probably forms P(NH₂)₂ClS—and then washing with water. The work of Chevrier (l.c.) and Schiff (l.c.) makes the isolation of these acids doubtful.

Phosphorus, sulphochlorobromide of; PHOSPHORUS SULPHOBROMOCHLORIDE, p. 147.

Phosphorus, sulphocyanide of, P(SCN), (Phosphorus thiocyanate. Phosphorus rhodanide). Mol. w. not determined; formula probably molecular. Prepared by mixing 5 parts Pb(SCN), with an equal weight of dry sand, gradually adding 1 part PCl₃, digesting for some time on a water-bath, and then distilling very carefully from a hard glass tube. A liquid which does not solidify at -20° , begins to boil $260^{\circ}-270^{\circ}$, and decomposes at a higher temperature, giving off CS,; vapour is spontaneously inflammable; S.G. 1.625 at 18°. Soluble in alcohol, ether, CHCl, CS₂, and C₆H₆. Slowly decomposed by cold water to H₃PO₃Aq and HSCNAq (Miguel, A. Ch. [5] 11, 349).

Phosphorus, sulphofluoride of, PSFs (Thiophosphoryl fluoride). This compound was prepared and fully described by Thorpe a. Rodgers in 1888 (preliminary notice in C. J. 53, 760; fuller account in C. J. 55, 306). Mol. w. 119.9.

V.D. 59 6 at ordinary temperature.

Formation.-1. By heating a mixture of AsF, and PSCl, in ratio AsF, : 4PSCl, in a sealed tube at 150° .—2. By heating PbF₂ mixed with P_2S_a to c. 120° - 150° ; or by using BiF₃ in place of PbF₂ and heating to a higher temperature. A mixture of red P, S, and a large excess of PbF₂ may also be used $(P_2S_5 + 3PbF_2 = 3PbS + 2PSF_3)$. 3. By heating S with PF_3Cl_2 to 115° (Poulenc,

C. Ř. 113, 75).

Preparation.—A quantity of P2S, (freshly prepared from well-washed and perfectly dry amorphous P and roll S) is quickly mixed with c. 31 parts of pure freshly fused PbF2, and the mixture is placed in a thin equal layer in a dry tube of lead or 'composition,' one end of which is fitted with a caoutchouc cork carrying a glass delivery tube dipping under dry Hg, and the other end is connected with an apparatus for supplying pure dry N. A rapid stream of N is passed through till the air is driven out, the tube being gently heated to get rid of traces of H2S (produced by moisture on P₂S₂); the N is stopped, and the tube is heated from behind forwards by a small flame; reaction begins c. 170°; the temperature should be kept as low as possible, it must not exceed 250°. The gas is collected in a dry glass gas-holder containing a few small pieces of quicklime, the gas being allowed to pass into the holder as soon as a sample is wholly absorbed by dilute potash or ammonia. Before collecting the gas, small quantities of dry N should be passed into the gasholder and repeatedly exhausted by the Sprengel pump (this removes air from the pores of the lime), and when 2 or 3 c.c. of the PSCl, have passed into the holder the gas should be sucked out by the Sprengel, and this repeated twice or thrice (the traces of N are thus removed). After standing a few days over the lime, which removes PF, and any traces of SiF4, the gas is pure PSF2.

Properties.—A transparent, colourless gas; liquefied at 7.6 atmos. at 8.8°, 9.4 atmos. at 10°.

10.8 atmos. at 13.8°, and 13 atmos. at 20.3°. PSF₃ is spontaneously inflammable in air or O; it is decomposed easily by heat or electric sparks to S, P, and fluorides of P. Somewhat soluble in ether. Not easily dissolved by H₂O, or by solutions of alkalis. No action on Hg, H₂SO₄, CS₂, or C₂H₂. Neither gaseous nor liquid PSF₃ acts appreciably on glass at ordinary temperatures.

Reactions.—1. A small stream of PSF, issuing from a Pt jet into air at once takes fire; when mixed with a large volume of air, explosion occurs. - 2. Oxidised rapidly and explosively by oxygen; probably with formation of PF, and SO_2 , the PF, and O then producing P_2O_5 and also POF_3 . The ignition-temperature of PSF, is very low, and the flame is a very cold (Full details are given by T. a. R., l.c., pp. 312-317.)—3. Slowly decomposed by water, thus: $PSF_3 + 4H_2O + Aq = H_2S + H_3PO_4Aq + 3HFAq$. 4. Solution of potash or soda slowly reacts, forming Na₃PSO₃ and NaF.-4. With ammonia gas produces a white, deliquescent solid, probably a mixture of $\mathrm{NH_4F}$ and $\mathrm{P(NH_2)_2SF}$. For action of water on this substance—perhaps resulting in formation of PS(NH2)2OH-v.T. a. R., l.c., p. 319.-5. Heated in a glass tube, PSF_s gives SiF., P, and S.—6. PSF, is decomposed by electric sparks passed from Pt terminals, with separation of S and P, and a gas which is probably PF₅.

Phosphorus, sulphoselenide of. All attempts to prepare this compound have led to negative results. Michaelis heated PCl₃ with Se; Baudrimont (A. Ch. [4] 2, 5) tried the reaction of PCl₃ with SeCl₂, PCl₃ with P₂Se, Sb₂Se₃, and PbSe, and

the action of P on SeCl, and SeCl,

Phosphorus, sulphoxide of, P₄O₆S₄. Mol. w. 347·52. Melts at 102°, and boils at 295°. V.D. 171·3 at 350°-400°. Formed by heating small quantities P₄O₆ with S in an atmosphere of CO₂ or N at c. 160° (Thorpe a. Tutton, C. J. 59, 1022).

Preparation.—A strong glass tube is closed at one end, and filled with dry CO₂ or N; from 3 to 5 g. freshly-distilled P₄O₆ is placed in the bottom of the tube, and 1.74 g. S. best in small crystals, are added for every 3 g. P₄O₆ used (ratio P.Os: 4S). The tube is sealed, and the lower half is immersed in a glycerin-bath, which is gradually heated. At c. 154°-165° the melted S is suddenly projected to the top of the tube, and in a few seconds the action is complete. The product is transferred to a similar tube, which is exhausted by a Sprengel pump, sealed, and warmed at its lower portion; sublimation begins at c. 90°, and proceeds best at 140°-150° with formation of long feathery needles; part of the P4O8S4 remains as a vitreous or crystalline mass

Properties and Reactions.—A white solid, occurring in feathery aggregations or a vitreous mass, or as colourless, isolated, tetragonal prisms; melts 102°, and boils 295° (cor.). Very deliquescent, smelling in air of H₂S; quickly dissolved by water, forming H₂S and HPO₂Aq, which soon changes to H₂PO₄Aq. Easily sol. in 2 vols. CS₂, from which it crystallises unchanged. Also sol. benzene, but with decomposition.

Phosphorus, sulphur acids of, salts of; v. beginning of article Phosphorus sulphides

(p. 145); PHOSPHOROUS SULPHIDE, Reactions 7 and 8 (p. 146); PHOSPHORIC SULPHIDE, Reaction 7 (p. 147); and Thio-oxyorthophosphates (p. 147).

Phosphorus, telluride of. Oppenheim (J. 1857. 214) obtained a black amorphous solid, giving off fumes of P₂O₃ in air by heating P with

powdered Te.

Phosphorus, thio-acids of, salts of; v. beginning of article Phosphorus sulphides (p. 145); Phosphorus sulphide, Reactions 7 and 8 (p. 146); Phosphorus sulphide, Reaction 7 (p. 147); and Thio-oxyosthophosphates (p. 147).

147); and Thio-oxyorthophosphates (p. 147).

Phosphorus. thio-amic acids of, and thio-amide of; v. Thiophosphoryl chloride, Re-

action 7 (p. 148).

Phosphorus, thiobromides of; v. Phosphorus sulphobromides, p. 147.

Phosphorus, thiobromochloride of; v. Phosphorus sulphobromochloride, p. 147.

Phosphorus, thiochlorides of; v. Phosphorus sulphochlorides, p. 148.

Phosphorus, thiocyanide of; v. Phosphorus sulphocyanide, p. 148.

Phosphorus, thioselenide of; v. Phosphorus sulphoselenide, ante.

M. M. P. M.

in this article.

Besides the phosphoric acids, three oxyacids of P, and some salts of a fourth acid, have been isolated; the acids in question are H₃PO₂, H₃PO₃, H₂PO₃ or H₃P₂O₆; and salts of H₄P₂O₅. No anhydride of H₃PO₂ is known; the hypothetical anhydride would be P₂O (P₂O+3H₂O = 2H₃PO₂); P₂O₃ is the anhydride of the acid H₃PO₃ (P₂O₃+3H₂O = 2H₃PO₃), and also the hypothetical anhydride of H₄P₂O₅(P₂O₃+2H₂O = H₄P₂O₅); P₂O₄ is the hypothetical anhydride of H₄P₂O₅(P₂O₄+2H₂O = H₄P₂O₅), but this oxide gives H₃PO₃Aq and H₃PO₄Aq with water (o. Phosphorus tetracylde, p. 141). The oxyacide of P do not correspond in composition with those of N (v. Table in Nitrogen group of elements, vol. iii. p. 575).

HYPOPHOSPHOROUS ACID SALTS H₃PO₂; M₁H₂PO₂ and M₁(H₂PO₂)₂. Hypophosphites are formed by decomposing phosphide of Ba, Ca, or Sr by water (Dulong, A. Ch. [2] 2, 141); by heating P with potash, milk of lime, or BaOAq (H. Rose, P. 9, 225, 361; 12, 77, 288); by passing PH₂ into solution of alkalis or alkaline hydrates (Winckler, P. 111, 443); and by heating P with syrupy H.PO.Aq to 200° (Oppenheim, Bl. [2] 1, 163). The acid is monobasic; it may be regarded as derived from the hypothetical anhydride P.O. There is no acid, nor are there salts, corresponding with hyponitrous acid HNO, but some salts are known of the form MPS; v. Thiometahypophosphites, at beginning of Phosphorus, sulphides or (p. 145). Selenometahypophosphites M'PSe have also been isolated (v. DIPHOSPHORUS SELENIDE, p. 144).

HYPOPHOSPHORUS ACID H₄PO₂. Melts at 17·4° (Thomsen, B. 7, 994; ? specimen pure). H.F. [H²,P,O²] = 139,970 (production of crystallised

acid); 187,660 (production of acid in liquid |

state); [H³,P,O²,Aq] = 189,800 (Th. 2, 225).

Preparation.—Ba(H,PO₂)₂;H₂O is prepared by warming water with BaO and P (in small pieces) in a basin till inflammable P hydride ceases to come off, filtering, removing excess of BaO by passing in CO₂, filtering again, and crystallising. 285 g. pure Ba(H₂PO₂).H₂O are dissolved in c. 5 litres water, 98 g. H₂SO₄ (c. 101.5 g. ordinary conc. acid), diluted with 3 to 4 times its weight of water, are added; after shaking for some time the pp. of BaSO, is allowed to settle, and the clear liquid is syphoned off, and evaporated in a porcelain dish, by boiling, till it is c. 10 of the original volume, when it is placed in a Pt dish; a thermometer is immersed in the liquid, evaporation is continued, the temperature being allowed to rise gradually to 110°; any solid which separates is filtered off while the liquid is hot, and the filtrate is evaporated, without being allowed to boil, at c. 110°. Thomsen (B. 7, 994) finally heats for 10 mins. to 130° or 138°, filters into a stoppered bottle, and cools below 0°; Geuther a. Ponndorff (J. Z. 10, Supplbd. 2, 45, 49) say that heating above 110° decomposes H₃PO₂, and that Thomsen's preparation therefore contained H.PO.

Properties.—Large, white, crystalline tablets; melting at 17.4°; remains liquid considerably above m.p. (Thomsen, l.c.). Decomposed by heat to PH, and H,PO. H,PO.Aq is a very energetic reducing agent. The acid is monobasic, forming

salts M¹H₂PO₂ and M¹(H₂PO₂).

Reactions.—1. Decomposed by heat, giving P hydride and H₂PO₄ (H. Rose, l.c.).—2. H₂PO₂Aq is readily changed to H₃PO₈Aq or H₂PO₄Aq by oxidisers; Cl. Br. HNO₃Aq, KMnO₄Aq, and several metallic oxides produce H₂PO₄Aq (Dulong, V. P. 2007, 100 and 10 l.c.; Rose, l.c.; P. de St. Gilles, A. Ch. [3] 55,374); PbO₂ forms PbHPO₃ (Wurtz, C. R. 18, 702); SO₂ forms H₃PO₃Aq and S (Geuther a. Ponndorff, l.c.). According to Engel (C. R. 110, 786) Pd charged with Hoxidises H,PO,Aq to H,PO,Aq with evolution of H.—3. H.PO, Aq reduces solutions of salts of gold, silver, and mercury, ppg. the metals; from copper sulphate solution, Cu₂H₂ is ppd. at 55°-60° (Wurtz, l.c.; Rammelsberg, l.c.).—4. Reduced by nascent hydrogen to PH₄ (Dusart, C. R. 43, 1126; Blondlot, C. R. 52, 1197).— 5. With hydriodic acid forms PH,I (G. a. P., l.c.).-6. Heated with conc. sulphuric acid, H.PO₄, SO₂, and S are produced (Wurtz, l.c.). 7. Heated with phosphorus pentachloride, POCl PCl, and HCl are formed; phosphorus trichloride produces P, H₂PO₂, and HCl; with phosphoryl chloride, P, PCl₃, HPO₂, and HCl are the products (Geuther, J. pr. [2] 8, 359)

HYPOPHOSPHITES MIH2PO2 and MII(H2PO2)2. These salts are formed in several reactions (v. beginning of this article); the greater number are prepared by reacting on the salt of Ba with the sulphates of other metals; the salts of Ba, Ca, and Sr are generally prepared by heating BaOAq, CaOAq, and SrOAq with P. The hypophosphites of Cd, Ca, Pb, and Tl crystallise without water; those of Ba, Li, Mn, Na, Sr, and U with one H₂O; those of Co, Mg, Ni, and Zn with 6H₂O (Rammelsberg, C. J. [2] 11, 12). They are decomposed by heat, evolving P hydride and H, and generally leaving pyro- and meta- phos-phate, the Co and Ni salts leave metaphosphate and phosphide, and the U salt leaves pyro- and meta- phosphate and phosphide (R., l.c.). The hypophosphites are easily oxidised to phosphites and phosphates, hence they react as reducing agents. Heated in solution with alkalis they give phosphites, and then phosphates, and evolve H (Wurtz; H. Rose, l.c.). The chief memoirs on hypophosphites are by H. Rose (P. 9, 225, 361; 12, 77, 288); Wurtz (A. Ch. [3] 7, 193; 16, 196); and Rammelsberg (C. J. [2] 11, 1, 13).

Ammonium hypophosphite NH, H, PO2. Hexagonal tablets; by decomposing the Ba salt by (NH₄)₂SO₄Aq; easily sol. absolute alcohol (Wurtz,

Rose).

Barium hypophosphite Ba(H₂PO₂)₂.H₂O. Lustrous monoclinic needles (Rammelsberg; Topsöe, W. A. B. 69 [2] 19). By heating BaOAq with P till gas ceases to come off, filtering, removing BaO from the filtrate by CO2, filtering, and crystallising. Loses H₂O at 100°. Soluble in 31 parts cold, and 3 parts boiling, water; insol. alcohol. Heated out of air leaves pyro- and meta- phosphate (Wurtz, Rose, Rammelsberg).

Calcium hypophosphite Ca(H₂PO₂)₂. Prepared like the Ba salt. Berlandt (Ar. Ph. [2] 122, 237) recommends 29 parts P, 47 parts CaO₂H₂, and 24 parts water. Bachmann and Martenson (J. 1864. 191) decompose Ca phosphide by boiling water. Thin, monoclinic tablets (Rammelsberg; Schabus, J. 1854. 325). bitter taste; unchanged in air; does not give off H₂O at 300°; heated to redness gives off H, PH₃, and H₂O, and leaves pyro- and meta- phosphate (Rammelsberg; Michaelis, J. 1872. 210). Sol. 6 parts cold water, not much more sol. hot water; insol. strong alcohol (Rose).

Cobalt hypophosphite Co(H2PO2)2-6H2O. Red octahedral crystals, isomorphous with Mg salt (Rose); by decomposing Ba salt by CoSO, Aq, filtering, and evaporating in vacuo (Wurtz). Forms a double salt with Ca(H,PO₂)₂ (Rose). Copper hypophosphite Cu(H,PO₂)₂. Obtained

with difficulty from Ba salt and CuSO, Aq; solu-

tion easily decomposes, giving Cu₂H₂ (Wurtz).

Lead hypophosphite Pb(H₂PO₂)₂. By adding PbO or PbCO₂ to H₃PO₂Aq (Rose, Wurtz).

Manganese hypophosphite Mn(H,PO),-H₂O. Small, rose-red crystals; from Ba salt and MnSO,Aq (Wurtz, Kammelsberg); or by boiling Ca salt with Mn oxalate (Rose).

Potassium hypophosphite KH2PO2. Prepared by decomposing the Ba or Ca salt by K2CO2Aq, filtering, evaporating to dryness, treating the residue with alcohol, and crystallising the alcoholic solution in vacuo; or by dissolving P in boiling KOHAq, evaporating, and dissolving out from the residue by alcohol; also by decomposing the Ba salt by K₂SO₄Aq (Rose, Wurtz). A white, semi-crystalline mass. Easily sol. water and alcohol; very hygroscopic (Dulong, A. Ch. [2] 2, 141). Burns when heated in air; oxidised violently by evaporation with HNO, Aq. Heated out of air evolves inflammable P hydride, and leaves pyro- and meta- phosphate (Rammelsberg).

Sodium hypophosphite NaH, PO, H,O. pared like the K salt. Small, rectangular tablets, easily sol, water and absolute alcohol. Evaporation of the solution is attended with explosions (Marquart, Ar. Ph. [2] 95, 284; Trommsdorff,

ibid. 99, 388).

The salts LiH,PO,H,O, Ni(H,PO,),6H,O,

Sr(H₂PO₂)₂, and Zn(H₂PO₂)₂ are also described; and salts of Al, Be, Cd, Cr, and Fe have been obtained, but their compositions are somewhat doubtful.

PHOSPHOROUS ACID AND SALTS M12HPO2, H_sPO_s; MIIHPO, M¹H₂PO₂, Mul (HPO3)3. P2O3 is the anhydride of H,PO3; the acid is produced by dissolving the oxide in cold water, but the oxide is not obtained by removing H2O from the acid. H3PO, is dibasic; the salt Na₃PO₈, said to be formed by adding a large excess of NaOH to H₃PO₃ (Zimmermann), has been shown by Amat (C. R. 108, 403) not to exist. The acid is formed by oxidising P slowly in moist air; the salts are obtained by reactions between the acid and metallic hydroxides, or by double decomposition from the alkali salts. Neither the acid nor salts of the acid corresponding with nitrous acid, HNO2, are known, nor have thio- salts of this form been isolated.

Phosphorous acid H_3PO_3 . Melts at 74° (Geuther a. Hurtzig, A. 111, 159); at 70·1° (Thomsen, J. pr. [2] 8, 359). H.F. $[H^3,P,O^3] = 227,700$ (crystallised acid); 224,630 (acid in liquid state); 227,570 (acid in aqueous solution) (Th. 2, 225). Heat of fusion = -3070; one gram-molecule of the liquid acid occupies 49·66 c.c. (Th. 2, 224).

Formation.—1. Along with H₂PO₂ and H₂PO₄, by the long-continued oxidation of P in moist air (v. J. Corne, J. Ph. [5] 6, 123).—2. By dissolving P in dilute HNO₂Aq and allowing to oxidise in air.—3. By the oxidation of H₂PO₂Aq by weak oxidisers.—4. By dissolving P₂O₃ in cold water (Thorpe a. Tutton, C. J. 57, 567).—5. By decomposing PCl₂ by H₂O₃ or by warming with H₂C₂O₄ (Hurtzig a. Geuther, A. 111, 159); or by leading Cl into warm water in presence of an excess of molten P (Droquet, P. 12, 628).

Preparation.—1. A small quantity of PCl, is shaken with cold water, the reaction is moderated by plunging the beaker into cold water; when the PCl, is all decomposed, a little more is added, and so on $(PCl_3 + 3H_2O + Aq = 3HClAq + H_3PO_3Aq)$. The solution is evaporated, the temperature being raised to 180° near the end of the process; the syrupy liquid thus obtained is placed over H2SO4, in vacuo, till it crystallises; addition of a crystal of HsPOs causes rapid crystallisation (Thomsen, B. 7, 996). 2. A stream of dry air is passed through PCl, at 60°, and then into two flasks, each containing c. 100 g. water at 0°; after c. 4 hours the first flask is filled with crystals, which are drained under a pump, washed with small quantities of ice-cold water, and dried in vacuo (Grosheintz, Bl. [2] 27, 483).—3. A mixture of 1 part PCl, and 2 parts dry H2C2O4 is heated in a flask with a reversed condenser until a clear liquid is obtained; the liquid is evaporated in a stream of CO2, and the crystals are washed and dried as in 2 (Hurtzig a. Geuther, A. 111, 159: PCl₃ + 8H₂C₂O₄ $= H_3PO_3 + 3CO + 3CO_2 + 3HCl$).

Properties.—A colourless, crystalline mass; obtained in transparent crystals by evaporating H,PO₂Aq in vacuo over H₂SO₄. Melts at 74° (H. a. G., l.c.); at 70·1° (Thomsen, l.c.). Heated above its m.p. decomposes to H₂PO₄ and PH₃. H₂PO₅ is very deliquescent, and oxidises very slowly in air; it acts as an energetic reducing

agent. H_3PO_3 is dibasic, forming salts $M^1H_2PO_3$ and $M_{12}HPO_3$.

Reactions.-1. Heated above its m.p., H.PO. decomposes to H₂PO₄ and inflammable P hydride (Vigier, Bl. [2] 11, 125; Rose, Davy, Gm.-K. 2, 115); according to Hurtzig a. Geuther (A.111, 159) Palso separates.—2. Slowly oxidised in air to H₂PO₄; dilute solutions of H₂PO₃ are more rapidly oxidised; according to Salzer (A. 232, 114), H₃PO₃Aq does not oxidise in air at ordinary temperatures. - 8. Converted into H.PO, by many oxidisers, e.g. HNO₃Aq, ClAq, BrAq, KMnO₄Aq, hypochlorites; with SO₂Aq, H,PO,Aq and H,S are formed along with S (Gm.-K. 2, 116).—4. Reduces many metallic salts in solution; e.g. AgNO, Aq to Ag, HgCl, Aq to HgCl and then to Hg, CuCl_Aq to Cu,Cl, and then to Cu (Rammelsberg, C. J. [2] 11, 13).—5. Nascent hydrogen forms PH₂ (Dusart, C. R. 48, 1126). 6. When crystalline H_sPO_s is heated with bromine to 100° in a sealed tube, H,PO, is formed along with HBr and PBr, or the products are HPO, and HBr, according to the relative quantity of Brused (Gustavson, J. 1867. 139). According to Ordinaire (C. R. 64, 363), a crystalline compound is obtained—perhaps bromo-phosphorous acid-by heating H.PO, and Br in the ratio H₃PO₃: 4Br.—7. By heating H₃PO₃ with iodine Gustavson (l.c.) obtained H₂PO₄, HI, PI₂, and PH₄I.—8. Heated with phosphorus trichloride to 170°, P, HCl, and H₂PO₄ or H₄P₂O₇ are formed (Geuther, J. pr. [2] 8, 359; Kraut, A. 158, 332; Gautier, C. R. 76, 49). At 79° HCl, H₄P₂O₇, and a compound PH₄O are formed, according to Gautier (l.c.) (v. Phosphorus, compounds of, with HYDROGEN AND OXYGEN, p. 134).—9. With phosphorus pentachloride, POCl, HCl, and PCl, are produced; and the products of the reaction with phosphoryl chloride are PCl, HPO, and HCl (Geuther, J. pr. [2] 8, 359).—10. By neutralisation with alkalis or alkaline carbonates salts of the form M2HPOs are produced.

Qualitative distinction between phosphorous and hypophosphorous acids. H₃PO₃Aq reduces CuSO₄Aq to Cu, while the reduction product with H₃PO₂Aq is Cu₂H₃.

PHOSPHITES M'H,PO,, M',HPO,, M''HPO,, M''HPO,, M''',(HPO,),. These salts are generally obtained by neutralising H,PO,Aq by metallic hydroxides or carbonates, or by double decomposition from the alkali salts. They are also formed by the slow oxidation of hypophosphites. Zimmermann's assertion that Na₂PO₂ exists (A. 175, 1) has been disproved by Amat (C. R. 106, 1351), who has shown that the normal Na salt is Na₂HPO₃. The alkali phosphites are soluble water; most of the others are insoluble. Phosphites are decomposed by heat, generally giving off H or PH, and leaving pyrophosphate and phosphide. Solutions of phosphites are scarcely changed in air, but they are readily oxidised to phosphates by energetic oxidisers. The chief memoirs on phosphites are those of H. Rose (P 9, 26, 224; 12, 77, 288), Rammelsberg (P. 131, 263, 359; 132, 481; B. 9, 1577), Kraut (A. 177 274), Wurtz (A. 58, 65).

Ammonium phosphites (NH.)H.PO, and (NH.)2HPO, 2H.O. The former is obtained by adding NH.Aq to H.PO.Aq till neutral to methyl orange, concentrating, and drying the crystals over H.SO.; melts 123°; above 150° gives off

NH₂ and PH₃, and leaves H₂PO₄; absorbs NH₃ at 80°-100°, forming (NH₄)₂HPO₃ (Amat, C. R. 105, 809). The salt (NH₄)₂HPO₃.2H₂O was obtained by Rose by adding slight excess NH,Aq

to the acid and crystallising; in vacuo at ordinary temperature, or at 100° in air, loses NH, and gives NH, H,PO, (Amat, l.c.).

Barium phosphites. The normal salt BaHPO, is obtained by adding BaCl,Aq to (NH,)HPO,Aq (Berzelius, Gm.-K. 2, 270), also by boiling BaHPO Aq with KOHAq. A crystal. by boiling BaHPO₂Aq with KOHAq. A crystaline powder; strongly heated gives Ba₂P₂O₇, H, and Ba phosphide (v. Rammelsberg, B. 9, 1577); very slightly sol. water; on boiling with water gives a basic and an acid salt (Dulong, Gm.-K. 2. 270). The acid salts (BaH (HPO)) BaH₂(HPO₃)₂, 2BaHPO₃, H₃PO₃,8H₂O, and 2BaHPO₃,3H₃PO₃,2H₂O are said to be formed by the action of H₃PO₃Aq on the normal salt (Rammelsberg, Rose, Wurtz).

Calcium phosphites. CaHPO₃.xH₂O; from CaCl₂Aq and (NH₄)₂HPO₃Aq (Wurtz, Rose, Rammelsberg). The acid salt CaH₂(HPO₄)₂.H₂O is obtained by dissolving marble in H₃PO₃Aq, and evaporating in vacuo (Wurtz).

Cobalt phosphite CoHPO3.2H,O. A reddish powder, obtained by CoCl₂Aq + (NH₄)₂HPO₃Aq, or by dissolving freshly ppd. CoCO, in H,PO,Aq; goes blue at 250° (Rose, Rammelsberg).

Copper phosphite CuHPO3.2H2O. By double decomposition from (NH4)2HPO8 (Rose); also by acting on Cu acetate by H₃PO₃Aq (Wurtz). Unchanged by boiling water (Rammelsberg), but reduced to Cu by boiling with H₃PO₃Aq (Rose).

Lead phosphites. The normal salt PbHPO, is obtained by ppg. Pb acetate by (NH₄)₂HPO₃Aq (Rose). A white powder; heated gives off H and PH₃, leaving Pb₂P₂O, and Pb phosphide. Insol. water (Amat, C. R. 110, 901). By adding NaH₂PO₃Aq to excess of Pb2NO₃Aq, the compound PbHPO₃.Pb(NO₃)₂ is obtained (Amat, *l.c.*). The *acid salt* PbH₂(HPO₃)₂ is formed by dissolving the normal salt in very conc. H_sPO_sAq, and drying the crystals at 100°; with water gives PbHPO, and H,PO,Aq (Amat, l.c.)

Magnesium phosphite MgHPO3.xH2O. By MgSO,Aq by Na HPO Aq boiling a double (Rammelsberg) Forms Mg₂(NH₄)₂(HPO₂)₄.16H₂O (Rammelsberg, Rose).

Manganese phosphite MnHPOs.H.O. A reddish powder; very slightly sol. water, more sol. solutions of Mn salts; loses H₂O at 200°. From NH₄ salt and a Mn salt; or by dissolving MnCO₃ in H₂PO₃Aq, and adding Na₂CO₃ till ppn. begins, but not enough to remove acid reaction (Rose, Rammelsberg).

Potassiur phosphites. The normal salt K.HPO, is formed from H.PO,Aq and KOHAq or K.CO.Aq; semi-crystalline, syrup-like mass; very hygroscopic; insol. alcohol (Wurtz, Rose). The acid salt K,HPO,2H,PO, was obtained by Wurtz (A. 58, 63). Amat (C. R. 106, 1351) obtained the acid salt KH,PO, by adding KOHAq or K,2CO,Aq to H,PO,Aq till neutral to methyl orange, concentrating, and cooling.

Sodium phosphites. The normal salt

under these conditions. Na₂HPO, melts at 53°; heat of solution at $13.5^{\circ} = -4600$ (Amat, C. R. 110, 191). Dehydrated by drying in vacuo, and then at 150° (A., l.c.). The acid salt 2NaH₂PO₂.5H₂O is produced by making H₂PO₂Aq neutral to methyl orange by NaOHAq or Na CO Aq, concentrating, and cooling. Melts 42°; dehydrated by long heating at 100°; heated above 150° forms $Na_4P_2O_5$; very sol. water; heat of solution at $15^{\circ} = -10,600$ (Amat, C. R. 106, 1351).

Zinc phosphite 2ZnHPO₃.5H₂O; from ZnSO₄Aq and ammoniacal solution of PCl₃ (Rose); also from PCl, Aq and ZnO (Rammels-

The following phosphites have also been described:—Di₂(HPO₃)₃ (Frerichs a. Smith, A. 191, 331); Fe₂(HPO₃)₃,9H₂O (Rose, Rammelsberg); La₂(HPO₃)₃ (F. a. S., l.c.); SnHPO₂ (Rose). Phosphites of Al, Be, Bi, Cd, Cr, ferrous Fe, Hg, Ni, and Sr have also been prepared, but their compositions are somewhat doubtful

(v. especially Rose, l.c.)

HYPOPHOSPHORIC ACID AND SALTS P are partly covered with water and left for some time in a large vessel, an acid liquid is obtained; this liquid was formerly called Pelletier's phosphatic acid (P., Crell. Ann. 1796 [2] 447). Dulong (A. Ch. [2] 2, 141) found 47.8 p.c. P and 52.2 p.c. O in this substance, and supposed it to be P_2O_9 . In 1856 Pagel (J. pr. 69, 24) showed that the acid liquid contained phosphoric and phosphorous acids; and in 1877 Salzer (A. 187, 322) proved the presence of small quantities of a new acid to which he gave the composition nH₂PO, and the name hypophosphoric acid. The acid has been found to be tetrabasic, hence the formula is written H₄P₂O₆. The hypophosphates are generally formed directly from the acid, or by double decomposition from the alkali salts. The hypophosphates are more stable than the hypophosphites and phosphites; normal salts pass into phosphates by taking up O; they are decomposed by heating strongly, giving of PH, or H, and leaving phosphate and phosphide.

HYPOPHOSPHORIC ACID H₄P₂O₆. Melts at c. 55° (Joly, C. R. 102, 110). Heat of solution

=3,850 (J., l.c.).

Formation .- 1. Along with HaPOs and H₃PO₄, by the slow oxidation of P in moist air (Salzer, A. 167, 322; 194, 28; 211, 1; 232, 114, 271).—2. The Ag salt is formed by oxidising P by dilute HNO, Aq in presence of AgNO, (Philipp, B. 18, 749); also by reacting on H₃PO₃Aq with AgNO₂Aq (Sänger, A. 232, 1); and the Cu salt by reacting on P with Cu(NO₂)₂Aq (Corne, J. Ph. [5] 6, 123).

Preparation.—Sticks of P are placed in glass tubes narrowed at the lower end; these are arranged inside a funnel, which is placed in the mouth of a bottle containing a little water; the bottle is set in a basin, and the whole is covered with a large bell-jar open at the top. The apparatus is placed in a cool place (best in Na₂HPO_{2.5}H₂O is obtained by dissolving H₂PO₂ a cellar) for some weeks or months; according in an excess of NaOHAq, and evaporating in to Joly (C. R. 101, 1058), from $\frac{1}{2}$ to $\frac{1}{2}$ of the P vacuo (Amat, C. R. 108, 408). Zimmermann (A. 175, 1) supposed that Na₂PO₃ is formed in winter, and not more than $\frac{1}{10}$ in summer.

The soid liquid is mixed with a cold conc. solution of Na acetate, the ppd. Na,H2P2O.6H2O is collected, washed with a very little cold water, then dissolved in water (sol. in c. 45 parts cold water), and ppd. by Pb acetate; the Pb2P2O6 is collected, washed with cold water, suspended in water, and decomposed by H2S; the filtrate is evaporated at a low temperature-not exceeding 30°-or in vacuo over H₂SO₄, until crystals of H₄P₂O₆ are deposited. Joly (C. R. 101, 1058, 1148) heats to boiling the acid liquid obtained by oxidising P, as described above, adds Na₂CO₂ till nearly neutral to methyl orange, and crystallises by concentration; he washes the crystals of Na₂H₂P₂O₆.6H₂O with cold water, and recrystallises from hot water, dissolves in hot water, adds an equivalent of BaCl₂ in hot water, washes the ppd. BaH₂P₂O₈, decomposes it by an equivalent of H₂SO₄ diluted with its own weight of water, filters after two or three days, and evaporates in vacuo over H₂SO₄. Drawe (B. 21, 3401) carries out the oxidation of P in presence of Na acetate solution.

Properties.—Small colourless crystals, which melt at c. 55° (Joly, C. R. 102, 110); very hygroscopic; solution is unchanged in air at ordinary temperatures. Heated to c. 70° suddenly decomposes to H₃PO₃ and HPO₃ (J., l.c.); at c. 120° gives H,P,O, and P hydride (J., C. R. 102,

Reactions and Combination.—1. Heat decomposes H₄P₂O₆ (v. supra).—2. Not oxidised by such oxidisers as H₂O₂Aq, ClAq, CrO₃Aq, HgCl₂Aq, &c.-3. Not reduced by such reducers as SO₂Aq, H₂S, or Zn and H₂SO₄Aq.—4. Potassium permanganate is slowly reduced in the cold, quickly on heating, by H,P2O6Aq to which H₂SO₄Aq has been added.—5. Heated with nitric acid, HaPOAAq is formed.—6. Heated with dilute sulphuric acid, H₃PO₃Aq and H₃PO₄Aq are produced .- 7. Silver nitrate solution gives a white pp., which does not blacken in light and is soluble in warm HNO3Aq (1:1).—8. By evaporating H₄P₂O₆Aq till the liquid had the composition P₂O₄.6H₂O₅ and then placing over H₂SO₄ in vacuo, Joly (C. R. 101, 1058; 102, 110) obtained crystals of the hydrate H,P2O6.2H2O, melting at 62°-62.5°.

HYPOPHOSPHATES. $M_1H_2P_2O_6$, $M^1_2H_2P_2O_6$, $M^1_3HP_2O_6$, $M^1_4P_2O_6$, $M^1H_2P_2O_6$, $M^1_2P_2O_6$. These salts have here examined chiral and $M^1_2P_2O_6$. salts have been examined chiefly by Salzer (A. 187, 322; 194, 28; 211,1; 232, 114, 271), and crystallographically by Haushofer a. Fresenius (Z. K. 1, 257, 620; 3, 605; 6, 113; 9, 254). The hypophosphates are much more stable towards oxidisers than the hypophosphites or phosphites. They reduce KMnO, Aq very slowly in the cold after addition of H2SO, Aq, and give white pp. with AgNO, Aq unchanged in light.

Ammonium hypophosphates. The normal salt (NH₄),P₂O₈,xH₂O is obtained by adding excess of NH₂Aq to the acid; on drying, loses NH₄ and effloresces. The acid salts (NH₄)₂H₂P₂O₄ and (NH₄)₃HP₂O₅ are formed, the former by boiling a solution of the normal salt, the latter by adding the proper quantity of H.P.O. to the former salt.

Barium hypophosphates. The normal 901).

salt Ba₂P₂O₆ is obtained from BaCl₂Aq and Na₆P₂O₆Aq; the acid salt BaH₂P₂O₆2H₂O is heating PbH₂(HPO₂)₂ in a dry vacuum to c. 140°

formed from NaH, P,O,Aq and BaCl,Aq (v. Joly, C. R. 101, 1058).

Cadmium hypophosphate Cd2P2O22H2O; and the double salt CdNa2P2O66H2O (Drawe, B. 21, 8401).

Calcium hypophosphates Ca.P.O.

CaH_P_O_6H_2O.

Cobalt hypophosphate Co_P_O_8H_2O; and the double salt 2CoNa_P_O_8H_O (Drawe, l.c.).

Copper hypophosphate Cu₂P₂O₆.6H₂O (Drawe,

Lead hypophosphate Pb₂P₂O₆.

Magnesium hypophosphates Mg₂P₂O₆.12H₂O and MgH2P2O84H2O; and the double salt Mg(NH₄)₂P₂O₆.6H₂O.

Nickel hypophosphate Ni₂P₂O₆.12H₂O; and the double salt NiNa2P2O6.12H2O (Drawe,

l.c.). Potassium hypophosphates K,P,O,8H,O, K₂HP₂O₆.3H₂O, K₂H₂P₂O₆, and KH₃P₂O₆

Silver hypophosphate Ag, P,O, (v. Philipp, B.

14, 749). Sodium hypophosphates. The acid salt Na₂H₂P₂O₆.6H₂Ō is obtained by slowly oxidising P in moist air, and adding Na acetate solution as described under Preparation of hypophosphoric acid (p. 154); sol. 45 parts cold, or 5 parts boiling, water; insol. alcohol; on strongly heating gives off H and P hydride and leaves NaPO₃. The normal salt Na₄P₂O₆.10H₂O is obtained by adding an equivalent of Na₂CO₈ to Na₂H₂P₂O₆Aq; sol. c. 30 parts cold water, much more sol. hot water; solution reacts alkaline, and gives a mixture of salts on evaporation. The acid salt Na₃HP₂O₆.9H₂O is formed by mixing the Na₄ and Na₄ salts in the ratio Na₂H₂P₂O₆. Na₄P₂O₆. The acid salt NaH, P,O, 2H,O is formed by adding the proper quantity of H,P,O,Aq to Na₂H₂P₂O₆. For reaction of the Na salts with different indicators v. Joly, C. R. 101, 1058.

Another acid salt Na, H, (P,O,)2.20H,O is said to be formed by adding rather more molecular weight than one NaOH4Na₂H₂P₂O₆.6H₂O.

Zinc hypophosphate Zn₂P₂O₆.2H₂O (Drawe, l.c.).

PYROPHOSPHITES. In 1887 Amat (C. R. 106, 1400) showed that when 2NaH₂PO₃.5H₂O is heated to 160° it loses 6H2O and gives the salt Na₂H₂P₂O₃. A few other salts of the form M₂H₂P₂O₃ have been obtained, but the acid has not been isolated. When the Ba salt is decomposed by an equivalent of H2SO, Aq, the solution shows the reactions of pyrophosphites, but the pyrophosphite gradually changes to phosphite (Amat, C. R. 108, 1056). The pyrophosphites in solution slowly change to phosphites, the more rapidly the more conc. the solution and the higher the temperature; the presence of an acid, e.g. H2SO4, greatly quickens the rate of change (A., l.c.). Addition of NaOHAq to a boiling solution of the Na salt produces Na₂HPO₈ (A., C. R. 106, 1400). Pyrophosphites in very dilute solutions give no pp. with Pb(NO₈)₂Aq until boiled; phosphites give an immediate white pp.; this reaction serves as a qualitative test for the two classes of salts (A., C. R. 110, 901).

It is insol. water, but is gradually changed by water to H₂PO₈Aq and PbHPO₃ (A., C. R. 110, 901)

Sodium pyrophosphite Na₂H₁P₂O₅ is formed by heating 2NaH₂PO₅.5H₂O, which has been partially dehydrated in vacuo, to c. 160°. V. sol. water; solution changes to NaH₂PO₅Aq, but by careful evaporation in vacuo over H₂SO₄ crystals of Na₂H₂P₂O₅ can be again obtained (A., C. R. 108, 1056). Presence of acid hastens the change to NaH₂PO₅Aq; this change is also effected by adding NaOHAq to boiling Na₂H₂P₂O₅Aq. Heat of solution of Na₂H₂P₂O₅ = 300 (A., C. R. 110, 191).

For thiopyrophosphites M₄P₂S₃ and thio-oxypyrophosphites M₄P₂O₄S₂ and M₄P₂O₄S v. Phos-Phobous Sulphide, Reactions 7 and 8, p. 146; and for selenopyrophosphites M₄P₂Se₃ v. Phosphonous selenide, p. 145). M. M. P. M.

PHOSPHORYL COMPOUNDS. Compounds of the radicle PO are generally called phosphoryl compounds; and compounds of the radicle PS are often called thio-phosphoryl compounds. These compounds are described in this Dictionary as phosphorus oxychloride, oxyftuoride, sulphochloride, &c. The phosphoryl compounds are PO(NH₂)₃, POCl₃, POBr₃, POBrCl₂ and POBr₂Cl, POF₃, PO.NH.NH., PON. The thiophosphoryl compounds are PS(NH₂)₃, PSCl₃, PSBr₃, PSBrCl₂, PSF₃. The compounds P₂O₃Cl₄ and P₂S₃Br₄ are sometimes called pyrophosphoryl chloride and pyrothiophosphoryl bromide, respectively.

M. M. P. M.

(a)-PHOSPHOTOLUIC ACID

C_eH₂Me(PO₂H₂).CO₂H [3:4:1]. Toluphosphinic acid. [262°]. Formed by oxidising m-xyleno phosphonic acid with hot alkaline KMnO₄ (Weller, B. 20. 1723: 21. 1492). Prisms, v. gol.

(Weller, B. 20, 1723; 21, 1492). Prisms, v. sol. hot water, v. e. sol. alcohol. Yields HPO, and m-toluic acid when heated.—PCl, yields oily C,H,Me(POCL).COCl. — PbHA'' aq: needles

(from dilute HOAc).—Ag₃A'''.

(g)-Phosphotoluic acid
C_sH_sMe(PO_sH_s).CO_sH [5:3:1]. [220°]. Formed
by oxidising s-m-xylene phosphonic acid (Weller). Yields HPO_s and m-toluic acid on heating.
—Ag_sA''': amorphous pp.

Chloride C.H.Me(POCl2).COCl. (249° at

147 mm.). Oil.

Phosphotoluic acid C_sH_sMe(PO_sH_s).CO_sH[4:x:1]. [278°]. Formed by oxidising p-xylene phosphonic acid. Needles, decomposed by heat into ITPO_s and p-toluic acid.

Chloride C_eH_sMe(POCl₂).COCl. [62°]. Crystalline mass.

PHOTO-CHEMISTRY v. Influence of light on chemical change, under Chemical Change, vol. i. p. 748; also next article (Photographio Chemistry); also Optical methods, a section of Physical methods, in this volume, p. 221.

PHOTOGRAPHIC CHEMISTRY. The photographic processes at present in use depend primarily upon the photo-chemical decomposition of certain metallic compounds in the presence of suitable oxidisable substances known as sensitisers. Broadly speaking, the photodecomposable compound may be regarded from a chemical point of view as an oxidising agent, of which the oxidising power is only brought

into action under the stimulus of light, the associated compound or sensitiser being at the same time oxidised by the oxygen or halogen thus liberated. Very many metallic compounds are thus susceptible to the influence of light, although only a few of these have up to the present time found practical application. The nature of the chemical change undergone in such cases may be well illustrated by the action of light upon ferric salts. An aqueous solution of ferric chloride, for example, is not acted upon by light because the associated substance (water) is not capable of reacting with the liberated chlorine with sufficient rapidity; in other words, water is incapable of acting as a sensitiser towards ferric chloride. But if some oxidisable organic compound is present, light then reduces the ferric to the ferrous salt. Thus an alcoholic solution of ferric chloride exposed for a few minutes to sunlight gives a blue colouration with potassium ferricyanide, indicating the presence of a ferrous salt, probably formed according to the equation: Fe $Cl_8 + C_2H_8O = Fe_2Cl_4 + C_2H_4O + 2HCl$. In a similar way paper coated with a solution of ferric chloride can be used to obtain prints, because the paper (or the size contained in it) plays the part of a sensitiser, and the exposed portions of the surface thus become covered with a ferrous instead of a ferric compound, and on brushing over with a solution of ferricyanide the print is developed in blue. In practice ferric oxalate is found to be the most sensitive ferric compound, the organic acid being the sensitiser: Fe₂(C₂O₄),

= $Fe_2(C_2O_1)_2 + 2CO_2$.

Although the chemical action of light upon metallic compounds generally is of the nature of reduction, and is accompanied by the simultaneous oxidation of the compound which plays the part of a sensitiser, there are many instances in which light promotes the action of atmospheric oxygen or other oxidising agent. Thus potassium iodide may be kept in a dry atmosphere in the dark for any length of time without undergoing change, but in the presence of light and moisture photo-chemical oxidation takes place,

and the salt becomes alkaline:

 $4KI + 2H_2O + O_2 = 4KOH + 2I_2$ Action of light upon silver salts. Of all known metallic compounds the salts of silver are the most sensitive to the action of light, and these salts accordingly form the basis of all existing photographic processes. The most familiar instance of the photo-decomposition of a silver salt is the well-known change in colour experienced by silver chloride on exposure to light, a phenomenon which has been familiar since the sixteenth century. The other silver haloids, viz. the bromide and iodide, are also decomposed by light, the latter only in the presence of some sensitiser capable of rapidly absorbing iodine. The absolute sensitiveness of the silver haloids cannot be estimated by the degree of darkening on exposure, because the product of photo-decomposition is of a darker colour in the case of the chloride than in the case of the bromide, and the product is darker in the case of the bromide than in that of the iodide. The sensitiveness is actually determined by two conditions, viz. the nature of the sensitiser and the state of molecular aggregation of the silver haloid. In most of the negative processes now in use, the bromide or iodide, or a mixture of the two. forms the sensitive surface.

The nature of the chemical change undergone on exposure to light has not yet been completely elucidated for all the silver haloids. It is known that the chloride loses chlorine on exposure, and that the darkened product contains less chlorine than the unaltered haloid. It is as yet uncertain whether the halogen which is given off is always evolved in the free state or not; it has been shown, however, that silver chloride on exposure to light gives off a gas which rapidly turns potassium iodide and starch paper blue (Meldola, Chemistry of Photography, p. 66). It is stated by many writers that the darkened product is a subchloride formed according to the equation: $4AgCl = 2Ag_2Cl + Cl_2$. There is, however, no satisfactory evidence of the existence of such a subsalt, and it seems more probable that the dark product consists of an oxychloride, in combination, or admixture, with unaltered chloride. According to some analyses by Hodgkinson, the oxychloride has the formula Ag, OCl2. In support of this conclusion it is found that the chloride does not darken in absolutely dry air, unless vapour of mercury is present (Abney), thus making it appear that moisture is essential for the decomposition. The latest investigation of the action of light on AgCl by Baker (C. J. 61, 728) confirms the view that the darkened product contains an oxychloride; Baker gives the formula Ag₂ClO. A second, white, oxychloride also probably exists (Richardson, C. J. 59, 536). Silver chloride is thus analogous to cuprous or thallous chloride, which also darken under the influence of light, with the formation of what are believed to be oxychlorides. Whatever may be the chemical composition of the darkened product, it is probable that the bromide and iodide give rise to the formation of analogous compounds, and that the invisible picture in the older processes is formed of these products on The una background of unaltered haloid. developed image on modern dry plates, coated with gelatin emulsion, may, however, have a different composition, owing to the presence of the organic sensitiser.

These are coloured Photosalts of silver. forms of the silver haloids prepared by the action of certain reducing agents upon silver compounds, and the subsequent conversion of the partially reduced products into the haloids by treatment These compounds with the necessary acids. are of a red or purple colour, and have been shown to contain less halogen than the un-altered haloids. It is probable that they consist of physical combinations of the oxyhaloid or hydrated oxyhaloid with the unaltered haloid; their discoverer (Carey Lea) has shown that they closely resemble, if they are not actually identical with, the products of the photo-chemical reduction of the silver haloids. If this be admitted, it follows that the product which forms the photographic image can be prepared

by purely chemical methods (Am. S. 1887).

Photographic processes. In all photographic processes the first operation is the exposure of a surface coated with a uniform film of the silver haloid to the image formed by a lens in a camera. The image thus depicted on the film is invisible, in the first place because of the very

slight difference in colour between the product of photo-reduction and the unaltered haloid, and in the next place because the short period of exposure does not produce a sufficient quantity of the product of reduction to render the latter visible. The invisible image thus consists of a layer of reduction-product of infinitesimal thinness, and the picture is afterwards made visible by the deposition of metallic silver on this reduction-product by the application of certain solutions known as 'developers,' the chemical action of which will be described subsequently.

Daguerreotype. This process is historically interesting as having been the first method of photography with a silver salt successfully applied to the production of a picture from the image formed in a camera. It takes its name from Daguerre, who announced the discovery in 1839. The process is no longer used; it has been superseded by more rapid methods. order to prepare a Daguerreotype plate, a plate of copper is silvered, and the polished surface is exposed to the vapour of iodine and bromine alternately. The sensitive film is therefore a mixture of silver iodide and bromide. Silver iodide can act to some extent as its own sensitiser, but the substratum of metallic silver is chiefly effective in this capacity as it is capable of rapidly absorbing the halogen liberated by the action of light upon the sensitive surface. The invisible image is developed by exposure to the vapour of mercury, which condenses on the product of reduction but not on the unaltered haloid.

Collodion processes. In these processes the sensitive film consists of a silver haloid formed by double decomposition in collodion as a vehicle. Collodion is a solution of the tri- and tetranitrate of cellulose (collodion pyroxyline) in alcohol and ether (v. vol. i. p. 716), and on coating a glass plate with this solution the solvents evaporate and leave a uniform layer of the pyroxyline attached to the plate. Some soluble iodide, NH, I or CdI, sometimes mixed with a small quantity of a bromide, is first dissolved in the collodion, the plate is coated with this salted collodion, and when dry is sensitised by immersion in a silver nitrate solution. sensitive surface in this process is therefore a silver haloid wetted with a film of silver nitrate solution, the latter playing the part of a sensitiser by absorbing the liberated halogen: $3I_2 + 6AgNO_3 + 3H_2O = 5AgI + AgIO_3 + 6HNO_3$. By washing out the soluble salts after the plate has been removed from the silver solution the sensitiveness is practically destroyed, because the pyroxyline by itself has no halogen-absorbing power. On drying such a washed plate, and then coating it with a solution of tannin or some organic sensitiser, the sensitiveness is partially restored. Plates thus treated can be used in the dry state, and dry-plate photography by this method was made practically successful in 1861 by Colonel Russell.

Emulsion processes. All the photographic methods at present in vogue are comprised in this group. The silver haloid is precipitated in a finely divided state in some vehicle, usually gelatin, and the plates coated with this emulsion are allowed to dry and are then ready for use. A brief description of a method for preparing a gelatin emulsion will suffice to make

clear the general mode of procedure. The soluble bromide (potassium or ammonium), sometimes mixed with a little iodide, is dissolved in water, together with a certain quantity of the gelatin, and to this solution the calculated quantity of silver nitrate (previously dissolved in water) is added little by little with constant agitation. The quantities are so adjusted as to leave no excess of silver nitrate. The silver haloid is by this means precipitated in an extremely fine state of division in the viscid gelatin solution, with which it forms an emulsion. A stronger solution of gelatin is then mixed intimately with the first dilute emulsion, and the whole is heated on a water-bath for about an hour to become 'ripened.' When cold, the semi-solid emulsion is washed by immersion in a fine state of division in cold water, so as to remove all soluble salts (nitrates formed by double decomposition, excess of soluble haloids, &c.). The washed emulsion is then mixed with the necessary quantity of strong gelatin solution to reduce it to the required consistency, and the plates are coated and allowed to dry. Many variations in the details of mixing have been introduced, but these involve no change in principle. Ripening can also be effected at ordinary temperatures by the action of ammonia.

Ripening of emulsions. The process of ripening above referred to is necessary in order to increase the sensitiveness of the silver haloid, since the latter when first precipitated is comparatively insensitive. The increase of sensitiveness is no doubt due, partially if not wholly, to a physical change in the state of molecular aggregation of the silver haloid, as it is accompanied by a growth in the size of the particles and by a change in the absorption spectrum, the unripened emulsion transmitting more of the red rays, and, therefore, absorbing less of this part of the spectrum, than the ripened emulsion. The extreme sensitiveness of the emulsions at present in use is thus due in great measure to the circumstance that the haloid is rendered sensitive to a much wider range of spectral colours than is tho case with ordinary silver bromide precipitated in a non-emulsifying medium. The increased sensitiveness of a ripened emulsion may also be ascribed, in part, to the formation of an actual compound of the silver haloid with the gelatin or some constituent thereof (Meldola, Cantor Lectures on Photographic Chemistry, 1891. 20,

Development of the photographic image. Acid development. In the wet collodion process the image is developed by the application of a solution of ferrous sulphate mixed with acetic acid and alcohol (to insure uniformity of flow). The film being already wet with silver nitrate solution, the action of the developer is simply to reduce this salt according to the reaction: $6AgNO_3 + 6FeSO_4 = 2Fe_2(SO_4)_3 + Fe_2(NO_3)_6 + 6Ag$. Metallic silver is thus contained as a potential deposit in the mixed solutions, but its precipitation is retarded by the acetic acid, which plays the part of what is technically called a 'restrainer.' The finely-divided metal is deposited only on the product of photo-decomposition (the invisible picture), the density of the deposit being proportional to the amount of decompo-

sition on each portion of the sensitive surface. The image thus continues to gain in density as long as there is silver to feed it, so that the picture is built up by accretion and is raised in relief on the surface of the film. If the film is treated with dilute nitric acid after development, the picture is dissolved off, leaving the film in the same condition as before exposure, thus proving that the image is purely superficial. The ferrous sulphate developer, which acts in the manner described, is a type of the class of acid developers.

Alkaline development. Gelatin emulsions contain no excess of silver nitrate, so that the foregoing process of development is inapplicable. The gelatin itself, being a bromine absorbent, is in this case the sensitiser. Development is effected in modern processes by means of an alkaline or neutral solution of some reducing agent, usually an easily oxidisable organic compound, such as ammonium pyrogallate, first introduced by Colonel Russell in 1862. In this method the developer acts directly as a reducing agent upon the product of photo-reduction, replacing the latter by its equivalent of metallic silver. The image at the same time gains in density by the further reduction of those portions of the unaltered silver haloid which are in immediate contact with the nascent silver being generated by the developer; the action is probably electrolytic, since the image goes on increasing in density as long as the developer can exert a reducing action. A very small amount of the silver deposit may also be due to the reduction of the minute quantity of silver actually dissolved out of the film by the ammonia or other solvent in the developer. In alkaline development the main portion of the silver image is therefore built up by growth from the silver haloid in the film, and after development the image can be dissolved out by dilute nitric acid, leaving its impression sunk in the gelatin, instead of leaving a plane surface, as in the case of a collodion picture similarly treated.1 The developers belonging to this class are all strong reducing agents capable of directly reducing the silver haloids, so that the action has to be moderated by means of restrainers, potassium bromide being generally employed for this purpose. The restraining action of this salt is probably due to its tendency to form a double potassio-silver bromide, which is more stable than the silver bromide itself. In addition to ammonium pyrogallate several other developers acting in a similar manner have come into use. Of these may be mentioned potassio-ferrous oxalate (Carey Lea, 1877), which acts according to the equation: = $3Fe_2(C_2O_4)_3 + 6KBr$. $3Br_2 + 6FeC_2O_4 + 3K_2C_2O_4$ The bromine in this case is derived primarily from the reduction-product. Other developers coming under this category are an alkaline solution of hydroquinone (quinol). hydroxylamine, phenylhydrazine, and an alkaline salt of amido-\$-naphthol-monosulphonic acid (Meldola, C. J. 39, 47), introduced by Andresen (Eikonogen). Among the most recently introduced developers are certain amido-phenols.

¹ An able investigation of the connection between the density of the deposit of reduced silver and the period of exposure, &c., is published by Harter a. Driffield (S. C. L. 2, 455; 10, 100; v. also other papers in same volume; also Armstrong, Conference No. of Journ. of the Camera Club, 1892, and discussion in same journal, July 1892).

amido-phenolic and carboxylic acids, and their alkyl derivatives, known as 'amidol,' 'metol,' &c. Those which contain the basic and hydroxyl substituents in the para-position seem to be the most effective (Hauff, Engl. Patents [1891] 15, 431; 20, 690).

In the processes of alkaline development the gelatin itself also acts the part of a restrainer by preventing too intimate a contact between the reducing solution and the silver haloid. Any viscous substance exerts this protecting influence (sugar, glycerin, &c.), and such compounds are sometimes called physical restrainers, to distinguish them from chemical restrainers, such as the acid in the ferrous sulphate developer or the potassium bromide in alkaline developers. A developer of very great reducing power is for this reason available in dry plate photography, and upon this fact largely depends the extreme sensitiveness of modern processes.

Development by vapour. The Daguerreotype picture was developed by exposure to mercury vapour, this metal having the property of condensing on the product of photo-reduction (the invisible image) but not on the unaltered silver haloid. It is not known whether the combination of the mercury with the reduction-product is of a chemical or physical nature. This method of development is at present the sole

representative of its class.

Fixing the picture. The image developed by the foregoing methods is always on a back ground of unchanged silver haloid, which must obviously be removed before the picture can be exposed to white light. In the early days of photography strong solutions of potassium, sodium, or ammonium chloride were used as fixing agents, these salts having the property of forming soluble double salts with the silver haloids. The action of these salts was, however, very imperfect, and a certain amount of undissolved silver haloid was always left in the film, which led on exposure to the gradual darkening and obliteration of the whole picture. Potassium cyanide has since been employed, this salt forming with the silver haloid an extremely soluble double cyanide: AgBr+2KCN =AgK(CN)₂+KBr. The fixing agent now generally preferred is sodium thiosulphate (Sir J. Herschel, 1839); this salt, if used in excess, forming the extremely soluble sodio-silver thiosulphate $Ag_2Na_4(S_2O_3)_3$, $2aq: 2AgBr + 3Na_2S_2O_3 = Ag_2Na_4(S_2O_3)_3 + 2NaBr$. If the fixing solution is dilute, the insoluble $NaAgS_2O_3$ is formed, which remains in the film and spoils the picture: $AgBr + Na_2S_2O_3 = AgNaS_2O_3 + NaBr$. After immersion in the fixing bath, the soluble salts are removed by thorough washing in a stream

Intensification, and reduction, of density. In cases where the image is wanting in density through under-exposure, bad light, insufficient development, &c., a process of intensification is resorted to. The process as applied to modern dry plates consists in substituting for the silver, of which the image is composed, some denser deposit. Thus, by immersing a negative in a solution of mercuric chloride, the image is bleached by conversion into a mixture of mercurous and silver chlorides: 2Ag+2HgCl₂=2AgCl+Hg₂Cl₂. By treating this mixture

with a solution of potassio-ferrous exalate the image is restored in silver and mercury, and thus 'in a condition of increased $2 \operatorname{AgCl} + \operatorname{Hg}_2 \operatorname{Cl}_2 + 4 \operatorname{FeC}_2 \operatorname{O}_4 + 2 \operatorname{K}_2 \operatorname{C}_2 \operatorname{O}_4$ density: = $2Ag + 2Hg + 2Fe_2(C_2O_4)_3 + 4KCl$. Many other intensifiers are known, some acting by simple substitution, such as auric or platinic chloride: $6Ag + 2AuCl_s = 2Au + 6AgCl$, or $4Ag + PtCl_s$ =Pt + 4AgCl; others giving rise to a mixture of products together denser than the original deposit, e.g. lead or uranium ferricyanide: $4Ag + 2Pb_3Fe_2(CN)_{12} = Ag_4Fe(CN)_0 + 3Pb_2Fe(CN)_0$ In cases where too dense an image has been obtained, and it is desired to thin it down, a reducing solution is applied; the chemical principle may be described as a conversion of the metallic silver into some compound which can be simultaneously removed by a solvent such as sodium thiosulphate. Thus a mixture of potassium ferricyanide with thiosulphate is often used for gelatino-bromide plates: $4Ag + 2K_{6}Fe_{2}(CN)_{12} = Ag_{4}Fe(CN)_{6} + 3K_{4}Fe(CN)_{6}$ The silver ferro-cyanide is dissolved off by the thiosulphate, as fast as formed: $Ag_{+}Fe(CN)_{6} + 6Na_{2}S_{2}O_{+}$ = $2Ag_2Na_4(S_2O_3)_3 + Na_4Fe(CN)_6$. When the necessary reduction of density has been attained, the action of the solution is stopped by washing the film with water.

Reversal of the photographic image and allied phenomena. Any influence which affects a sensitive film in a way similar to the action of light gives rise to a developable image. Thus mechanical pressure-marks produced by rubbing a film with a glass rod can be developed by ferrous oxalate &c., so as to show a corresponding set of dark streaks. It is probable that the friction in thiscase induces a minute amount of chemical change between the sensitive haloid and its associated: sensitiser (gelatin, &c.), this small quantity of reduction-product being afterwards exaggerated by the cumulative action of the developer. Whether the compound resulting from this mechanicalaction is identical with that produced by the action of light is at present uncertain, but the associated sensitiser appears to be as essential tothe success of the phenomenon in this case as in the ordinary formation of the invisible picture by the action of light. The sensitiser, as has already been explained, is an essential part of any photographic system, and this is best shown by the so-called 'reversal' of the photographic image by over-exposure or other means. It has been found that a very strong light allowed to act on a sensitive film for too long a period gives a. weaker image on development than is produced with a shorter exposure, and that under some circumstances the image is even 'reversed,' i.e. the high lights come out lighter, on development. than the shadows. The explanation of this phenomenon is probably to be found in the circumstance that while the silver haloid is losing halogen under the influence of light, the sensitiser is simultaneously becoming halogenised, so that when the charge of halogen in the latter exceeds a certain amount the chemical change sets. in in a reversed direction, i.e. the halogenised sensitiser begins to re-halogenise the product of photo-decomposition and thus to destroy the invisible image. That this explanation is, in the main, correct in principle, is shown by the fact that the most sensitive processes are just those which-

are most prone to give rise to reversal, because it is in such films that the sensitiser becomes most rapidly halogenised. A preliminary exposure to diffused light also promotes reversal, because this imparts a certain initial charge of halogen to the sensitiser. Then, again, oxidising agents favour reversal, because these lessen the halogen-absorbing power of the sensitiser; on the other hand, reducing agents prevent reversal, because they retard or prevent the halogenation of the sensitiser. Thus plates exposed in ozone or in solutions of K2Cr2O7 and K2Mn2O8 are easily reversed, while nitrites, sulphites, &c., prevent reversal (Abney). Atmospheric oxygen in many cases probably plays a part in the process by assisting in the liberation of halogen from the sensitiser. Thus, by way of illustration, if a plate be given a preliminary exposure to diffused light, and be then coated with a solution of potassium iodide, a completely reversed image can be obtained on exposure in the camera and development. The chemical reaction is in this case: $4KI + 2H_2O + O_2 = 2I_2 + 4KOH$. The iodine is here absorbed by the reduction-product, and thus reverses the picture; the surface of the film is in fact the sensitiser in this case, and the potassium iodide is the photo-sensitive com-

Waterhouse has recently found that a small quantity of a thio-carbamide added to the developer produces a 'reversed' image (Photo. News, 1890. 727, 743, 804; v. also Journ. of Camera Club, July 1892; also Rawlins, S. C. I.

10, 18).

Printing processes. It is obvious that the photographic image, after normal development, represents the object with its lights and shadows reversed, i.e. the picture is a negative one. positive print is obtained by exposing suitable sensitive surfaces under such a negative.

Silver prints. A very general method of obtaining silver prints depends upon the use of a sensitive surface (usually paper) consisting of a mixture of silver chloride and an organic compound of silver with albumen, this surface being obtained by first coating the substratum with a solution of albumen containing ammonium chloride, and then (when dry) floating on a solution of silver nitrate. The printing is carried on to the required depth of colour, and the visible print consists of the reduction-products of silver chloride, and of the organic silver compound, which is also susceptible of photo-reduction. To correct the reddish colour of these mixed reduction-products the print is 'toned' by immersion in a solution of auric chloride, kept neutral by the addition of sodium carbonate, acetate, or phosphate, chalk, borax, &c. The function of these salts is to neutralise the hydrogen chloride liberated by the reduction of the auric chloride by the reducing compounds which constitute the picture. The gold is thus precipitated in a state of fine division only on the products of photodecomposition (the picture), and imparts the desired tone. The accumulation of acid in the toning bath retards the precipitation of gold, and

¹ Potassium bromide similarly acts as a reversing agent shough to a less extent. It is for this reason that every trace of this sait has to be washed out of a gelatino-bromide emulsion after the process of ripening, since the presonce of the soluble bromide materially diminishes the sensitiveness of the emulsion.

the deposit is not of a good tone; hence the necessity for the presence of one of the salts referred to. The unaltered silver chloride &c. is removed by fixing with thiosulphate, and washing. In some recent processes the albumenised silver paper is replaced by paper coated with gelatino-bromide or chloride emulsion, a short exposure being given, and the invisible (positive) picture being developed in the usual manner.

Printing with iron and uranium salts. Ferric and uranic salts in the presence of organic sensitisers become reduced on exposure to light, and the ferrous or uranous compounds thus formed can be developed by treatment with potassium ferricyanide or other salts which give coloured products with the ferrous or uranous. but not with the unreduced, salts. Thus paper coated with ferric oxalate gives a faint image of the ferrous salt, which comes out as a deepblue print on development with ferricyanide: $6\text{FeC}_2\text{O}_4 + 2\text{K}_6\text{Fe}_2\text{Cy}_{12} = 2\text{Fe}_3(\text{Fe}_2\text{Cy}_{12}) + 6\text{K}_2\text{C}_2\text{O}_4.$ Many printing processes depending upon the foregoing principles have come into use, and will be found described in works on practical photography.

Platinotype. Prints in finely-divided platinum can be obtained by exposing a surface coated with a mixture of ferric oxalate and potassium chloroplatinite under a negative. The ferrous salt thus produced does not react with the chloroplatinite till the picture is developed by immersion in a solution of potassium oxalate, which dissolves the ferrous oxalate, with the formation of a double salt, which simultaneously reduces the chloroplatinite:

 $3K_2PtCl_4 + 6FeC_4O_4$ = 3Pt + 2Fe₂(C₂O₄)₃ + Fe₂Cl₈ + 6KCl. The soluble salts are then removed by washing with dilute hydrochloric acid, and finally with water.

Pigment printing. A mixture of gelatin with potassium dichromate undergoes a chemical change on exposure to light, in the course of which the dichromate is reduced and the gelatin at the same time becomes insoluble in water. The chemical composition of this insoluble gelatin is unknown, but it appears to contain an oxide of chromium as an essential constituent (Eder). Many printing processes are based upon this property of gelatin, such, for example, as the so-called carbon and pigment prints, in which the finely divided carbon or pigment is intimately mixed with the gelatin solution and the mixture sensitised by the addition of dichromate. After exposure under a negative the picture is developed by warm water, which dissolves away those portions of the tissue unacted upon by light. Many of the photo-mechanical printing processes de-

pend also on this property of gelatin.

Photo-etching processes. When a layer of asphalt or bitumen is spread over a surface and exposed under a design, those portions of the film which are acted on by light become insoluble in hydrocarbon oils, so that the design can be developed by such solvents, and the surface, if of metal, can be converted into a printing block by etching with acid. The change experienced by the bitumen is probably the result of photo-chemical oxidation. The processes based on this property are much in vogue at the present time under various modifications. This action of

light upon bitumen furnished the earliest successful permanent reproduction of the camera picture (Joseph Nicephore Niepce, 1824).

Bibliography.—It has not been considered advisable in the present article to give the authority for every statement, as most of the results achieved by photographers are to be found in publications rarely consulted by chemists. The following general works contain practically all that is known of photographic chemistry at the present time : -Robert Hunt, Researches on Light, 2nd edit., 1854; E. Becquerel, La Lumière, Paris, 1867; J. W. Draper, Scientific Memoirs, a collection of reprints, 1878; J. M. Eder, Ueber die Reactionen der Chromsäure u. der Chromate auf Gelatine, Gummi, Zucker, &c. Wien, 1878; J. M. Eder, Ausführliches Handbuch der Photographie, Halle, 1884-9; H. W. Vogel, La Photographie des Objets Colorés avec leurs Valeurs Réelles, Paris, 1887; Pizzighelli and Hübl, La Platinotypie, Paris, 1887; W. de W. Abney, Treatise on Photography, 5th edit., 1888; by the same author, Instructions in Photography, and Photography with Emulsions; Chapman Jones, Introduction to the Science and Practice of Photography, 1888; R. Meldola, The Chemistry of Photography, 1889; H. W. Vogel, Handbuch der Photographie, Berlin, 1890-1. R. M.

PHOTOSANTONIN v. SANTONIN.

PHRENOSIN C₄₁H₇₉NO₅. A substance occurring, according to Thudichum (*J. pr.* [2] 25,

19), in the brain.

PHTHALACENE C₂₁H₁₆. [173°]. Formed by reduction of phthalacone carboxylic ether with HI and P at 170° (Gabriel, B. 17, 1390). Crystals (from HOAc). Yields C₂₁H₁₅Br [184°], which is oxidised by K₂Cr₂O, and HOAc to C_nH₁₈BrO [c. 200°]. Fuming HNO_s and HOAc yield di-nitro-phthalacene, which separates from hot nitro-benzene in yellow needles. K2Cr2O, and HOAc oxidise phthalacene to 'phthalacene oxide' $C_{21}H_{1,0}$ [211°-214°], which yields an oxim $C_{21}H_{1,1}(NOH)$ [266°]. PHTHALACENIC ACID $C_{22}H_{18}$.CO₂H. [247°].

Formed by heating phthalacene-oxide with sodalime (Gabriel, B. 17, 1399). Crystals.—A'Ag.

CARBOXYLIC PHTHALACONE ACID OnH11O2(CO2H). [281°]. Formed by dissolving its ether in conc. H₂SO, and pouring into water (Gabriel, B. 17, 1389). Minute yellow needles, sol. hot alcohol. The acid yields a dioxim C₁H₁₁(NOH)₂CO₂H [273°] and the salts KA'aq and NaA'aq. Zinc-dust and NaOHAq yield a tetrahydride C₂₂H₁₈O₄, melting above 280°, which gives AgA', crystallising in needles.

Ethyl ether EtA'. [211°]. A product of the action of phthalic anhydride and NaOAc on acetoacetic ether at 140°. Yellow needles, sol. hot HOAc. Yields a di-nitro- derivative [above 280°] and a dioxim C₂₁H₁₁(NOH)₂CO₂Et [264°]. Zinc-dust and HOAc reduce it to the white

tetrahydride C₂₂H₁₂(OH)₂CO₂Et [218°]. PHTHAL-ALCOHOL v. DI-OXY-XYLENE. PHTHALALDEHYDIC ACID C₂H₂O₃ i.e. CHO.C.H..CO.H.[1:2]. [97°]. Formed, together with C_1 , H_{10} , [221°], by heating a-bromophthalide with water (Racine, B, 19, 778; A. 239, 78; C. R. 106, 947). Formed also by boiling penta-ω-chloro-o-xylene with water (Colson a. Gautier, Bl. [2] 45, 509). V. sol. water, alcohol, and ether. Reduces ammoniacal AgNO.

Phenyl-hydrazine yields C₁₄H₁₀N₂O [105°]. Alco holic NH_s forms C₂₄H₁₈N₂O₆ [187°], while an alcoholic solution of aniline gives C_{1,H1,NO}, [174°]. Ac₂O at 200° forms CHO.C₆H₁,CO₂Ac [240°]. Ures forms NH₂, CO.N:CH.C.H., CO.₂H [240°].—CaA'₂ 2aq.—AgA': slender needles. Ethers MeA'. [44°].—EtA'. [66°]. Anhydride O.₂H., O.₃. [221°]. Got by

heating the acid with bromo-phthalide.

[120°]. OximCH(NOH).C.H..CO.H. Formed by adding hydroxylamine hydrochloride to a cold aqueous solution of the acid. In an alcoholic solution the product is C₆H₄ CO. O [120°], which is converted by heat first into CNC,H,CO,H and then into phthalimide (Allendorff, B. 24, 2346, 3264).

Isomeride v. p-Aldehydo-Benzoic acid. Diphthalaldehydic acid v. DI-PHTHALYL-LAC-TONIC ACID.

PHTHALAMIC ACID v. PHTHALIC ACID. PHTHALAMIDE v. PHTHALIC ACID.

ISOPHTHALAMIDINE C.H. N. i.e. $C_8H_4(C(NH).NH_2)_2$ [1:3]. Formed from C.H. (C(NH).OEt), and alcoholic NH, (Luckenbach, B. 17, 1432). Small needles, insol. benzene and ether, v. sol. alcohol and water. Its aqueous solution soon decomposes, giving off NH3.-B"H_Cl_: needles, v. sol. water.—B"H_PtCl_.

C.H.(C(NOH).NH.)₂. [193°]. Formed from C.H.Cy₂ [1:8] and hydroxylamine (Goldberg, B. 22, 2976). Prisms (from alcohol) containing aq?, v. sol. hot water.

Phthalanil v. Phenylimide of Phthalic ACID. PHTHALBENZO-TOLUIDE v. Phthalyl-AMIDO-TOLYL PHENYL KETONE

PHTHAL-o-CYANO-BENZYL-IMIDE o-Cyano-benzyl-phthalimide.

PHTHALEÏNS. Colouring-matters obtained by condensation of phthalic anhydride with phenols (e.g. Phenol-Phthalein and Fluor-escein). They may be reduced to colourless 'phthalins,' which are re-oxidised by air to phthaleins.

PHTHALIC ACID C,H,O, i.e. [1:2] C.H.(CO.H). Mol. w. 166. [184°] (Lossen, A. 144, 76); [203°] (Ador, A. 164, 230; Baeyer, A. 269, 184). S. 54 at 14°; 18 at 99° (Graebe, A. 238, 321); S. (alcohol) 10 at 15° (Bourgoin, Bl. [2] 29, 247); S. (ether) 684 at 15°. H.C. 771,600. H.F. 187,400 (Stohmann, J. pr. [2] 43, 540); 158,000 (von Rechenberg). S.H. (from

75° to 119°) ·256 (Hess, A. Ch. [2] 35, 410).

Formation.—1. By the action of nitric acid on naphthalene, naphthalene dichloride, alizarin, purpurin, munjistin, naphthoquinone, o-toluic acid (Laurent, A. Ch. [2] 61, 113; Marignac, A. 42, 215; Schunck, A.66, 197; Wolff a. Strecker, A. 75, 12, 25; Stenhouse, A. 130, 334; Liebermann a. Dittler, B. 6, 945; Piccard, B. 12, 579; Beilstein a. Kurbatoff, A. 202, 215).—2. By the action of various oxidising agents on naphthalene (Lossen, A. 144, 71; Hermann, Z. [2] 4. 551; Depouilly, C. R. 56, 82; Häussermann, D. P. J. 223, 810).—3. A product of the action of MnO₂ and H₂SO₄ on benzene (Carius, Z. [2] 4, 705; A. 148, 60).—4. By oxidising o-toluic acid with KMnO₄ (Weith, B. 7, 1057).—5. By oxidising isoquinoline with alkaline KMnO. (Hoogewerff a. Van Dorp, R. T. C. 4, 285).-6. By heating salicylic acid with H2SO4 and K.FeCy. (Guyard, Bl. [2] 29, 247).—7. By heating resorein or salicylic acid with formic acid and H₂SO₄ (G.).—8. By the action of a hot solution of cuprous potassium cyanide upon o-diazobenzoic chloride (from anthranilic acid), and subsequent saponification of the product (Sandmeyer, B. 18, 1499).

Properties.—Trimetric plates. Yields an anhydride when heated. Insol. chloroform.

Reactions .- 1. Distillation with lime yields benzoic acid and benzene. - 2. Sodium-amalgam reduces it to a dihydride.—3. Chromic acid mixture oxidises it to CO₂ (Fittig, A. 156, 242).-4. The acid aniline salt A"H(NH,Ph) of orthophthalic acid loses H2O when its aqueous solution is boiled, phenyl-phthalimide crystallising out. The para- and meta- phthalic acids do not react in this way; the reaction therefore serves as a means of separating ordinary phthalic acid from its isomers (Michael a. Palmer, B. 19, 1376; Am. 9, 202)

Salts.—(NH4)HA": prisms, v. sol. water. NaHA" 2aq: prisms (Wislicenus, A. 242, 89).-Na₂A": pearly plates. — K₂A". — CaA" aq.—BaH₂A"₂: prisms (Hermann, A. 151, 78).— BaA": v. sl. sol. water (Carius, A. 148, 64).— Ba,A"aO: monoclinic prisms.—CuA"aq.—PbA". -Ag₄A": crystalline pp. — Aniline salt: needles [146°] (Clarke, B. 12, 1066).

Di-methyl ether Me₂A" (280° i.V.) at 734

mm. (Graebe, B. 16, 860). H.F. 164,600 (Stoh-

mann, J. pr. [2] 40, 353).

Mono-ethyl ether EtHA". Formed by heating phthalic anhydride with absolute alcohol at 100° (Michael, Am. 1, 413). Liquid, m. sol. water. Decomposed by heat into phthalic anhydride and alcohol.—Ba(EtA")2.—AgEtA".

Di-ethylether Et.A". (295° cor.). Got by passing HCl into an alcoholic solution of phthalic acid (Graebe a. Born, J. 1866, 411). Formed also from phthalyl chloride and alcohol. Liquid. Not attacked by hydroxylamine (Jeaurenaud, B. 22, 1273). When warmed with NaOEt and EtOAc it yields the compound $C_0H_4 < CO > CH.CO_2Et$ [75°-78°] (Wislicenus, A. 246, 349). PCl, followed by sodium malonic ether forms the acid C21H16O16 [c. 180°] (Zelinsky, B. 20, 1010).

Phenyl ether Ph2A". [70°]. Formed by heating phthalyl chloride with phenol (Schreder, B. 7, 704; von Gerichten, B. 13, 419). Colour-

less crystals.

Anhydride C,H,O, i.e. C,H, COOO. Mol. w. 148. V.D. 5.26 (calc. 5.13) (Troost, C. R. 89, 851). [128°]. (284.5° i.V.). H.C.p. 784,000. H.F. 106,000 (Stohmann, J. pr. [2] 40, 139). Formed by heating phthalic acid alone or with AcCl (Laurent; Anschütz, B. 10, 326). Formed also by the action of lead nitrate on phthalyl chloride (Lachovitch, B. 17, 1283). Long needles, v. sol. alcohol and ether, sl. sol. cold water. Reactions. - 1. When heated with phenols it yields phthaleins, with elimination of water (Baeyer, B. 7, 968).—2. Heated with NaOAc and HOAc it yields phthalyl-acetic acid C₁₀H₄O₄, while phenyl-acetic acid gives benzylidene-

phthalide. Phenyl-acetonitrile gives rise to C'H'<CO'ŏ -C:CPh.CN [165°], and isobutyric acid and ZnCl₂ at 250° give a ketone C₁₁H₁₀O₂ [96°]. 3. Boiling with NaOAc and phenoxyacetic acid yields C₆H₄:C₂O₂:CH.OPh [143°] while p-tolyloxy-acetic acid and sodium acetate give rise to C.H.: C.O.: CH.OC.H.Me [174°] (Gabriel, B. 14, 922).-4. Acetoacetic ether gives CeH, Bz, and phthalacone carboxylic ether.—5. On heating with succinic acid and sodium succinate it yields CO_2 and $C_{18}H_{10}O_4$ [above 300°].—6. Di methyl-quinoline and zinc chloride at 200° give $C_6H_4:C_2O_2:C_{11}H_6N$ [238°] (Beyer, J. pr. [2] 33, 407).-7. Ethenyl-amido-phenyl-mercaptan and $ZnCl_2$ at 200° give $C_eH_4 < N > C.CH < OO > C_eH_4$ orystallising in yellow needles [above 320°] (Jacobson, B. 21, 2630).—8. Cyanethine forms, on heating, C₆H₁₈N₂.N:C₂O₄:C₆H₄ [128°] (E. von Meyer, J. pr. [2] 39, 262).—9. Coal-tar picoline and zinc chloride at 200° form pyrophthalone C_sH₄O₂:CH.C_sH₄N [above 260°] (Jacobsen a. Reimer, B. 16, 2604) crystallising from alcohol in yellow plates.—10. Benzene and AlCl_s yield o-benzoyl-benzoic acid, and other aromatic hydrocarbons act in like manner (Friedel a. Crafts, C. R. 92, 833).-11. Benzyl chloride and zincdust at 75° yield $C_{50}H_{46}$ (?) [73°] (Wislicenus, A. 248, 68).-12. On heating with anvines and amides, phthalic anhydride yields derivatives of the imide and amide of phthalic acid; thus acetamide yields phthalimide and HOAc, while ethylamine yields ethyl-phthalimide. Secondary amines form derivatives of the amic acid and of the amide (Piutti, A. 227, 181; G. 16, 1, 251).— 13. Amido-acids are converted by phthalic anhydride into their phthalyl derivatives (Reese, A. 242, 1).—14. On fusion with sinc-dust phthalio anhydride gives diphthalyl. Zinc-dust and HOAc yield a mixture of phthalide, diphthalyl dihydride C.H.

CH, [199°], which is re-CO,H.C,H,.CH,.CH, duced by HIAq to C₂H₄(C₆H₄.CO₂H)₂ (Wislicenus, B. 17, 2178).—15. Tri-amido-phenol (picramic acid) forms (C₈H₄O₂:N)₈C₆H₂·OH [above 800°] converted by potash into the compound (CO₂H.C₆H.CO.NH)₂C₆H₂OH [above 300°], from which nitric soid produces the quinone (C₆H₄O₂:N)₂C₆H₂O₂ [277°] reduced by SO₂ to $C_0H_4O_2:N)_2C_0H_2(OH)_2$ [above 310°] (Piutti, G. 16, 254).

C,H, [229°] and

Chloride C_eH₁ COl₂>0. Phthalyl chloride. [0°] (Wischin, A. 143, 259). (275°) at 726 mm. S.G. $\frac{20}{4}$ 1.4089. $\mu_a = 1.569$ (Brühl, A. 285, 14). Formed by heating phthalic acid with PCl₅ for 2 hours at 190° (H. Müller, Z. 1863, 257; Graebe, A. 238, 320). Oil, absorbs moisture from the air, forming phthalic anhydride. Slowly converted into phthalic acid by water or Na₂CO₂Aq. Reduced by zinc and HClAq to phthalide. Sodium-amalgam and HOAc form C_eH₄(CH₂·OH)₄. PCl₅ forms two isomeric chlorides C_eH₄CCl₂>O and C_eH₄CCl₂, one melting at 88° and boiling at 275°, the other melting-

by aniline into C.H. CONPh) NPh [153°] (von Gerichten, B. 13, 417; Claus, B. 19, 1188). Excess of PCl, yields chloro-benzoic acid, CCl, and other products (Claus). Aqueous NH, forms $C_8H_4O_2(NH_2)_2$ [90°] not identical with phthalamide, but converted by HClAq into C₈H₄O₂(NH) [145°] which is changed by fusion into the isomeric phthalimide [228°] (Auger, A. Ch. [6] 22, 303; cf. Kuhara, Am. 3, 26). Dry NH, acting on the benzene solution behaves in like manner. Hydroxylamine yields C₆H₄:C₂O₂:NOH [230°] (v. vol. ii. p. 738). o-Amido-phenyl mercaptan hydrochloride forms $C_{20}H_{12}N_2S_2$ [112°] captan hydrochloride forms C_{20} L_{12} L_{22} L_{22} L_{23} L_{24} L_{25} L_{25} Lfollowed by water gives, in like manner, C12H14O2 [54°] (250°).

Semi-nitrile v. o-Cyano-benzoic acid.

Amic acid C_eH₄(CO.NH₂).CO₂H. Phthalamic acid. [c. 140°] (Auger); [149°] (Aschan, B. 19, 1401). Formed by heating phthalic anhydride with alcoholic NH, or phthalimide with baryta-water. Prepared by evaporating phthalic anhydride with NH,Aq, and decomposing the resulting ammonium salt by HCl (Auger, Bl. [2] 49, 349). Prisms, sl. sol. water, v. sol. alcohol. Converted by heat into phthalimide, and by hot water into hydrogen ammonium phthalate. Excess of HClAq forms phthalic acid.—NH, A'.-KA'.—BaA'₂.—BaA'₂ aq.—AgA': needles, v. sl. sol. water (Landsberg, A. 215, 197).

Isoamyl-amic acid O₄H₁(CO₂H).CO.NHC₃H₁₁. Isoamyl-phthalamic acid. [115]. Formed by warming isoamyl-phthalimide with KOHAq (Neumann, B. 23, 998). Crystalline pp.—AgA'.

Phenyl-amic acid C.H.(CO.H).CO.NHPh. (CO.H).CO.NHPh. Phenyl-phthalamic Phthalanilic acid. [192°]. Got by boiling phenyl-phthalimide with NH, Aq and some alcohol (Laurent a. Gerhardt, A. Ch. [3] 24,

188). Plates, sl. sol. cold water, v. sol. alcohol. Phenyl-ethyl-amic acid C_{1e}H_{1s}NO₅ i.e. C₆H₄(CO₂H).CONPhEt. Phenyl-ethyl-phthalamic acid. Got by dissolving phthalic anhydride in ethyl-aniline (Piutti, G. 13, 545; A. 227, 185). Heavy oil, v. sl. sol. water.

Di-phenyl-amic acid C₆H₄(CO₂H).CONPh₂. Di-phenyl-phthalamic acid. [148°]. Made from diphenylamine and phthalic anhydride (P.). Hard prisms, sl. sol. ether, v. sol. alcohol.—AgA'.

o-Tolyl-amic acid

 $C_{\mathfrak{g}}H_{\mathfrak{q}}(CO_{\mathfrak{p}}H).CONHC_{\mathfrak{g}}H_{\mathfrak{q}}Me.$ Got by boiling o-tolyl-phthalimide with NH, Aq (Kuhara, Am. 9, 51). Needles.—Ag₂C₁₅H₁₁NO₃: pp.

m-Methyl-benzyl-amic acid $C_{\bullet}H_{\bullet}(C_{\circ}H)$.CO.NHCH, $C_{\bullet}H_{\bullet}Me$. [13 [131°]. from m-xylyl-phthalimide and caustic soda (Brömme, B. 21, 2700). Needles, sol. alcohol.— AgA': white pp.

ψ-Cumyl-amic acid $C_bH_4(CO_2H).CO.NHC_gH_2Me_g.$ Phthal- ψ -cumidic acid. [179°]. Needles, v. sol. alcohol, sl. sol. ether (Fröhlich, B. 17, 1808).

Naphthyl-amic acids C_eH₄(CO₂H).CO.NHC₁₀H₁. The (a)-acid [185°] is got from (a)-naphthyl-phthalimide. The (β) -Vol. IV.

at 47° and boiling at 262°; both are converted | acid crystallises from alcohol in tables (Piutti. G. 15, 480).

Phenyl-oxy-ethyl-amic acid C₆H₄(CO₂H).CO.NH.C₂H₄.OPh. [125°]. Formed by warming phenyl-oxyethyl-phthal-imide with KOHAq (Schmidt, B. 22, 3255).

p-Tolyl-oxy-ethyl-amio acid C_eH₄(CO₂H).CO.NH.CH₂.CH₂.O.C_eH₄Me. [137°]. Formed from the corresponding imide which is got by heating potassium phthalimide with C₆H₄Me.OC₂H₄Br (Schreiber, B. 24, 191).

Phenyl amido-phenyl-amic acid C₀H₄(CO₂H).CO.NH.C₃H₄.NHPh. [120°-11] Powder (Gabriel, B. 22, 2223). [120°-130°].

Benzoylethyl-amic acid

C₈H₄(CO₂H).CO.NH.CH₂.CH₂Bz. Propiophenonephthalamic acid. [140°]. Made from the imide (Schmidt, B. 22, 3251). Needles.—AgA': pp.

Di-sulphido-di-ethyl-amic acid (C₄H₄(CO₂H).CO.NH.CH₂.CH₂)₂S₂. [130°]. Formed from sulphocyano-ethyl-phthalimide

Scales. HClAq at 180° forms S₂(C,H,NH₂).

Uramic acid C₆H₄(CO₂H).CONH.CO.NH₂.

Phthaluric acid. S. 35 at 99°. Formed by heating phthalic anhydride with urea at 115° (Piutti, A. 214, 19; G. 12, 173). Silvery scales, sol. alcohol. May be converted by POCl, into C₆H₄ CO.NH CO, which yields AgC₉H₅N₂O₃.

Salts.—NaA' 2aq.—BaA', —AgA': needles. Thio-uramic acid

C₆H₄(CO₂H).CO.NH.CSNH₂. [172°]. Got by heating phthalic anhydride with thio-urea (P.). Silvery needles (from alcohol).—BaA'27aq.

s-Amide C.H.(CO.NH₂)₂. [220°] (Bulow, A. 236, 188 Phthalamide. Formed from phthalimide and cold NH, Aq (Aschan, B. 19, 1399). Minute crystals, insol. cold water, alcohol, and ether. Boiling water converts it into Alkaline KOBr at 80° forms phthalimide. $C_6H_4 < \stackrel{CO.NH}{NH.CO}$ (Hoogewerf a. Van Dorp, R. T. C. 10, 9).

 $C_6H_4 < CO \longrightarrow 0$ u-Amide [c. 90]. Formed from phthalyl chloride and NH,Aq (Auger, A. Ch. [6] 22, 304). Long transparent prisms, v. sol. water, sol. hot alcohol. AgNO₃ ppts. C₉H₄O₂NAg in white lustrous scales, NH₃ being eliminated.

Phenyl-ethyl-amide $C_{b}H_{4} < CO \xrightarrow{C(NPhEt)_{2}} O.$ [141°]. Formed by heating the ethyl-aniline salt of phenyl-ethyl-phthalamic acid above 200° (Piutti, G. 13, 547; A. 227, 187). Prisms, insol. water, sol. alcohol and ether. Not attacked by KOHAq, but split up by potashfusion into phthalic acid and ethyl-aniline.

Di-phenyl-amide C.H.: C.O. (NPh.), 'Di-phenylamine-phthalein.' [288°]. Formed from phthalyl chloride and diphenylamine (Lellmann, B. 15, 830). Formed also by boiling phthalic anhydride with diphenylamine (P.). Needles (from alcohol). H2SO, containing HNO, forms an intense violet solution.

m-Phenylene-diamide $C_0H_1O_2:N_1H_2:C_0H_1$. [179°]. Formed, together with $(C_0H_1O_2:N)_2C_0H_1$ [252°], by fusing m-phenylene-diamine with phthalic anhydride (Bieder mann, B. 10, 1160). Nodules.

M

p-Phenylene-diamide $C_{\rm e}H_{\rm s}:C_{\rm 2}O_{\rm 2}:N_{\rm 2}H_{\rm 2}:C_{\rm 6}H_{\rm 4}:$ [182°]. Converted by warming with dilute HClAq into a base $C_{\rm 24}H_{\rm 25}N_{\rm 6}O_{\rm 4}$ and $(C_{\rm 6}H_{\rm 4}:C_{\rm 2}O_{\rm 2}:N)_{\rm 2}C_{\rm 6}H_{\rm 4}$ [295°] (Biedermann, B. 10, 1163).

u-Imide $C_{\bullet}H_{\bullet} < C(NH) > 0$. u-Phthal-

imide. [c. 145°]. Formed by the action of HClAq on the u-amide (Auger). Slender needles, exactly resembling the s-imide, into which it changes when heated. Differs from the s-imide by decomposing moist BaCO, in the cold, hence it may possibly be o-cyano-benzoic acid.

s-Imide C_sH₄ CONH. Phthalimide.

[228°]. Formed by heating $C_6H_4(CO_2H)(CO_2NH_4)$ or C₀H₄(CO₂H).CONH₂ (Laurent, A. 41, 110; A. Ch. [2] 61, 121; [8] 23, 119; Lansberg, A. 215, 181). Got also by heating the u-imide, and by the action of CuCy₂ and KCy on o-diazobenzoic acid (Sandmeyer, B. 18, 1499). Sixth of the sublimed. sided prisms (from ether). May be sublimed. Reduced by tin and HCl to phthalidine CaHaO(NH) (Graebe, B. 17, 2598). When distilled with steam over heated zinc-dust it yields $C_{18}H_{11}N$ [100°] (Gabriel, B. 13, 1684). Boiling Ac₂O forms C_2H_1 : C_2O_2 :NAc [182°-185°] (Aschan, B. 19, 1898). Amyl alcohol and Na reduce phthalimide to o-methyl-benzyl-amine (Bamberger, B. 21, 1888). Yields benzonitrile when distilled with lime (Reese, A. 242, 5). Alkaline KOBr at 80° forms o-amido-benzoic acid (Hoogewerff a. Van Dorp, R. T. C. 10, 8) C.H.O.NK. Formed from phthalimide and alcoholic potash (Cohn, A. 205, 801; Gabriel, B. 20, 2225). Plates, con-A. 205, 301; Gabriel, B. 20, 2225). Plates, converted into potassium phthalamate by boiling water. Converted by alkyl iodides into alkylphthalimides. [1:2]C_cH₄(CH₂Br)₂ gives rise to C_cH₄(OH₂N:C_cH₂O_c)₂[253], while the m-isomeride melts at 237°. Reacts with epichlorhydrin, forming C_cH₄O_cN.C_cH₅O [204°] (Goedeckemeyer, B. 21, 2689) with (a) dichlorhydrin, forming (C_cH₄O₂N.CH₂CH.OH[204°], and with \gamma-bromonto butyronitrile, forming C_cH₄O₂N.CH₂-CH₂-CH₂-CN.CH₂-(C₁H₁O₂N)₂Ba 4aq: plates. — (C₂H₁O₂N)₂Cu 4aq. — (C₃H₄O₂N)₂Cu aq. — (C₃H₄O₂N)₂Cu 4aq. — (C₃N₄O₂N)₂Cu 4

Imidoxim $C_8H_8N_2O_2$ i.e. $C_6H_4 < \frac{C(NOH)}{CO} > 0$.

[250°]. Formed by heating o-cyano-benzoic ether with an alcoholic solution of hydroxylamine at 90° for a long time (Müller, B. 19, 1498). Needles (from dilute alcohol), insol. benzene. Boiling with HClAq and FeCl, converts it into phthalimide.

Methyl - imide C,H,O2:NMe. (286°). Formed from potassium-phthalimide and MeI at 150° (Graebe, B. 17, 1174; A. 247, 302). Needles, reduced by tin and HCl to C.H.NO [120°] (800°), which yields a gold salt [196°].

Ethyl - imide C.H.:C.O.NEt. Ethylphthalimide. [79°]. (282°) at 726 mm. Formed by distilling a solution of phthalic anhydride in aqueous ethylamine; and formed also from potassium phthalimide and EtI (Michael, B. 10, 1645; Graebe, A. 247, 802; Wallach a. Kamenski, B. 14, 171). Needles. Br at 140° forms $C_{10}H_aBr_aNO_2$ [o. 189°]. Bromo-ethyl-imide

[83°]. C₈H₄O₂:N.CH₂.CH₂Br. Formed from potassium phthalimide and C.H.Br. (Gabriel, B. potassium phthalimide and C.H.4Br. (Gabriel, B. 20, 2225; 21, 566; 22, 1137). Needles. Converted by aniline into phenyl-amido-ethyl-phthalimide [100°], and by p-toluidine into both (C.H.O.:N.C.H.).NC.H.4Me [200°] and C.H.O.:N.C.H.4NHC.H.4Me [96°]. \psi - Cumidine forms C.H.O.:N.C.H.4NHC.H.4Me.[146°], while (a)- and (B)-naphthylamines form compounds C.H.O.:N.C.2H.4NHC.H.7 [158°] and [141°] respectively (Newman, B. 24, 2196). Potassium sulphocyanide forms C.H.O.:N.C.H.SCy [108°] sulphocyanide forms C₈H₄O₂:N.C₂H₄SCy [108°] (Coblenz, B. 24, 2131).

Sulphydro - ethyl - imide C₈H₄O₂:N.C₂H₄SH. [77°]. Formed from the bromo-ethyl-imide and KSH at 100°. Converted by NaOEt and glycollic chlorhydrin into syrupy C₈H₄O₂:N.C₂H₄.S.C₂H₄OH, whence POCl₃ forms C₈H₄O₂:N.C₂H₄.S.C₂H₄Cl [77°], while POBr₂ forms C₈H₄O₂:N.C₂H₄.S.C₂H₄Br [90°]. Reacts with C₈H₄O₂:N.C₂H₄Br and NaOEt, forming (C₈H₄O₂:N.C₂H₄)₂S [129°], which is oxidised by bromine water to the corresponding sulphoxide [191°], and by chromic acid to the sulphone [256°]. Iodine in alcoholic solution forms (C₂H₁O₂:N.C₂H₄)₂S₂ [139°], whence B"H₂Cl₂ [203°] (Gabriel, B. 24, 1122, 3098). n-Propyl-imide C₂H₁O₂:NPr. [66°].

(283°). Crystals (Gabriel, B. 24, 8105).

Isopropyl - imide [85°]. (278°). β - Bromo - propyl - imide [C₈H₄O₂:N.CH₂.CHBr.CH₃. [105°]. Formed from the allylimide and HBr (Seitz, B. 24, 2627). Converted by KSH into C₈H₄O₂:N.CH₂.CHMe.SH [88°], which is oxidised in alcoholic solution by I to $(C_4H_4O_2:N.CH_2.CHMe)_2S$ [161°]. Potassium sulphocyanide forms $C_4H_4O_2:N.CH_2.CHMe.SCy$ [89°-98°].

γ - Bromo - propyl - imide C,H,O,:N.CH,CH,CH,Br. [73°]. Got from trimethylene bromide and potassium phthalimide Gabriel, B. 21, 2671; 23, 90). Needles (from ligroïn). Converted by sodium ethyl-malonic ether into C₈H₄O₂:N.C₂H₆.CEt(CO₂Et)₂ [62°], and by sodium benzyl-malonic ether into C₈H₄O₂:N.C₂H₆.C(CH₂Ph)(CO₂Et)₂ [c. 110°] (Aschan, B. 23, 3692). Potassium sulphocyanide forms CaH4O2:NO3HaSCy [98°].

Ethylene - di - imide $(C_8H_4O_2.N)_2C_2H_4.$ [232°]. Formed by heating potassium phthalimide with ethylene bromide at 200° (G.).

Butyl - imide C₈H₄O₂:NC₄H₉. Formed by distilling cupric phthalyl-amido-hexoate (Reese, A. 242, 16). Tables (from dilute alcohol).

Isobutyl-imide C,H,O,:N.CH,Pr. [93°]. Formed from isobutyl bromide and potassium phthalimide (Neumann, B. 23, 999). Plates.

Isoamyl-imide C,H,O,:N.C,H,1. (808°). Solidifies when strongly cooled (N.).

Allyl-imide C,H,O,:NC,H,. Allyl-phthal-imide. [71°]. (295°). Tables. Yields a dichloride C,H,O,:NC,H,Cl, [93°], and a dibromide [114°]. When nitrous acid is passed into its solution in cold benzene, and the product heated to 145°, there is formed C₂H₄O₂:NC₂H₆NO₃, crystallising in plates [178°] (Neumann, B. 23, 1000).

Acetonyl · imide C₈H₄O₂:N.CH₂Ac. [117°] Made by heating potassium phthalimide with chloro-acetone at 120° (Goedeckemeyer, B, 21, 2684). Plates and needles. Gives rise to an oxim [172°] and also to a phenyl-hydrazide CH₂.C(N₂HPh).CH₂N:C₈H₄O₂ [152°].

Desyl · imide C₈H₄O₂:N.CHPhBz. [158°].

Desyl-imide C₈H,O₂:N.CHPhBz. [158°]. Formed from phenyl bromo-benzyl ketone and potassium phthalimide (Neumann, B. 23, 994).

Yellowish crystals, v. sl. sol. alcohol.

Phenyl-imide C_6H , C_O NPh. Phthalanil. [205°]. Formed by distilling phthalic acid (1 mol.) with aniline (1 mol.), and by heating phthalimide with aniline (Laurent; Doebner, A. 210, 267; Piutti, B. 16, 1322); Michael a. Palmer, Am. 9, 202). Needles (from alcohol), insol. water. Converted by heating with BzCl and ZnCl₂ into $C_8H_4O_2$:N. C_8H_4Bz [183°] (Doebner, A. 210, 267). The p-chloro-, p-bromo-, p-iodo-, and m-nitro- phenyl-imides of phthalic acid melt at 195°, 204°, 228°, and 243° respectively (Gabriel, B. 11, 2260).

Benzyl·imide C₆H₂O₂:NCH₂Ph. [116°]. Formed from potassium-phthalimide and benzyl chloride (Gabriel, B. 20, 2227). Needles (from alcohol). The o- and m- nitro-benzylimides

melt at 219° and 155° respectively.

o-Cyano-benzyl-imide C₆H₄:C₂O₂:NCH₂C₆H₄.CN. [182°]. Prisms (from HOAc) (G.). The p-isomeride [184°] is converted into CO₂H.C₆H₄.CO.NH.CH₂.C₆H₄.CO₂H [255°] by NaOHAq (Günther, B. 23, 1059).

Tolyl-imides C₆H₄:C₂O₂:NC₆H₄Me. The o-, m-, and p-compounds melt at 182°, 158°, and 204° respectively (Michael, B. 10, 579; Piutti, 4. 227, 205; Fröhlich, B. 17, 2679; Kuhara,

Am. 9, 52).

m-Methyl-bensyl-imide C_oH₄:C₂O₂:N.CH₂C_oH₄Me. [118°]

Formed by heating potassium phthalimide with C₂H₄Me.CH₂Br at 200° (Brömme, B. 21, 2700). White needles, v. sol. alcohol. The compound (C₂H₄:C₂O₂:N.CH₂)₂C₂H₄ [237°] is formed by using [1:8] C₂H₄(CH₂Br)₂.

ψ-Gumyl-imids C₁,H₁₅NO₂ i.e. C₅H₄;C₂O₂;NC₅H₂Me₅. [148°]. (above 360°). Formed by heating phthalic anhydride with ψ-cumidine (Fröhlich, B. 17, 1802). Trimetric crystals. Converted by ammonia into the compound C₅H₄(CONH₂).CONHC₅H₂Me₅ [218°], crystallising in needles, while methylamine and allyl-amine yield the corresponding compounds C₅H₄(CONHC₅H₂).CONHC₅H₂Me₅ [215°] and C₅H₄(CONHC₅H₅).CONHC₅H₂Me₅ [179°].

s-Tri-methyl-phenyl-imide C₀H₄:C₂O₂:NC₀H₂Me₂. Phthalmesidil. [171°]. Got by boiling mesidine with phthalic anhydride (Eisenberg, B. 15, 1017). Silky needles (from alcohol), insol. water. Yields, on nitration, the compounds C₀H₄O₂:NC₀HMe₃(NO₂) [210°] and C₀H₄O₂:NC₀Me₃(NO₂)₂ [242°].

Isocymyl-imide C₂H₄O₂:NC₁₀H₁₂. [145°]. Formed by heating the anhydride with m-isocymidine (Kelbe a. Warth, A. 221, 169). Yields C₂H₄O₂:NC₁₀H₁₂:NO₂ [167°] on nitration.

Naphthyl-imides O₆H₄:C₂O₂:NO₁₆H₇. The (a)-compound [166°] (P.); [182°] (M.) and its (β)-isomeride [216°] are got by heating the corresponding naphthylamines with phthalic an-

hydride (Piutti, G. 15, 479; Maschke, C. C. 1886, 824).

Tri-nitro-phenyl-imide

C_sH₁O_s:N.C_sH₂(NO_s)_s. [259°]. From potassium phthalimide and pioryl chloride (Schmidt, B. 22, 3257). Stair-like groups of crystals.

o-Oxy-phenyl-imide
C₅H.;C₅O₂:NC₅H₄OH. Oxyphthalanil. [220°].
Formed by heating phthalic anhydride with
o-amido-phenol (Ladenburg, B. 9, 1528). Converted by sodium carbonate solution into CO₂H.C₅H₄OO.NH.C₅H₄OH [223°].

p-Oxy-phenyl-imide. [288°]. Got in like manner from p-amido-phenol (Piutti, G. 16, 252). Yields C.H.; C.O.; N.C.H.OAc [239°] and CO.H.C.H., CO.NH.C.H.OH [289°]. HNO, forms C.H.; C.O.; N.C.H.(NO.), OH [210°], which yields

an acetyl derivative [177°].

m-Nitro-phenacylimids C_xH₁O₂:N.CH₂-ÖO.C_xH₂NO₂. [204°]. Formed from ω-bromo-m-nitro-acetophenone and potassium phthalimide (Schmidt, B. 22, 3249). Pointed crystals (from HOAc).

Phenyl hydrazide C₆H₄:C₂O₂:N₂HPh. [178°]. Formed from phthalyl chloride and phenyl-hydrazine in ethereal solution (Pickel, A. 282, 233). Yellow needles, insol. water, sol.

hot alcohol and chloroform.

Hydrides of phthalic acid. When reduced by sodium-amalgam in a solution kept acid by HOAc phthalic acid yields trans (1, 2)-dihydride. Neutral sodium phthalate reduced by sodiumamalgam without addition of acid yields the stable (4,5)-dihydride (Baeyer, A. 269, 154; cf. Graebe a. Born, A. 142, 830). Phthalic soid, suspended in water, is reduced by sodium-amalgam to a lactonic acid which yields a crystalline lactone [255°]; phthalide, and the (1,2)-dihydride are also formed. Both the (1,2)-dihydrides reduce aqueous AgNO₂ (sic) on warming, forming at once a black pp. Boiling aqueous cupric acetate gives off CO₂ and forms a white pp. which on addition of HOAc deposits Cu₂O. The liquid contains benzoic acid (Baeyer, A. 269, 151). None of the other dihydrides of phthalic acid are attacked by cupric acetate. Boiling ammoniacal AgNO, gives with: the (1,2)-dihydride a black pp.; the (4,5)- and (1,6)- dihydrides a brown pp.; the (3,6)-dihydride a white pp. not turning frown. The (1,2)- (1,6)- and (4,5)dihydrides give benzoic acid and CO, when feebly oxidised. The (3,6)-dihydride gives phthalic acid (Baeyer, A. 269, 179). The hexahydrides are most stable, the tetra-hydrides next, and the dihydrides least stable (Baeyer, A. 269, 169). The cis anhydrides are more stable than the *trans* anhydrides. The *trans* acids are more stable than the *cis* acids. The anhydrides of the dihydrides are well crystallised, and yield the corresponding acids when boiled with water. The anhydrides are got by means of AcCl or Ac₂O. The (1,2)-dihydride will not stand AcCl. The (4,5)-dihydride needs long boiling with AcCl. Boiling Ac₂O often changes the trans to the cis variety. The analysis of the cis variety appropriate and the cis of the cis hydrides of the cis melt at lower temperatures than their trans isomerides (Baeyer, A. 269,

In the following hydrides the two carboxyls are in the positions 1 and 2. The small numbers following Δ indicate the position of C

x 2

atoms supposed doubly united to the adjacent C atoms.

acetate gives a flocculent pp., sol. HOAc. Reactions .- 1. Boiling NaOHAq changes it entirely to the (4,5)-isomeride. Nine hours boiling with water does the same.—2. KMnO. hot ammoniacal AgNO_s, and aqueous AgNO_s at 100° are at once reduced.—3. Aqueous cupric acetate forms a green pp., which gives off CO2. On adding HOAc the liquid becomes clear and contains benzoic acid, Cu,O being ppd.-4. Oxidised by boiling CuSO, to benzoic acid (Baeyer, A. 269, 191).—5. The acid takes up Br (4 at.) and the product is reduced by zinc-dust and HOAc to the original dihydride.—6. Combines with HCl, hence cannot be etherified by alcohol and HCl.-7. Combines with HBr (2 mols.) in aqueous solution at 100°. The product could not be crystallised, is reduced by sodium-amalgam to a hexahydride, while zinc and HOAc do not form an unsaturated acid.

Cis (1,2)-dihydride. [175°]. S. 1.8 at 10°. Does not accompany the trans form in the product of reduction of phthalic acid (Baeyer, A. 269, 192). The trans acid boiled with Ac₂O (but not AcCl) for 7 minutes is changed to the anhydride of the cis isomeride. The lead salt of the cis acid is insol. HOAc, while that of the trans acid is soluble therein. Large colourless many-faced prisms. Readily converted into anhydride by warming with Ac₂O. The anhydride [100°] crystallises from ether in colourless needles, and is re-converted into the acid by boiling water. The cis acid is converted into the (4,5)-dihydride by boiling NaOHAq, by 9 hours' boiling with water, and even (unlike the trans isomeride) by standing for 2 days with 15 p.c. NaOHAq. Beacts with silver and copper salts like the trans acid.

Δ^{9,6} or (4,5) · *Dihydride*. CH_CH:C.CO₂H [215°]. S. ·8 at 25°; ·2 at 10°. Got by reducing phthalic acid (60 g.) with sodium-amalgam (1200 g.) (Baeyer, A. 269, 195; cf. Astié, A. 258, 187). Lustrous triclinic crystals.

Reactions.—1. Not affected by evaporating with boiling NaOHAq, but by long warming with very conc. aqueous or alcoholic KOH it is partially changed to the Δ^{2,4} or (1,6)-dihydride.—2. Reduced by sodium-amalgam in a current of CO₂ at 100° to Δ² or (1,4,5,6)- and cis and trans Δ⁴ or (1,2,3,6)-tetrahydrides.—3. KMnO₄ oxidises it to oxalic and phthalic acids (no succinic acid).—4. Alkaline K₃FeCy₆ on boiling forms benzoic acid. Cold dilute H₂SO₄ and MnO₂ also form benzoic acid.—5. PCl₃ gives phthalic anhydride.—6. Br in the dark forms C₃H₂Br₂O₄ [185°].—7. With HBr (2 mols.) it combines forming crystalline di-bromo-phthalic acid bexabydride (Baeyer, A. 269, 198), which is re-

duced by sodium amalgam to phthalic acid trans-hexahydride. Accll converts the acid into an anhydride [157°]. The silver salt boiled with water is reconverted into the (4,5)- dihydride. Alcoholic potash converts dibromophthalic acid hexahydride into the $\Delta^{2,4}$ or (1,6)- dihydride.

Methyl ether MeA'. (250°). An hydride [84°]. Got by boiling the (4,5) acid with AcCl (Baeyer, A. 269, 196). The product is evaporated over soda-lime and H₂SO₄ in vacuo, when the anhydride separates in large tables or prisms, sl. sol. ether, v. sol. chloroform. Hot water readily dissolves the anhydride, converting it into the parent acid. Na₂CO₃Aq forms an orange-red solution, which bleaches litmus as long as any anhydride is undissolved, the blue colour afterwards returning when all is dissolved. On adding H₂SO₄ to the solution a resin is ppd. Heating on the water-bath resinifies the anhydride, forming some phthalic anhydride.

Δ^{2,4} or (1,6)-dihydride. CH CH₂:CH.CO₂H [180°]. Formed by boiling the di-hydro-dibromide of the (4,5)- dihydride of phthalic acid CH₂:CHBr.CH.CO₂H with a solution of KOH (1 pt.) in MeOH (2 pts.) for 1½ hours (Baeyer, A. 269, 199). Formed also in small quantity by heating the (4,5)- acid with conc. aqueous or alcoholic KOH or NaOH. Rosettes of prisms (from water). More soluble in water than the (4,5)- dihydride. Hot cupric acetate forms a white pp. which dissolves on cooling.

Reactions.—1. KMnO, and Na₂CO₃Aq form oxalic acid in the cold, and some phthalic acid. 2. Ammoniacal AgNO₃ forms a white pp. turning brown on warming. -3. Cold dilute H, SO, and MnO, form benzoic acid and CO,-4. Boiling alkaline K₃FeCy, also forms benzoic acid.—5. Converted by Ac₂O in the cold into the anhydride which crystallises from chloroform in cubes [104°] and is reconverted into the acid by warm water. The anhydride dissolves in Na₂CO₂Aq and bleaches litmus meanwhile. The anhydride is converted by heat into the anhydride of the (3,6)- dihydride. — 6. Sodiumamalgam reduces it in the cold (difference from isomerides) to the cis-(1,2,3,6)- tetrahydride.-7. Combines with HBr (2 mols.) when heated at 100° with a solution of HBr in HOAc (Baeyer, A. 269, 200). The product [190°] crystallises from ether in plates, and is reduced by sodiumamalgam to the trans hexahydride.

Δ^{1,4} or (8,6)-dihydride CH.CH₂C.CO₂H CH.CH₂C.CO₂H CH.CH₂C.CO₂H. [153°]. S. 1.7 at 6°. Formed by boiling the Δ^{2,4} or (1,6)-dihydride (1 pt.) with Ac₂O (2 pts.) for 6 minutes, and allowing the product to evaporate over soda-lime and H₂SO₄. The anhydride which separates is boiled with water (Baeyer, A. 269, 204). Large crystals with many facets. The Pb salt is sl. sol. HOAc. Cupric acetate gives a green pp., sl. sol. HOAc. The Ag salt is not reduced by boiling, even in presence of ammonia (unlike any of its isomerides).

Reactions.—1. When heated at 100° for a long time it is partially converted into anhydride. Evaporation of its aqueous solution also partially converts it into anhydride. The anhydride [135°] is also got by heating the acid with Ac₂O. It crystallises in plates.

Readily sublimes in feathery plates. Does not bleach litmus.—2. Sodium-amalgam does not reduce it even when gently warmed (Baeyer, A. 269, 205).—8. Dilute H₂SO₄ and MnO₂ oxidise it to phthalic acid. Boiling alkaline K. FeCy, also forms phthalic acid.—4. KMnO, is at once decolourised.—5. Boiling (10 p.c.) NaOHAq forms (4,5) and (1,6)- isomerides.

 Δ^1 or (3,4,5,6) - Tetrahydride

CH₂·CH₂·C.CO₂H. [120°]. Formed by dissolving its anhydride in boiling water (Baeyer a. Astié, A. 258, 203; 269, 176). Got also by distilling the tetrahydride of pyromellitic acid (Baeyer, A. 166, 346). Monoclinic leaflets (containing aq). When heated at 100° for some time it melts, forming the anhydride. KMnO, oxidises it to adipic acid. Boiling conc. KOHAq changes it to the Δ^2 acid. Oxidised by Br and alkalis to di-oxy-phthalic acid hexahydride (tartrophthalic acid). BaA" aq : crystalline pp.

Methyl ether Me2A". Combines with Br,

forming two dibromides [84°] and [124°]. Anhydride $C_aH_aO_3$. [74°]. Formed by heating the \$\textstyle \archef{a} \text{ acid or its anhydride at 215° for some} time. Plates (from ether), v. sol. ether.

 Δ^2 or (1,4,5,6) - Tetrahydride

CH₂.CH₂.CO₂H. [215°]. S. 88 at 10° H.C.p. 881,600. H.F. 215,400 (Stohmann, J. pr. [2] 43, 540). Got by reducing a boiling solution of sodium phthalate with sodium-amalgam (Baeyer, A. 258, 175). Formed also, together with the trans Δ^4 isomeride, by reducing the (4,5)-dihydride in the same way. Prisms (from water). Bromine vapour yields a dibromide [225°]. MeOH and HCl form an oily methyl ether, which yields a crystalline dibromide [74°]. oxidises it to oxalic and succinic acids.

Anhydride C_aH_aO₃. [79°]. Formed from the acid and AcCl. Prisms. Changed by heat

into the anhydride of the A' acid.

Trans- Δ^{\bullet} or (1,2,3,6) - Tetrahydride CH.CH, CH.CO, H $[218^{\circ}].$ S. .145 at 6°. CH.CH2.CH.CO2H Formed, together with the (1,4,5,6)-tetrahydride, by reducing the (4,5)-dihydride in boiling solution by sodium-amalgam (Baeyer, A. 258, 210; 269, 161). Leaflets (from water). colourises KMnO, at once. Yields a methyl ether Me_2A'' [40°], which forms a dibromide [117°]. Anhydride $C_sH_sO_s$. [140°]. Got from the acid and AcCl. Needles (from ether). Changed by heat into the cis- isomeride.

Cis- Δ^4 or (1,2,3,6) - Tetrahydride [174°]. S. 9 at 6°. Formed by adding sodiumamalgam at 0° to a solution of the Na salt of the (1,6)-dihydride in a current of CO₂ (Baeyer, A. 269, 202). Formed also by boiling the *trans* Δ^4 tetrahydride with Ac₂O for 15 minutes. Large prisms. Boiling Ac2O forms the anhydride, which crystallises from ether in tables [59°].

(or fumaroid) Transhexahydride C₆H₁₀(CO₂H)₂. [221°]. S. 23 at 20°. Formed by reducing the hydrobromides of the di- and tetra- hydrides with sodium-amalgam or with zino-dust and HOAc (Baeyer, A. 166, 350; 258, 214; 269, 161; cf. Mizerski, B. 4, 558). By reducing the (3,4,5,6)-tetrahydride in a hot solution a mixture of trans- and cis- hexahydrides is obtained. Leaflets (from water). May be distilled unchanged if quickly heated, but yields the cis- anhydride when slowly heated. Not oxidised by cold aqueous KMnO. Yields a dimethyl ether [33°], m. sol. ligroin.—PbA" aq: plates.

Anhydride. [140°]. Formed from the acid and AcCl. Long needles (from ether). Changes when heated into the cis-isomeride.

Cis- (or maleoid) hexahydride. [c. 192°]. Got from its anhydride, which is formed by heating the anhydride of the trans-isomeride at 220° for eight hours. Short four-sided prisms, more soluble than the trans- isomeride. attacked by cold aqueous KMnO,. Conc. HClAq at 180° changes it to the trans-variety. The Ba and Zn salts are less sol. hot than cold water.

Anhydride. [32°]. Isophthalic acid [Isophthalic acid [1:3] C₆H₄(CO_.H)₂. 300°]. S. 013 at 25°; 22 at 100°. I H.C.v. 769,100. H.C.p. 768,800. H.F. 190,200 (Stoh-

mann, J. pr. [2] 40, 138).

Formation.—1. By oxidising m-xylene with K₂Cr₂O, and H₂SO₄ (Fittig a. Velguth, Z. [2] 3, 526; A. 148, 11; 153, 268).—2. By fusing sodium formate with potassium m-sulpho-benzoate (V. Meyer, A. 156, 265; 159, 1).—3. The ether is formed by the action of ClCO2Et and sodiumamalgam on m-di-bromo-benzene (Wurster, A. 176, 149).—4. By fusing potassium formate with potassium benzoate (Richter, B. 6, 876), mbromo-benzoate (Ador a. Meyer, A. 159, 16), or di-sulpho-benzoate (Barth a. Senhofer, A. 159, 228).-5. By oxidation of m-toluic acid (Weith a. Landolt, B. 8, 715).—6. By saponification of its nitrile.—7. By heating the hydrides of prehnitic and pyromellitic acids with H2SO4 (Baeyer, A. 166, 334; Suppl. 7, 4).—8. By strongly heating NaOBz (Conrad, B. 6, 1395).—9. By oxidation of colophony with dilute nitric acid (Schreder, A. 172, 93).—10. By the action of a hot aqueous solution of cuprous potassium cyanide upon m-diazo-benzoic chloride, and saponification of the resulting nitrile (Sandmeyer, B. 18, 1498).

Preparation.—m-Xylene is converted, by heating with bromine at 125°, into C₆H₄(CH₂Br)₂, which is boiled with alcoholic potash, and the resulting C₆H₄(CH₂OEt), oxidised with chromic acid mixture (Kipping, B. 21, 46).

Properties.—Long slender needles (from water), m. sol. alcohol. May be sublimed.

Salts.—K,A".—BaA" 6aq. Triclinic crystals (by spontaneous evaporation) (Lossen, A. 266, 30). V. sol. water.—BaA" 4aq? Needles (by cooling hot saturated solutions) .- CaA" 2 taq:

needles.—Ag₂A'': amorphous pp.

Methyl ether Me₂A''. [65°]. Needles
(from dilute alcohol). H.F. 178,300 (Stohmann,

J. pr. [2] 40, 353).

Di-ethyl ether Et.A". [0°]. (285°). Di-phenyl ether Ph.A". [120°]. Got by

boiling the chloride with phenol. Long needles. Chloride C.H. (COCI). [41°]. (276°). Formed by heating the acid with PCl, at 200°

(Schreder, B. 7, 708; Münchmeyer, B. 19, 1849). Amide C.H. (CO.NH₂). [265°] (B. Beyer, J. pr. [2] 22, 851); [above 270°] (Luckenbach, B. 17, 1431). Got from the chloride and NH₂. Plates, sl. sol. water and alcohol.

Nitrile CoH₄Cy₂. [158°]. Formed by distilling potassium cyanide with potassium bens-

ene m-disulphonate (Barth a. Senhofer, A. 174, 236; Meyer a. Michler, B. 8, 672; Nölting, B. 8, 1112; Luckenbach, B. 17, 1428), or with potassium m-chloro- or bromo- benzene sulphonate (Meyer a. Stüber, A. 165, 165; Limpricht, A. 180, 92). Formed also by heating the oxim of isophthalic aldehyde C₆H₄(CH:NOH)₂ with excess of AcCl at 100° for a long time (Münchmeyer, B. 20, 508), by the dry distillation of calcium m-cyano-benzoate (Brömme, B. 20, 521), and by boiling an alcoholic solution of C₆H₄(CS.NH₂)₂ with an aqueous solution of lead acetate (Luckenbach). Small needles (from alcohol), insol. water, m. sol. ether. Converted by alcohol and dry HCl into phthalimido-ethyl ether $C_eH_4(C(NH).OEt)_2$ [66°]. MeOH and HCl yield $C_eH_4(C(NH).OMe)_2$ [c. 62°]; while mercaptan and HCl form $C_eH_4(C(NH).SEt)_2$, which, like the two preceding bodies, forms a crystalline hydrochloride (Luckenbach).

Semi-nitrile v. m-CYANO-BENZOIC ACID. Tetrahydride of Isophthalic acid

C₆H₈(CO₂H)₂. [199°]. Formed by reduction of isophthalic acid by boiling its alkaline solution with sodium-amalgam (Baeyer, B. 19, 1806). Needles, v. sol. hot water.—Ag₂A": white pp. Methyl ether Me₂A". Oil.

Isomeride v. TEREPHTHALIC ACID.

References.—Amido-, Bromo-, Bromo-nitro-, CHLORO-, IODO-, NITRO-, OXY-, and OXY-AMIDO-PHTHALIC ACIDS.

Homo-phthalic acid v. CARBOXY-PHENYL-ACETIC ACID.

Diphthalic acid v. DIPHTHALYLIC ACID.

PHTHALIC ALCOHOL v. DI-OXY-XYLENE. PHTHALIC ALDEHYDE C₈H₄O₂ i.e. C₈H₄(OHO)₂[1:2]. Mol. w. 184. [52°]. Formed by boiling C₈H₄(CHCl₂)₂ (1 mol.) with NaOHAq (4 mols.) (Colson a. Gautier, Bl. [2] 45, 509; A. Ch. [6] 11, 29). Formed also by oxidising C₆H₄(CH₂.OH)₂ (Hjelt, B. 19, 411). Solid, v. sl. sol. water. Coloured blue by NH, Aq. Slowly oxidised by air.

Oxim C₈H₄(CH:NOH)₂. [245°]. Small

needles (Münchmeyer, B. 20, 509).

Isophthalic aldehyde $C_6H_4(CHO)_2[1:3]$. [90°]. Formed by boiling $C_0H_4(\ddot{C}H\ddot{C}l_2)_2$ [1:3] with water (Colson a. Gautier, Bl. [2] 45, 509; V. Meyer, B. 20, 2005). Needles. Reduces silver solution with difficulty. Gives a violet-red tint with rosaniline reduced by SO2. Oxidised by KMnO4 to isophthalic acid.

Oxim C₆H₄(CH:NOH)₂. [212°] (Münchmeyer); [180°] (Meyer). Plates (from hot alcohol). Converted by AcCl at 100° into C₄H₄Cy₂. Yields the ethers C₄H₄(CH:NOMe)₂ [77°] and

C₆H₄(CH:NOEt)₂ [165°].

Reference. -- OXY-ISOPHTHALIC ALDEHYDE.

Isomerides .- TEREPHTHALIC ALDEHYDE and PHTHALIDE.

PHTHALIC ALDEHYDE ACID v. ALDEHYDO-BENZOIC ACID and PHTHALALDEHYDIC ACID.

PHTHALIDE $C_8H_6O_2$ i.e. $C_6H_4 < \stackrel{CH}{CO} > 0$. Lactone of w-Oxy-o-toluic acid. [73°]. (290°

Formation. -1. By reducing phthalyl chloride with zinc and HClAq (Kolbe a. Wischin, C. J. 19, 339), or with Mg and HOAc (Baeyer, Z. [2] 5,899; 10,123,1445; 11,637).—2. Occurs among the products of reduction of phthalic an-

hydride by zinc-dust and HOAc (Wislicenus, B. 17, 2178).—3. By passing bromine-vapour over o-toluic acid at 140° (Hjelt, B. 19, 412).

Preparation.—Phthalimide (1 pt.) is reduced to phthalidene by means of tin (13 pts.) and HCl. The tin is ppd. by zinc, and sodium nitrate added to the filtrate. The yellow pp. of the nitrosamine of phthalidine is filtered off, washed, and heated with dilute NaOH. The phthalide is ppd. by HCl and distilled. Yield on the phthalic anhydride -70 p.c. of the theoretical (Graebe, B. 17, 2599).

Properties.—Needles (from hot water), sl. sol. cold water, v. sol. alcohol and ether. Does not combine with NaHSO₃ (Hessert, B. 11, 238), or reduce ammoniacal AgNO₃. Does not react with

hydroxylamine.

Reactions.—1. Oxidised by alkaline KMnO. to phthalic acid. -- 2. Alkalis and alkaline carbonates form ω-oxy-o-toluic acid, v. vol. iii. p. 781. - 3. Sodium-amalgam forms hydrophthalide C₈H₈O₂, a viscid mass, v. sol. alcohol and ether, and also 'phthalylpinacone' C16H18O. [197°].—4. Ammonia on heating forms phthalimidine.—5. Aniline at 210° forms phenylphthalimidine.—6. Phthalic anhydride on heating forms diphthalyl.—7. KCy at 185° forms C₀H₄(CH₂CN).CO₂H [116°], which forms CaA' 22aq, and yields carboxy-phenyl-acetic acid on saponification (Wislicenus, B. 18, 172; 233, 112). KCy at 200° yields $C_{18}H_{14}N_2O_3$ (?) [240°-245°].-8. Boiling HIAq forms o-toluic acid. 9. Br at 140° forms exo-bromo-phthalide. at 160° yields phthalyl chloride (Racine, A. 239, 79).—10. NaOEt acting on an ethereal solution of oxalic ether and phthalide forms an ether $C_{12}H_{10}O_8$ [122°] crystallising from alcohol in needles, and forming with phenyl-hydrazine the compound $C_{18}H_{16}N_2O_4$ [159°] (Wislicenus, B. 20, 2062; A. 246, 342).

Phenyl-hydrasine compound C₁₄H₁₄N₂O₂i.e. CH₂(OH).C₆H₄.CO.N₂H₄Ph. [174°]. Readily formed by warming phthalide with phenyl-hydrazine for a few hours (Meyer a. Münchmeyer, B. 19, 1707, 2132; Wislicenus, B. 20, 401). Sol. hot water and alcohol, sl. sol. ether. Silvery needles. Partially resolved into the parent substances by fusion. Very unstable towards acids and alkalis. H2SO4 forms a colourless solution, turned reddish-violet by FeCl.

References. — AMIDO-, BROMO-, CHLORO-, NITRO-, and OXY-PHTHALIDE

PHTHALIDE CARBOXYLIC ACID v. Oxy-CARBOXYL-PHENYL-ACETIC ACID

PHTHALIDE SULPHONIC ACID

 $SO_3H.C_6H_4 < CH_2 > 0.$ Formed by warming phthalide with fuming H₂SO₄ (20 p.c. SO₅ extra) (Hoenig, B. 18, 3453), Needles, v. sol. alcohol, insol. ether. - BaA' - CuA' 22aq: light-blue prisms.

PHTHALIDINE is PHTHALIMIDINE. PHTHALIMIDINE C.H.NO i.e.

 $C_6H_4 < CH_2 > NH.$ [150°]. (337°) at 730 mm. Formed by reducing phthalimide with tin and HCl, and by heating phthalide in a current of NH₄ (Graebe, B. 17, 2598; 18, 1408; A. 247, 290; Barbier, C. R. 107, 918). Needles or prisms, sl. sol. cold water, v. e. sol. alcohol and ether. Not volatile with steam. Oxidised by KMnO₄ to phthalimide. $C_{16}H_0N_2O_2Br_8$ [150°]. Dis Bromine forms Distillation over zincdust forms phenyl-isoquinoline. Yields a nitroderivative C₆H₄ CO NH [210°], converted by oxidation into phthalide. Yields also a di-nitro- derivative [195°]

ั เวี๋ 50 º า. Salts. - B'HCl. Thin white needles, v. sol. water.—B'_2H_PtCl₃.—B'_3HAUCl,. [176°].—B'C₆H₂(NO₂)₂0H. [140°].—C₆H₆AgNO. Acetyl derivative C₈H₆AcNO. [151°].

Needles (from dilute HOAc).

C₈H₆(NO)NO. Nitrosamine[156°]. Yellow needles (from water or alcohol). Converted by NaOHAq into ω-oxy-o-toluic acid, and by NaSH into thio-phthalide C.H.OS [60°].

Reference. - DI-CHLORO-PHTHALIMIDINE. pseudo-Phthalimidine C₈H,NO probably $C_0H_4 < CO^2 > NH$. This compound is formed, as hydrochloride, by heating w-chloro-o-toluylamide C₆H₄(CH₂Cl).CO.NH₂ to 150°-160°. The picrate B'C₆H₂(NO₂)₃OH forms a crystalline yellow pp.; the salt B'₂H₂Cl₂PtCl₄ 2aq forms flat orange-yellow needles (Gabriel, B. 20, 2234). ISO-PHTHALIMIDO-ETHYL ETHER

[66°]. $C_6H_4(C(NH).OEt)_2$ [1:3]. The hydrochloride is got by passing dry HCl into a mixture of isophthalic nitrile (1 mol.) and absolute alcohol (2 mols.) dissolved in benzene (Luckenbach, B. 17, 1431). The free base crystallises in small needles, v. sol. alcohol and ether. It decomposes on heating into alcohol and isophthalic nitrile.—B"H,Cl. [270°]. Crystalline. PHTHALIMIDYL-BENZYL v. BENZYLIDENE-

PHTHALIMIDINE

PHTHALIMIDYL-PROPIONIC ACID C₁₁H₉NO₃ i.e. C₆H₄ CO.NH -Ç:CH.CH,.CO,H

[225°]. Formed by dissolving the dilactone of phenyl ethyl ketone dicarboxylic acid in NH₃Aq (Roser, B. 18, 3119). Long yellowish needles, v.

sol. hot alcohol.—BaA'₂.—CaA'₂ aq.—AgA'.

Lactone C₁₁H₂NO₃. [c. 205°]. Formed by evaporating the dilactone mentioned above with conc. NH_sAq on the water-bath. Small tables, v. sol. alcohol, sl. sol. cold water. Reconverted by boiling HClAq into the dilactone. Dissolves easily in cold aqueous alkalis, forming salts of a dibasic acid C₁₁H₁₁NO₄.

PHTHALONITRILE v. Nitrile of PHTHALIC

PHTHALOPHENONE v. DI-PHENYL-PHTHAL-

IDE. o-Phthalophenone C.H.Bz. [1:2]. [146°]. Got by oxidising o-di-benzyl-benzene (Zincke, B. 9,

31). Tables (from alcohol).

Isophthalophenone C_aH_aBz_z [1:3]. Phenylene di-phenyl diketone. [100°]. Formed by the action of benzene and AlCl, on isophthalyl chloride (Ador, B. 13, 320). Plates (from alcohol). Yields two dinitro- derivatives [200°] and [100°]?

Mono-oxim C.H.s.C(NOH).C.H.Bz. [201°] (Nölting, B. 19, 146). Nodules, v. sol. alcohol.

Di-oxim (C₆H₂.C(NOH))₂C₆H₄. [70°-75°] (Münchmeyer, B. 19, 1849). Small crystals. p-Phthalophenone C₆H₄Bz₂[1:4]. Terephthalophenone. (a)-Di-benzoyl-benzene. [160°]. Formed by oxidising a di-based benzene. Formed by exidising p-di-benzyl-benzene with CrO, and HOAc (Zincke, B. 9, 81; Wehnen, B.

9, 309). Got also from terephthalyl chloride, benzene, and AlCl₂ (N.). Plates (from benzene). PCl₅ yields C₂₀H₁₁Cl₄ [92°]. Sodium-amalgam forms di-oxy-di-benzyl-benzene [171°].

Mono-oxim. [213°]. Nodules.

Di-owim. [235°]. Crystals. PHTHALOXYL-AMIDO-ACETIC ACID

C₁₀H₂NO₅ i.e. CO₂H.C₆H₄.CO.NH.CH₂.CO₂H. Glycocoll-phthaloylic acid. [106°]. Formed by the action of alkalis on phthalyl-amido-acetic acid (Reese, A. 242, 6; Gabriel a. Kroseberg, B. 22, 426). Six-sided plates (containing aq).—Na,A".—K,A": hygroscopic needles.—BaA".—Ag,A": plates, sl. sol. hot water.

PHTHALOXYL-AMIDO-BENZOIC ACID CO₂H.C₆H₄.CO.NH.C₆H₄.CO₂H. [277°]. Formed by oxidation of the p-tolylimide of phthalic acid

with KMnO, (Michael, B. 10, 576).

PHTHALOXYL-AMIDO-HEXOIC ACID CO₂H.C₆H₄.CO.NH.C₄H₇Me.CO₂H. Occurs two varieties, the active [132°] and inactive [153°], which are got by boiling the corresponding phthalyl-amido-hexoic acids with NaOHAq (Reese, A. 242, 20). Both acids are decomposed by boiling-water into leucine and phthalic acid.

PHTHALURIC ACID v. Uramic acid of

PHTHALIC ACID.

DIPHTHALYL C16H8O4 i.e.

 C_6H_4 $C_{O.OOO}$ C_6H_4 . [335°]. Formation.—1. By heating phthalyl chloride with reduced silver at 150° (Ador, A. 164, 229). 2. By fusing phthalic anhydride with zincdust; and, together with other products, by reducing phthalic anhydride with zinc-dust and HOAc (Wislicenus, B. 17, 2178).—3. By heating phthalide with NaOAc and phthalic anhydride or thio-phthalic anhydride (Graebe a. Guye, B. 17, 2851; A. 228, 126; 233, 241).—4. By boiling phthalaldehydic ether with alcoholic KCN (Goldschmidt a. Egger, M. 12, 60).

Properties.-Needles, insol. water, v. sl. sol. alcohol and ether. May be sublimed in a current of CO₂. Combines with Br forming C₁₆H₂Br₂O₄ [c. 225]. PCl₃ at 160° forms the chloride C₁₆H₂Cl₂O₄ [245°]. Bromine and water at 100° form bromo-diphthalyl, which crystallises from benzene. Alkalis form diphthalyllactonic acid.

Hydride C16H10O, i.e.

-Ç**H**.ÇÄ- $C_{e}H_{\bullet} \subset CO.O CO.O C_{e}H_{\bullet} [229^{\circ}] (W.); [250^{\circ}]$ (Hasselbach, A. 243, 249), a product of the action of zinc-dust and HOAc on phthalic anhydride (Wislicenus). Needles (from alcohol), v. sol. HOAc. Potash forms di-oxy-di-phenyl-ethane dicarboxylic acid.

References .- TETRA-CHLORO-, NITRO-, and OXY- DIPHTHALYL.

PHTHALYL-ACETIC ACID C, H,O, i.e.

-C:CH.CO₂H. [c. 246°] (G.); [above C.H. < CO.O 260°] (Roser, B. 17, 2620). Formed by boiling phthalic anhydride (10 pts.) with Ac₂O (20 pts.) and NaOAc (2 pts.) (Michael a. Gabriel, B. 10, 391, 1551, 2199). Formed also by dissolving acetophenone ow-dicarboxylic acid in H.SO. (Gabriel, B. 17, 2521). Needles (from nitrobenzene), insol. water, v. sl. sol. hot alcohol.

Reactions.-1. Aqueous NaOH (1 mol.) forms a solution of its Na salt, but excess of alkali converts it into acetophenone ow-dicarboxylic acid. 2. Distillation in vacuo splits it up into CO, and methylene-phthalide.—3. Bromine in HOAc gives CBr_s.CO.C₆H₄.CO₂H [160°]. Dry Br gives C.H.O.: CBr.CO.H [c. 2356].-4. Chlorine acting on the HOAc solution forms COl, CO.C,H, CO,H [144°].—5. H₂SO₄ at 100° forms tri-benzoylene-benzene (v. vol. i. p. 485).—6. Ammonia gives rise to phthalimidyl-acetic acid C₁₀H₂NO₃ i.e. -Ç:CH.CO₂H (?) C,H,CO.NH which crystallises from water in needles [c. 200°] and yields CaA', aq, BaA', 4aq, and AgA' (Roser, B. 17, 2623; Gabriel, B. 18, 2451).—7. NMeH,Aq at 0° forms NHMe.CO.C,H,.CO.CH,CO.H [145°], which is converted by concentrated H.SO, into -C:CH.CO₂H [c. 212°], crystallising $C_eH_4<_{CO.NMe}$ from alcohol in silky needles, split up by heat into CO2 and C6H4 CO.NMe, a crystalline mass which is converted by bromine-water into $C_{10}H_{10}BrNO_2$ [126°] (Gabriel, B. 18, 2453).—8. Aqueous *ethylamine* forms $C_{20}H_{24}O_5N_2$ crystallising from ether in needles [129°], converted by —Ç:СН́.СО₂Н _[180°] cold H2SO4 into C6H4 CO.NEt -C:CH₂ (Merand by heat into liquid C.H. CO.NEt tens, B. 19, 2368).—9. Propylamine forms, in the cold, $C_{25}H_{28}N_2O_4$ [103°], crystallising in prisms (M.).—10. Aniline at 100° forms C₁₈H₁₈NO₂ [192°], crystallising from benzene in cubes, converted by cold H₂SO₄ into C₁₈H₁₁NO [265°] and by heating at 204°-230° into the iso-- C:CH₂ [100°] (Mertens, B. 19, meric C_eH₄ CO NPh 2871) .- 11. Secondary and tertiary amines have

(q. v.).
Salt.—AgC₁₀H₂O₄: amorphous pp.
PHTHALYL-DIACETIC ACID

 C_0H_4 CO.O CO.O

no action .- 12. Sodium-amalgam forms the

lactone of oxy-carboxy-phenyl-propionic acid

i.e. C_eH_4 CO.O. C:CAc.CO₂Et. [124°]. Formed from sodium acetoacetic ether and phthalyl chloride (Fischer a. Koch, B. 16, 651; Bülow, A. 236, 185). Prisms (from alcohol). Decomposed by H_2SO_4 at 65° into HOAc and phthalyl-acetic acid. Alcoholic KOH forms deliquescent crystalline $K_2C_{1e}H_{1e}O_7$. Phenyl-hydrazine acetate forms $C_{2e}H_{1e}N_2O_4$ [238°], which is reduced by zinc-dust and HOAc to carboxy-benzyl-acetoacetic ether

[92°].
PHTHALYL ALCOHOL v. DI-w-OXY-0-

PHTHALYL-AMIDO-ACETIC ACID

C₁₆H,NO₄ i.e. C₂H₄:C₂O₂:N.CH₂:CO₂H. Phthalyl glycocoll. [192⁵]. Formed by heating phthalic anhydride (2 pts.) with glycocoll (1 pt.) (Drechsel, J. pr. [2] 27, 418; Reese, A. 242, 1). Crystals (from water), decomposed by boiling HClAq into phthalic acid and glycocoll. Cold NaOHAq forms phthaloxy-amido-acetic acid (q. v.).—NaA'aq.—NH,A'. [206⁵].—CaA', 2aq.—CuA', 3aq.—AgA'.

-Pt(N₂H₆A')₂. Colourless prisms; got by evaporating a solution of the acid with oxide of plato-diammonium.

Ethyl ether EtA'. [105°] (R.); [113°] (G.). (above 300°). Formed from Ag₂A" and EtI (R.), and also by heating potassium phthalimide with CH₂Cl.CO₂Et (Goedeckemeyer, B. 21, 2688; Gabriel, B. 22, 426). Needles, v. sol. benzene.

PHTHALYL - p - AMIDO - BENZENE SUL-PHONIC ACID C₀H₄:C₂O₂:N.C₆H₄.SO·H. The Na salt, got by heating phthalic anhydride with sodium p-amido-benzene sulphonic acid at 250°, crystallises from water in silky needles (Pellizzari, A. 248, 158; G. 18, 314). It is converted by boiling ammonia into phthalimide and sodium p-amido-benzene sulphonate. The Ba and NH₄ salts are also crystalline.

PHTHALYL - o - AMIDO - BENZOIC ACID C_sH₄:C₂O₂:NC_sH₄:CO₂H. [217°]. Formed by heating anthranilic acid with phthalic anhydride (Gabriel, B. 11, 2261). Prisms, sol. HOAc.

Phthalyl-m-amido-benzoic acid. [276°] (G.); [282°] (P.). Formed from phthalic anhydride and m-amido-benzoic acid, and also by saponifying its ether [152°], which is formed by heating phthalic ether with m-amido-benzoic acid (Pellizzari, B. 18, 216; A. 232, 147). Needles. The amide is described in vol. i. p. 158.

Anilide. [209°]. Prisms (Piutti, B. 16,

1322).

FHTHALYL-AMIDO-ETHANE SULPHONIC ACID C₈H₄O₂:N.C₂H₄.SO₃H. Salt. — KA'½sq. Formed by heating potassium amido-ethane sulphonate with phthalic anhydride (Pellizzari, G. 18, 324). Monoclinic crystals; a:b:c=7.908:1:2.594; β=60° 42′. Decomposed by alcoholic NH₈ into phthalimide and taurine.

PHTHALYL - AMIDO - HEXOIC ACID $C_{14}H_{15}NO_4$ i.e. $C_8H_4O_2$: $N.CH(C_4H_5).CO_2H$. [116°]. [a]_D = -21:87° in a 5 p.c. alcoholic solution at 22°. Formed by fusing leucine with phthalic anhydride (Reese, A. 242, 9). Needles, v. sl. sol. hot water. Lævorotatory. Converted by distillation into an inactive variety [142°], which yields $PtN_2H_6A'_23\frac{1}{2}aq$. Conc. HClAq at 150° forms leucine and phthalic acid. NaOHAq forms $PtN_2H_6A'_23^2$ and $PtN_2H_6A'_23^2$ aq. NaA'.— NH_4A' . [160°-165°].— $PtN_2H_6A'_23^2$ aq.

PHTHALYL-AMIDO-NAPHTHALENE SUL-PHONIC ACID C₈H₄:C₂O₂:N.C₁₆H₆.SO₃H. The salt KA'3aq is got by heating sodium (a)naphthylamine sulphonate with phthalic anhydride (Pellizzari a. Matteuci, G. 18, 321). It crystallises in needles, sol. water.

PHTHALYL- γ -AMIDO-n-PROPYL-MALONIC ETHER C_6H_4 : C_2O_2 :N.CH $_2$ CH $_2$ CH $_2$ CH $_2$ CH(CO $_2$ Et) $_2$ [48°]. Formed by the action of sodium malonic ether on the γ -bromo-propyl-imide of phthalic acid (Gabriel, B. 23, 1767). Monoclinic plates (from ligroin); $a:b:c=3\cdot 260:1:1\cdot 644$; $\beta=81^\circ$ 4′. Converted into δ -amido-valeric acid by heating with HClAq at 190°.

Phthalyl-amido-di-propyl-malonic ether C_aH₄O₂:N.OH₂.CH₂.CPr(CO₂Et)₂. [57°]. Formed in like manner from propyl-malonic ether (Aschan, B. 23, 8698). Prisms, insol. water, y. sol. alcohol and ether.

water, v. sol. alcohol and ether.
DI - PHTHALYL - DI - AMIDO - QUINONE
C₂₂H₁₂N₂O₂ i.e. C₂H₂O₂(N:C₂O₂:C₄H₄)₂[1:2:4:6].
[277°]. Prepared by heating tri-amido-phenol

hydrochloride with phthalic anhydride, dissolving the resulting $C_eH_2(OH)(N:C_eO_2:C_eH_4)_2$ [above 300°] in potash, and treating the product $C_eH_2(OH)(NH.CO.C_eH_4.CO_2H)_2$ [above 800°] with nitric acid (S.G. 1.48) (Piutti, G. 16, 254). Yellow needles (from HNO₂). Forms a compound [174°] with phenyl-hydrazine. H_2S passed into its solution in dilute HOAc forms crystalline $C_eH_2(OH)_2(N:C_2O_2:C_eH_4)_2$, not melted at 310°. PHTHALYL-AMIDO-SUCCINIC ACID

PHTHALYL-AMIDO-SUCCINIC ACID

C₁H₄NO₆ i.e. C₈H₄O₂:N.CH(CO₂H).CH₂CO₂H.

[225°]. Formed by heating aspartic acid with phthalic anhydride for 1 hour at 180° (Piutti, G. 14, 473; 16, 2). Tufts of prisms (from water). Decomposed into phthalic and aspartic acids by heating with HClAq. Boiling aniline forms aspartic acid and the phenylimides of phthalic and phthalyl-amido-succinic acids. Diphenylamine forms three di-phenyl-amic acids C₈H₄O₂:N.C₂H₃(CONPh₂)(CO₂H) [180°], [204°], and [194°]; the first and third crystallises with aq, and all three yield a salt AgA' and are decomposed by potash-fusion into aspartic acid, phthalic acid, and diphenylamine. Salt.—CuA'₂4aq: blue prisms.

Phenylimide C₈H₄O₂N:CH CO.NPh (264°). Formed from the acid and aniline. Small needles (from HOAc), nearly insol. alcohol.

Tetra-phenyl-diamide

C₈H₄O₂:N.C₂H₄:C₂O₂(NPh₂)₂. Two isomerides, [273°] and [286°], are got by heating the acid with diphenylamine for 5 hours at 190°. Both are split up by HClAq at 200° into phthalic acid, diphenylamine, and aspartic acid.

PHTHALYL CHLORIDE v. Chloride of

PHTHALIC ACID.

PHTHALYL-ETHANE v. Anhydride of PHENYL ETHYL KETONE o-CARBOXYLIC ACID.

Di-phthalyl-ethane C₁₈H₁₀O₄ i.e.

 $C_eH_4 < CO.O CO.O C_eH_4$. Ethine-di-

phthalyl. [above 350°].

Formation.—1. Together with other products by the condensation of phthalic anhydride with succinic acid (Gabriel, B. 10, 1559; 19, 837).—2. By the action of conc. H₂SO₄ upon di-phenyl ethylene diketone di-o-carboxylia acid C₆H₄(CO₂H).CO.CH₂.CO.C₆H₄(CO₂H), or its ether.—3. By further elimination of H₂O from the (α) or (β) anhydrides C₁₈H₁₂O₅ of the latter acid (Roser, B. 17, 2770; 18, 3115).

Properties.— Yellow needles (from nitrobenzene), insol. water and alcohol. By boiling with alkalis it is converted into di-phenyl ethylene diketone di-o-carboxylic acid. Nitrous acid forms C₁₈H₁₀N₂O₈ a crystalline body decomposing at 160°, and converted by boiling HOAc into

C₁₈H₉NO₆ [c. 240°].

Isomeride $C_{18}H_{10}O_4$. Formed as a by-product in preparing the preceding body by heating phthalic anhydride with succinic acid and NaOAc. Red needles with green lustre, not melted at 280°. Insol. water and alcohol, v. sol. hot aniline and nitrobenzene. Weak acid, forming unstable violet salts.

PHTHALYL-ETHYL-HYDROXYLAMINE v.

vol. ii. p. 740.

PHTHALYL-HYDROXYLAMINE v. vol. ii. p. 738.

 $\mathbf{C}_{16}\mathbf{H}_{10}\mathbf{O}_{6}$ i.e. Diphthalic DI-PHTHALYLIC ACID CO₂H.C₆H₄.CO.CO.C₆H₄.CO₂H₇(?) Diphthalic acid. [272°]. Formed by oxidation of diphthalyl (Ador, A. 164, 236), of (\$\beta\$)-dinaphthyl-diquinone $C_{20}H_{10}O_4$ (Korn, B. 17, 3021), and of di-phenylethane dicarboxylic acid (Dobreff, A. 239, 68). Formed also by the action of alcoholic potash on diphthalyl dibromide (Graebe, A. 228, 132; 242, 221). Minute tables, almost insol. water, alcohol, and ether. Converted by boiling conc. KOHAq into phthalic acid. Dilute NaOHAq at 110° forms di-phenyl-carbinol tricarboxylic acid. Reduced by HI to (CO₂H.C_eH₄)₂C₂H₄. Alcoholic hydroxylamine hydrochloride forms, on heating, $C_{18}H_{18}NO_5$ [152°] and $C_{16}H_8N_2O_4$ [286°]

Salts: BaA" 2aq: plates.—Ag.A".

Methyl ether Me₂A". [192°]. Formed from Ag₂A" and MeI. Lemon-yellow plates.

By passing HCl into a hot solution of the acid in MeOH there is formed a colourless is omeric body [276°], partially converted into the methyl ether by heating with MeOH at 200°, and split up by HClAq at 150° into diphthalylic acid and

MeCl.

Ethyl ether Et₂A". [155°]. Got from Ag₂A" and EtI. Lemon-yellow needles. The colourless is omeride [174°] is formed by ethylation with alcohol and HCl.

Anhydride C₁₆H₈O₅. [165°]. Formed by heating the acid with Ac₂O at 200°. Crystals

(from HOAc), v. sol. chloroform.

C_eH₄ C(NH).OO.CO C_eH₄? [above 360°]. Got by warming di-phthalyl-lactonic acid with NH₂Aq and by heating phthalimidine with NeOAc and phthalic applydride or phthalimide

DIPHTHALYLIMIDE C16H, NO, i.e.

NaOAc and phthalic anhydride or phthalimide at 220° (Graebe, A. 228, 137; 233, 246). Needles (from HOAc) forming a yellow solution in NaOHAq.

 $\begin{array}{ll} \textbf{DIPHTHALYL-LACTONIC ACID } C_{16}H_{12}O_{\bullet} \text{ i.e.} \\ C_{6}H_{4} < & CO_{.}O \end{array}$

aldehydic acid. Formed by warming diphthalyl, in an atmosphere of H, with alcoholic potash (Graebe a. Schmälzigang, A. 228, 126; of. Ador, A. 164, 229). Crystals (from alcohol). When heated for some time at 220° it splits up into water and diphthalyl, which then melts above 300°. When quickly heated it decomposes at 235°-240° with partial fusion. Conc. H₂SO₄ converts it into diphthalyl. Its alkaline solution is yellow, but becomes colourless through absorption of atmospheric oxygen, yielding diphthalic acid.

PHTHALYL-MALONIC ETHER C₁₅H₁₄O₆, i.e. C₆H₄CO.O C:C(CO₂Et)₂· [75°]. S. (ether) 7·1 at 9°; 58·8 at 35°. One of the products of the action of phthalyl chloride on sodium malonic ether (Wislicenus, A. 242, 23). Triclinic prisms (from ether), v. sol. alcohol.

Reactions.—1. Decomposed by long boiling with water into phthalic acid and malonic ether.

2. Hot potash forms malonic and phthalic acids. A solution of caustic potash at 0° forms

C₀H₄CO₀O(OH).CK(CO₂Et)₂ whence acids

liberate oily 'phthalyloxymalonic' ether, which quickly decomposes into phthalic anhydride and

malonic ether.—8. NaOEt forms the compound C.H.< -C(OEt).CNa(CO,Et), crystallising in prisms, v. e. sol. water, yielding the cupric salt Cu(C₁,H₁₀O₂)₂ 2aq and, on adding an acid, oily C₁,H₂₀O₂. The dry salt, NaC₁,H₁₀O₂ is converted by EtI at 100° into oily C₁,H₁₆EtO,, upon which alcoholic potash reacts forming the salt C_eH_•< -C(OK).CEt(CO₂K)₂ ppd. as very deliquescent plates on adding alcohol to its aqueous solution, yielding amorphous Ag₃C₁₈H₉O₇, and splitting up on acidification into ethyl-malonic acid and phthalic acid .- 4. Zinc-dust and HOAc yield o-carboxy-benzyl-malonic ether (v. vol. i. p. 705).—5. Alcoholic NH, forms the diamides of malonic and phthalic acids.—6. Sodium-malonic ether forms yellow $C_{22}H_{24}Na_2O_{10}$, decomposed by boiling water into phthalyl-dimalonic ether, malonic ether, and phthalic acid.

 $C^{\circ}H^{\circ} \subset CO.\tilde{O}$ Seminitrile -C:CCy.CO₂Et [c. 175°]. Formed from phthalyl chloride and sodium cyano-acetic ether in dry Et,O (Muller, C. R. 112, 1140). White substance, sol. benzene. Phthalyl-di-malonic ether $C_{22}H_{26}O_{10}$, i.e.

C.H.< -C(CH(CO₂Et)₂)₂· [49°]. Formed from sodium-malonic ether and phthalyl chloride (Wislicenus, A. 242, 23, 80). Prisms (from alcohol), insol. water.

Reactions.—1. Potash forms a yellow solution containing C22H24K2O10, which separates as orange needles (containing 2aq) when Et2O is added to a solution of the ether in alcoholic potash. Boiling aqueous potash forms phthalyl-diacetic acid C₁₂H₁₀O₆ [158°].—2. Alcoholic NaOH forms, in like manner, lemon-yellow C₂₂H₂₄Na₂O₁₀ 2aq, decomposed by hot water into malonic and phthalyl-dimalonic ethers and sodium phthalate. EtI at 100° converts the Na derivative into $C_2H_{24}Et_2O_{10}$, which yields ethyl-malonic ether on boiling with water.—3. The di-sodium derivative C22H24Na2O10 is converted by treatment with Ac.O, phthalic anhydride, or phthalyl chloride into 'phthaloxy-dimalonic ether' C₂₂H₂₄O, or C_eH₄ CO.C(CO₂Et)₂ (?), S. (alcohol) ·57 at 14°, which crystallises from ether in needles melting at 117° when slowly heated and at 106° when quickly heated. 'Phthaloxy-dimalonic ether' is also one of the products of the action of phthalyl chloride on sodium malonic ether (Wislicenus, A. 242, 28, 61). It forms a yellow solution with aqueous K₂CO₂ and KOH. Alcoholic potash forms $C_{22}H_{25}\acute{K}O_{16}$, which is the monopotassium derivative of phthalyl-dimalonic ether. Zinc-dust and HOAc reduce 'phaloxy-dimalonic

ether' to oily C₂₂H₂₆O₆.

PHTHALYL-PROPIONIC ACID C₁₁H₆O₄ i.e. C.H. < CO.O [245°-248°]. C:CH.CH2.CO2H Formed by boiling phthalic anhydride (5 pts.) with propionic anhydride (10 pts.) and sodium propionate (1 pt.) for 45 minutes (Gabriel a. Michael, B. 11, 1013, 1679). Slender needles. Converted by boiling KOHAq into phenyl ethyl ketone o-carboxylic acid. Reduced by sodiumamalgam to C.H. -CH.C₂H₄.CO₂H which yields BaA', and AgA' and is converted by

boiling baryta-water into oxy-carboxy-phenylbutyric acid $CO_2H.C_0H_4.CH(OH).C_2H_4.CO_0H.H_2SO_4$ at 100° forms $C_{20}H_{14}O_3$ [237°].

Salt .- AgA': white powder.

Amide C,1H,O,NH, [195°]. Leaflets.

PHYCITE v. ERYTHRITE.

PHYLLIC ACID C₇₂H₆₄O₁₆ (?). [170°]. S.G. 1.014. Extracted by alcohol from the leaves of the cherry-laurel, apple, almond, and elder (Bougarel, Bl. [2] 28, 148). Crystalline granules, decomposing at 180°. The K salt crystallises in needles, sl. sol. water, sol. alcohol.

PHYLLOCYANIN v. CHLOROPHYLL.

PHYSALIN C₁₄H₁₆O₅. Occurs in the leaves of the winter cherry (*Physalis Alkekengi*), from which it may be extracted by chloroform (Desaignes a. Chautard, J. Ph. [8] 21, 24). Yellowish amorphous powder, with bitter taste, v. sl. sol. cold water and ether, v. sol. alcohol. Softens at 180°, and decomposes at a higher tempera ture. Sl. sol. acids, m. sol. NH3Aq. Salt .-Pb_s(C₁₄H₁₅O₅)₂O₂: white pp., got by adding Pb(OAc) and NH3Aq to the alcoholic solution.

PHYSETOLEIC ACID C₁₆H₈₀O₂. [30°]. Occurs in sperm oil, and is perhaps identical with hypogwic acid (Hofstädter, A. 91, 177). Stellate groups of needles. Not affected by nitrous acid.

-BaA'2: crystals, sol. hot alcohol.

PHYSICAL METHODS USED IN CHE-MISTRY. The object of this article is to present a general account of the chief physical methods of inquiry which are made use of in attacking chemical questions. In each section of the article it has been sought to lay down the principles of the method discussed, and to present, in a general way, the more important results that have been obtained. No section claims to give a complete account of the subject with which it deals. The following articles, which are not specially referred to in their alphabetical positions in the present article, should be consulted: AGGREGATION, STATES OF, vol. i. p. 87; Atomic and Molecular Weights, vol. i. p. 336; CHEMICAL AND PHYSICAL PRO-PERTIES OF BODIES, CONNECTIONS BETWEEN, vol. i. p. 780; Densities, relative, vol. ii. p. 370; Dissociation, vol. ii. p. 385; Molecular con-STITUTION OF BODIES, vol. iii. p. 410.

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271 (reference only).

I. CAPILLARITY, METHODS BASED ON. The term 'capillarity' is given to that property of bodies which is the cause of the rise of liquids in narrow tubes, the spherical shape of raindrops and soap-bubbles, the spreading of oil on the surface of water, and many other phenomena.

The generally accepted theory by which these phenomena are connected and explained is due mainly to Laplace, and rests on the assumption that the parts of a body are held together by attractive forces which are insensible except at very small distances. We have evidence of the existence of these forces in the cohesion of bodies, in the latent heat of evaporation, which is merely the energy that has to be supplied to separate the parts from each other, and in the fact that at high pressures, when the parts of which a gas is composed are so near together that they never get altogether away from each other's influence, the compressibility is found to be greater than is required by Boyle's law. Though Laplace made no reference to molecules it is now usually supposed that the attractions in question are the forces acting between the molecules.

Imagine a molecule of a liquid surrounded by a sphere, whose radius is the distance at which the force ceases to be sensible. If the molecule is so far from the surface of the liquid that this sphere is entirely immersed, it will be on an average equally attracted all round, and no work will have to be expended to move it, but if it be so moved that part of the sphere is outside the surface, there will be a resultant force tending to draw it back again, so that to bring a molecule into the surface requires an expenditure of work. Now if we distort a given volume of liquid so as to increase its surface we bring more molecules into the surface, and we increase the potential energy of the liquid by an amount proportional to the increase of surface, or the total potential energy arising from this source is proportional to the extent of surface. Hence, since in the position of equilibrium of any system the potential energy is a minimum, the liquid will of itself assume such a shape that the surface is as small as possible consistently with other conditions, such as the action of gravity.

It is plain that what has been said is not restricted to liquids, but applies equally to solids and to the surface separating two different media. In the latter case a molecule at the surface of separation is acted on by both media, but unless the attractions are equal there will be a resultant force acting on it, and it will have potential energy by virtue of its position.

Capillary phenomena are often regarded as due to the external film of the substance being in a state of tension. It is not easy to imagine a tension at the surface of a solid, but the conception leads mathematically to the same results as the assumption of the existence of surface energy, and is practically only another way of stating the same thing, for we have seen that on the hypothesis of molecular attractions we arrive at the result that the surface tends to become a minimum, and this is equally well

described by saying that it behaves as though it were in a state of tension. Further, the measure of the tension—that is, the total force acting across a line of unit length—is numerically equal to the potential energy per unit area, which is easily proved as follows. Imagine a rectangular piece of the surface a cm. long and b cm. wide increase it to a rectangle a' cm. long and b cm. wide. Then we have produced (a'-a)b sq. cm. of new surface, and if E is the potential energy per sq. cm. we shall have done E(a'-a)b units of work. On the other hand, if T is the surface tension, the force acting on the side b will be Tb, and we have moved it through a'-a cm., and hence have done Tb(a'-a) units of work. These two measures of the work must be equal, and hence E = T; and since the rectangle can be taken anywhere on the surface and with its sides in any direction, it follows that the surface tension is the same at every point and in any direction.

It is to be observed that this tension differs from that of a stretched piece of indiarubber, for instance, in the fact that it does not depend on the amount of stretching. No matter how much a soap film is extended, the tension remains the same until the film becomes so thin that there are only a few molecules in the thick-

The surface tension, then, is a consequence of the molecular attractions which tend to draw the molecules as close together as possible. In order to balance this tendency and to allow the substance to be in equilibrium, Laplace imagined the existence of an internal hydrostatic pressure, which, from the value of the latent heat, is calculated by Stefan to be about 1800 atmospheres in the case of water. The potential energy due to this pressure is proportional to the volume, which is constant in most capillary phenomena, and hence has no effect on the equilibrium position.

The forms of crystals are probably conditioned to some extent by capillary forces. Most of the properties of crystals being different in different directions, the potential energy of a surface may be expected to depend on its direction, and those faces will be formed which have least energy. Further, as a large crystal has less surface in proportion to its volume than a small one, the familiar phenomenon of the large crystals absorbing the small ones is explained by the loss of potential energy resulting from the operation (v. Liveing, Camb. Phil. Trans. 14, 370).

If two immiscible fluids are in contact with a solid, the surface separating them meets that of the solid at a constant angle, called the angle of contact. The most important case is where one of the fluids is air and the solid is glass. Many experiments have been made to determine this angle, the conclusion being that in most cases it is probably zero, but on this point see Magie (W. 25, 429), Quincke (W. 27, 219), and Worthington (P. M. [5] 20, 65).

The constant which is most commonly used is the surface tension as defined above, but Quincke and some others use another called the specific cohesion, denoted by a^2 , which is twice the surface tension divided by the density of the liquid, or the height to which the liquid

would rise in a tube of unit radius.

The following are the principal methods that have been used in determining these constants:

1. By determining the rise of liquids in capillary tubes (Quincke, P. 139, 8; Frankenheim, J. pr. 23, 401).—2. By weighing or measuring the drops from a rod or pipette (Quincke, P. 135, 621; Duclaux, A. Ch. [5] 13, 76; Linebarger, Am. S. 44, 83).—3. By measurement of large drops or bubbles (Quincke, P. M. [4] 41, 245; Eötvös, W. 27, 448; Worthington, P. M. [5] 20, 51).—4. By determining the force required to detach a disc or ring from the surface of a liquid (Weinberg, Z. P. C. 10, 34).

Mendeléeff states as one of the characteristics of a perfect liquid that its surface tension should be a linear function of its temperature, and Selby $(P.\ M.\ [5]\ 31,\ 430)$ has given thermodynamical reasons for this relation. The subject has been investigated experimentally by Frankenheim, Weinberg, and others, and it is found that the equation $\gamma=a-bt$ holds approximately, where γ is the surface tension, t the temperature, and a and b are constants; hence by determination of γ at two temperatures we can calculate roughly the

temperature $\frac{a}{b}$ at which the surface tension is

zero—that is, the critical temperature. Eŏtvös extends this result by showing that the rate of change with temperature of γv^{i} , where v is the molecular volume of the liquid (and hence γv^{i} is proportional to the energy of the amount of surface which contains a given number of molecules), has a constant value which is the same for all liquids.

The only measurements of the surface tensions of pure liquids which have been carried out extensively are those of Schiff (A. 223, 47; and G. 14, 368). Schiff determined the surface tensions of a large number of organic substances at their boiling-points, and divided the observed values by the molecular weights, denoting the quotient by N. This quantity was found to be in general unchanged by the substitution of one carbon atom for two hydrogen atoms, of one oxygen for three hydrogens, and similarly for other elements, so that each atom had its hydrogen equivalent, and compounds with the same total of hydrogen equivalents gave the same value for N. Taking N for ordinate, and x, the total hydrogen equivalent, for abscissa, Schiff plotted a curve from which he obtained the relation

 $\log N = 2.8155 - .00728x - \log x$. This equation enables us to calculate the surface tension of a liquid compound from its formula.

There are many exceptions to the law that each atom has a fixed hydrogen equivalent. For instance, C must be put equal to 8H in the free fatty acids instead of 2H as in most compounds; Cl generally has the value 7H, but when several chlorine atoms are attached to different carbons in a compound it has the value 6H; Br is usually equivalent to 13H, but sometimes to 11H, and so on.

A few preliminary measurements of the capillary constants of the surface separating water and organic liquids which do not mix with it, have been made by Linebarger (Am. S. 44, 88), by allowing the liquid to drop through the water, upwards or downwards, according to its specific

gravity, and counting the drops from a given volume. The results already published show that the introduction of two methyl groups in the meta-position into a benzene ring does not affect the surface tension. while if they are in the para-position the surface tension is greatly diminished.

The surface tension of an aqueous solution of a salt is greater than that of water, and increases proportionately to the amount of salt present. Quincke (P. 160, 337, 560) found that for solutions of chlorides of equivalent concentrations—thatis, containing the same amount of chlorine per c.c.—the constant of proportionality is the same; but Volckmann (W. 17, 353), on repeating the work, concluded that the agreement is not within the errors of experiment. Traube (J. pr. [2] 31, 192) showed that the capillary constant of a 10 p.c. solution of water in alcohol is not raised, like that of water, but is lowered, by the presence of a dissolved salt.

Determinations of the surface tensions of solutions of organic substances in water have not hitherto led to any important general laws. Such substances lower the surface tension but not proportionally to the concentration, as appears from the work of Duclaux (A. Ch. [5] 13, 76), and of Traube (B. 17, 2994; J. pr. [2] 31, 177; 84, 292). The former gives the law that if aqueous solutions of two alcohols, or of two acids, have the same surface tension, the percentage compositions of the two solutions will have a constant ratio. Traube's work confirms this result, but leads to nothing farther of interest beyond the fact that solutions of isomerides have not generally the same surface tensions.

J. W. C.

II. CRYSTALLOGRAPHIC METHODS; v. CRYSTALLISATION, vol. i. p. 278; and Isomorphism, vol. iii. p. 88.

III. DIALYSIS AND DIFFUSION, METHODS BASED ON. When a solid is dissolved in a solvent a movement of the particles occurs from the places where the solution is more concentrated to the places where it is less concentrated, and continues until the concentration is uniform throughout. Similarly when gases which do not react chemically are mixed, movements of the particles take place until the gases are equally distributed throughout the space. The mixing of gases or liquids, by reason of the movements of their particles, is called diffusion. When the diffusion of a liquid is accompanied by a total or partial separation into unlike bodies, the process is generally called dialysis; this process is usually effected by allowing the diffusion to take place through an animal or vegetable membrane. The chemical applications of diffusion are chiefly connected with the diffusion of substances in solution, and generally in solution in water.

Graham (T. 1850. 1, 805; 1851. 483) was the first to measure the rates of diffusion of different compounds, in aqueous solution, without a separating membrane. He nearly filled glass jars with the various solutions, carefully poured water on the top of the solutions, placed the jars in glass dishes, and filled these with water until the water extended in the dishes about 3 c. above the tops of the jars. The dishes were set aside for some time; when the process was to be stopped, glass plates were slid over the mouths of the jars.

which were then removed, and the quantities of | substance in the liquids outside the jars, called the diffusates by Graham, were determined.

Graham found that the quantities which diffused varied much according to the compositions of the diffusing substances. quantity of a specified substance which diffused in a determinate time was found to be nearly proportional to the concentration of the original solution. Graham also found that diffusion was able to effect a tolerably complete separation of two salts whose rates of diffusion differed considerably. He looked on his results as showing that nearly equal quantities of chemically similar salts diffused in equal times.

A few years after Graham's fundamental experiments, Fick, reasoning from Fourier's theory of the conduction of heat, came to the tentative conclusion that the quantity of a salt which diffuses through a stated area is proportional to the difference between the concentrations of two areas infinitely near one another. Assuming the truth of this statement, a definition was obtained for the diffusion-constant of a salt as the quantity of a salt which diffuses through unit area in unit time, when unit difference of concentration prevails throughout unit distance. A long series of researches by Beilstein (A. 99, 165), Simmler a. Wild (P. 100, 217), Stefan (W. A. B. 79, 161), and especially by Weber (W. 7, 469, 536) and Graham (T. 1861. 183), has fully confirmed Fick's law, and has given measurements of the diffusion-constants of many substances. Attention should be drawn, in this connection, especially to Graham's method of allowing salts to diffuse in water gelatinised by starch, gum, &c. (T. 1861. 183; v. also Voigtländer, Z. P. C. 3, 316). In 1880, Long (W. 9, 613) made a number of determinations of rates of diffusion; he divided the numbers obtained by the molecular weights of the salts used, and thus obtained figures which represented the number of molecules of each salt which diffused under the same conditions. The results showed somewhat regular arrangements of the molecular diffusion-values. For instance, Long found that the haloid compounds of K had nearly the same value, that the sulphates of Mg, Zn, Mn, Co, Ni, and Cu had approximately equal values, and so on.

Marignac (A. Ch. [5] 2, 546 [1874]) followed up Graham's observations on the simultaneous diffusion of pairs of salts, and arrived at the important result that the order of the rates of diffusion of the salts of any acid is independent of the nature of the acid, and that the order of the rates of diffusion of the salts of a metal is independent of the nature of the metal. Marignac was thus able to construct a table showing the order of the diffusion-coefficients of acid radicles, on the one hand, and of metals, on the other hand. The table is as follows:-

Cl, Br, I H NO₃ ClO₃, ClO₄, MnO₄ K, NH. Αg \mathbf{F} CrO. Ca, Sr, Ba, Pb, Hg 80, Mn, Mg, Zn Cu

Crystalloids and colloids. Graham observed that the rates of diffusion of different substances | recent work. J. M. van Bemmelen has especially

differ much. He found certain substances which diffused in water with very great slowness; these substances included gums, tannin, albumen, caramel, &c. Inasmuch as the substances which diffused comparatively rapidly generally assumed crystalline forms when they solidified, while substances which diffused very slowly solidified in amorphous forms, Graham called the former crystalloids and the latter colloids. The solution in water of crystalloids is usually accompanied by thermal changes; the solution; boil and freeze at temperatures different from the boiling- and freezing-points of water, and the properties of the solutions differ considerably from those of the solvent. On the other hand, the solution in water of a colloidal substance is not attended with any marked changes.

The solution of a colloidal substance allows the diffusion through it of a crystalloid, in solution, but scarcely permits the diffusion of another colloid. If, then, a solution containing a crystalloid and a colloid is separated from water by a colloidal membrane, such as animal or vegetable parchment, the crystalloid will diffuse through this membrane into the water outside, but the colloid will be retained in the interior liquid; in this way colloids can be separated from crystalloids by diffusion; this process was called dialysis by Graham.

By means of dialysis, Graham prepared many compounds in a colloidal, or jelly-like form (T. 1861. 183). Most inorganic colloidal compounds were found to exist in two forms; one soluble in much water, and the other gelatinous and insoluble in water. For instance, an aqueous solution of silicic acid, containing 14 p.c. of this acid, was obtained by adding a solution of sodium silicate to excess of dilute HClAq, and dialysing (by pouring into a flat saucer formed of parchment paper which was floated on pure water) for some days, until the liquid inside the dialyser ceased to give a reaction with AgNO3Aq; the liquid in the dialyser was then concentrated; by boiling in a flask. The solution of silicic acid gelatinises after a few days, or at once by addition of a trace of an alkaline or earthy carbonate, or by a few bubbles of CO2, or by certain soluble colloids, such as gelatin or soluble The gelatinised colloidal silicic acid alumina. is insoluble in water.

Graham prepared soluble and gelatinised colloidal forms of Al₂O₂, Fe₂O₃, Cu₂FeCy₄, Cr₂O₃, stannic acid, &c. Other soluble inorganic colloidal compounds have been obtained in recent

Graham looked on colloids as very different in their constitution from crystalloids. He regarded colloids as prone to undergo changes which take place very slowly; he thought it possible that the molecule, or molecular aggregate, of a colloid is formed 'by the grouping together of a number of smaller crystalloid molecules.' Colloids, according to Graham, are capable of loosely combining with various proportions of water; this process of 'gelatinous hydration' was regarded by Graham as being 'as truly chemical as that of crystalline hydration.'

Graham's views on the nature of colloids have been confirmed, on the whole, by more studied the hydration of colloids. He gives the name hydrogels to those gelatinous hydrates which contain varying quantities of water not agreeing with any definite formula; he describes the reactions of hydrogels with gases and liquids to form what he calls absorption-compounds, and discusses the part played by such compounds in the soil (L. V. 35, 69; Abstract in C. J. 54, 985). For van B.'s work on various individual colloids v. R. T. C. 7, 37, 69, 75, 87, 106, 114; Abstracts in C. J. 54, 1157-1162.

Picton (C. J. 61, 187) and Picton a. Linder (C. J. 61, 114, 148) have recently prepared a number of soluble colloidal forms of metallic sulphides, such as CuS, HgS, As₂S₈. colloidal solutions were obtained (1) by pouring solutions of the metallic salts into H₂SAq, into which H2S was continuously passed, and dialysing, after removal of excess of H.S by a current of H; (2) by passing H2S into water with metallic hydrates in suspension; (3) by suspending freshly ppd. metallic sulphides in water and passing in H₂S. All the solutions contained combined H₂S, and were therefore solutions of hydrosulphides. The solution of As₂S₈ contained about 5 g. As, S, per litre, and that of HgS about 10 g. per litre. These colloidal solutions were shown to contain solid particles. In some cases the particles were visible through a powerful microscope; in other cases the particles were proved to be present by passing a ray of bright light through the liquids, and showing the scattering of polarised light which thus occurred. Solutions of 'colloidal molybdic acid,' and colloidal silicic acid containing free HCl, seemed to be free from solid particles. Solution of colloidal antimony sulphide showed no particles under the microscope, but the presence of particles was revealed by the passage of a beam of light; after keeping for about ten days, particles had formed sufficiently large to be seen by the help of the microscope; and after some weeks the Sb was all ppd. as Sb₂S₃. In this case the passage could be followed from a liquid containing very minute particles, whose presence was detected only by the fact that they scattered light, to a liquid containing particles sufficiently large to be seen under the microscope. On the other hand, a solution of CrOl, in water containing a little chromous acetate was able to scatter light, and therefore contained solid particles; but after a few days the particles were no longer present. In this case the passage could be traced from a liquid containing very minute particles to a liquid free from such particles.

The colloidal solutions did not diffuse, with one exception—namely, arsenious sulphide. The diffusible solution was obtained by running As₂O₂Aq into H₂SAq, into which H₂S was continuously passed, and removing excess of H₂S by a current of H. This solution showed no particles under the microscope, but as it scattered polarised light particles were present in it. The liquid was put into a small wide-mouthed bottle, which was placed in a beaker and covered with water; a distinct amount of As₂S₂ had diffused in one day, and after eleven days about 12½ p.c. of the As₂S₂ was found in the diffusate. This colloidal solution therefore presented the interesting phenomenon of a liquid containing solid particles capable of scattering polarised

light from a beam passed through, and yet able to diffuse in exactly the same way as true solutions undergo diffusion.

Picton a. Linder consider that their experiments establish 'a good prima facis case for the belief that there is a continuous series of grades of solution passing without break from suspension to crystallisable solution.' They look on the very small particles in some of these colloidal solutions as large molecular aggregates, and they think that these aggregates become very small in the solutions which can diffuse, and that the forces by which the aggregates are held in solution 'become more definitely those of chemical attraction.' (In connection with colloidal solutions v. Paterno, Z. P. C. 4, 457; and Barus a. Schneider, Z. P. C. 8, 278.)

Diffusion of gases. The fact was observed by Dalton (P. M. 24, 8) that if a heavier gas is placed in a bottle which is connected with another bottle containing a lighter gas, and placed beneath the first bottle, after some days the gases will be equally mixed in both bottles. The same fact was observed, and some measurements were made, by Berthollet (Mém. S. d'A. 2,463). Graham (Q. J. S. 1829. 74; P. M. 1833. 175, 269, 851) made a great many measurements of the rates of diffusion of different gases. For most of these he employed a diffusiometer, which consisted of a glass tube about 20 cm. long and about 11 cm. diameter, having a plug of plaster of Paris in one end extending inwards about 1 cm., and graduated from this end downwards. The tube was filled with the gas under examination and placed in water; when the level of the water in the tube had become constant, the total volume of gas now in the tube was measured, and the amounts of air and original gas contained in the tube were determined. With gases lighter than air there was a decrease in the contents of the tube, as the lighter gas passed out through the porous plate more quickly than air passed in; with gases heavier than air there was an increase in the gaseous contents of the tube, as air passed in more quickly than the heavier gas passed out.

The conclusion which Graham arrived at was that 'the diffusion or spontaneous intermixture of two gases in contact is effected by an interchange in position of indefinitely minute volumes of the gases, which volumes are not necessarily of equal magnitude, being, in the case of each gas, inversely proportional to the square root of the density of that gas.'

Modifications in Graham's apparatus have been made, and many determinations of the rates of diffusion of gases have been conducted; the results have fully confirmed Graham's law, which may be stated in the form $c:c' = \sqrt{\bar{a}'}: \sqrt{\bar{a}}$, where c and c' are the diffusion-rates of two gases whose relative densities are d and d'.

It is evident that Graham's law of diffusion gives a means for finding the molecular weights of gases; inasmuch as the law enables measurements to be made of the relative densities of gases, and the density of a gas, referred to hydrogen, multiplied by 2 is (approximately) the molecular weight of that gas. For a description of an instrument for this application of the law of diffusion v. Bunsen's Gasomet. Methoden, p. 160.

M. M. P. M.

IV. DYNAMICAL METHODS. Several methods used in attacking chemical problems may be put together under this general title: v. Chemical change, vol. i. p. 731; Affinity, vol. i. p. 67; Aggregation, States of, vol. i. p. 87; DISSOCIATION, vol. ii. p. 385; EQUILIBRIUM, CHEMICAL, vol. ii. p. 434; MOLECULAR CONSTITUTION OF BODIES, THEORIES OF, vol. iii. p. 410.

V. ELECTRICAL METHODS.

Historical.—The history of the science of electricity divides itself into two well-defined periods, the boundary between which lies at the close of the last century, and is marked by Volta's discoveries concerning the production of electricity when different substances are brought into contact. The investigations in the older period, when only the phenomena of frictional electricity, characterised by small quantity and high tension, were known, showed no connection with chemical problems. It is true, Deimann and Paets van Trostvijk had decomposed water by means of the electrical machine; this effect was, however, assigned as due more to the high temperature of the electric spark than to a specific property of electricity.

It is only with Galvani's discovery of the electricity which appears when different substances are brought into contact, and with the scientific investigation of this discovery by Volta, that the period of electro-chemistry begins. This branch of science is thus of nearly exactly the same age as the current century. It was by means of the pile as constructed by Volta that chemical changes were recognised to be essential phenomena concurrent with the passage of an electric current through certain substances. Directly this apparatus became known, Nicholson and Carlisle (1800) used it for the decomposition of water, and since then the fact that there is a close connection between chemical and electrical phenomena has always been present to the mind of investigators. The mysterious and unexpected mode of action of this apparatus soon revived the hope that by means of it the problem of vital activity might be fathomed. Consequently, the electric current was made to pass through various animal fluids, such as blood, protein, &c., with the object of following the changes produced by this influence, and thus obtaining information concerning the processes taking place in the organism. It was found that, along with other effects, a basic reaction was always observed at one pole, and an acid reaction at the other. On further investigation it was found that the acid and the base still appeared, even when water was taken instead of the animal fluids, and thus the electric current seemed to be a means for producing acids and bases from pure water. This is the point at which the classical researches of Humphry Davy began. In order to decide whether acids and bases were really produced from pure water by means of electricity, he repeated the experiments. He soon recognised that the vessels in which the water was contained exerted a determinant influence on the results; he proved that very small quantities of the substance of the vessels were always dissolved by the water, and that vessels of glass, clay, basalt, &c., were subject to this influence. The electric current had the power of decomposing the very small quantities of saline matter

present into acids and bases, and of accumulating these at the poles, where they could be detected easily. Only vessels of gold proved to possess the necessary resistance, and when these were used not the least quantity of either acid or base was obtained from pure water.

After Davy had thus become acquainted with the great power of decomposing compounds possessed by the electric current, he proceeded at once to submit the most diverse substances to its influence. By means of the large batteries of the Royal Institution, which had been constructed according to his plans, he succeeded in obtaining great effects; he decomposed the alkalis, and isolated the metals potassium and

sodium.

At the present day it is difficult to imagine the impression which this discovery made on his contemporaries. It was not scientific circles only that were full of it; the public at large and the daily press occupied themselves most diligently with this fact. Everyone who could procure some dozens of copper and zinc plates tried to repeat the experiment, and gave an account of it. Napoleon, who then had just nearly reached the zenith of his power, proceeded at once to have larger batteries constructed in order to smooth the way for similar discoveries by the French scientists. He also offered great prizes for scientific works dealing with voltaic electricity.

Davy's discovery was of great importance for the development of the science of chemistry, because it enabled the alkalis to be classed with other basic metallic oxides. Chemical classification was thus simplified considerably. The investigations of Davy were, however, without influence on the knowledge of chemical affinity, sound as were the views held by this man of genius concerning the relation between chemical

and electrical processes.

It was at this same time that the two Swedish naturalists Berzelius and Hisinger carried out work which then, it is true, did not attract anything like the same attention as had justly been aroused by Davy's investigations, but which had an even more lasting influence on the later development of scientific chemistry. Davy's experiments dealt chiefly with the fact that the electric current split compounds into their constituents, and aimed at the isolation of these constituents. Berzelius and Hisinger went a step farther; they tried to grasp the laws underlying this decomposition, and from these they developed a theory concerning chemical compounds. The generalisations under which Berzelius and Hisinger comprised their results were as follows:

(a) Chemical compounds are decomposed by the electric current, and their components collect

at the poles.

(b) The combustible substances, the alkalis, and earths go to the negative pole; oxygen, the acids, and oxidised substances go to the positive

pole.

The fact that Berzelius had experimented chiefly on the salts of the alkalis determined the theoretical conception of electrolysis. Since acids and bases appeared at the poles when alkali salts were electrolysed, acids and bases were considered to be the components of salts. Berzelius assumed further that a similar binary division prevailed throughout the whole domain of chemical com-

pounds. He conceived every atom as endowed with a definite quantity of positive or negative electricity, and this led him to distinguish between positive and negative elements, and to arrange the elements in a series, beginning with the most positive down to the most negative. It was thus that for Berzelius the combination of elements with each other was simply an act of electrical attraction; according to him the thermal and optical effects which are produced along with chemical combination are due to the same cause as the corresponding phenomena which accompany the electric spark. It is true that at this point Berzelius, with the caution peculiar to him, himself brings forward the objection that when the opposite electricities have neutralised each other a further cause of their keeping together is no longer present. It seems, how-ever, that he did not consider this difficulty of sufficient importance to give up his theory for it.

When a positive and a negative atom interact, their electricities, according to Berzelius, are in general not completely neutralised, as the quantities of electricity present differ according to the nature of the atoms. The compound formed thus retains a surplus of positive or of negative electricity, and acts therefore similarly to an element, but less intensely. It is in this way that compounds can again combine with each other to form compounds of a higher order, and so on, and thus is brought about a binary

constitution of all compounds.

This electro-chemical theory of Berzelius, which he developed more fully at a later time, has exerted a very marked influence on the progress of chemistry, since it has impressed on this science the form which was the only one recognised from 1810 to 1840. It was characteristic of this phase that after this first investigation Berzelius did not again undertake any experimental work on the action of electricity on chemical compounds. The places wherein the Berzelian theory was weak from the physical point of view were not considered at all, as the theory was used only for the purposes of chemical classification; no attempt was made to explain by means of it the problems of chemical affinity.

After an almost uncontested reign of twenty years' duration the theory of Berzelius proved itself insufficient to follow the progress of the science. Since it was deduced from the phenomena of the decomposition of compounds by electricity, it was not surprising that it could not represent the chemical relations of organic compounds, which, as a rule, are not decomposed by electricity. Investigations on this subject proved more and more conclusively that in chemical compounds individual atoms could be substituted by other atoms or groups of atoms, quite independently of the 'electro-chemical' nature of the elements; in the most conspicuous and best-known examples it was a question of the substitution of 'positive' hydrogen by 'negative' chlorine. Such a process Berzelius considered to be quite impossible in the light of his theory

But it was not this proposition of the electro-chemical theory alone, but the whole foundation of Berzelius's system, which was made doubtful and proved to be untenable by the newer development of organic chemistry. The theory of the binary constitution of chemical compounds was no longer capable of being brought into accord with facts, which rather led to the unitary conception of substances. Closely connected with this development is the establishment of the conception of the chemical molecule, a conception which assigned a sharplydefined existence to the combinations of the elementary atoms, and which led to the view that these structures would split up in the most diverse ways, according to the nature of the influences to which they were subjected, although no such division appeared pre-existing in the molecule.

At the same time (after 1840) at which purely chemical facts had proved Berzelius's system insufficient for the domain of organic chemistry, the insufficiency of its physical foundations was also made apparent. This happened in consequence of Faraday's funda-

mental work.

A lively discussion concerning the cause of the production of electricity in the galvanic pile had been carried on ever since the days of Volta. While Volta and his successors sought for the cause of the electric tension at the extremities of the pile in the contact of the metals, a number of other workers held to the opinion, as first expressed by Fabbroni, that the cause of galvanic electricity was to be found in the chemical processes which take place in the pile. It is not possible to enter here into the history of the contest which has lasted up to the present day. Faraday attempted to solve the problem, and in so doing he discovered the electrolytic law (1834) which goes by his name.

This law affirms, firstly, that when electricity passes through a body which is decomposed by it—that is, through an electrolyte—the quantity of substance decomposed is proportional to the quantity of electricity that has passed through. Secondly, the law affirms that when the same quantity of electricity passes through different electrolytes, the quantities of the different substances which are thereby decomposed are to each other in the ratio of their chemical equiva-

lents.

These two generalisations primarily supported the chemical theory of galvanic electricity, as according to them a galvanic current is never possible without a corresponding chemical process. At the same time, however, they were in unresolvable contradiction to the foundations of the theory of Berzelius as conceived by him; since, if one and the same quantity of electricity is always necessary in order to decompose chemically equivalent quantities of any substances, it cannot be true that different quantities of electricity cause their combination, in which act they neutralise each other. Berzelius was keenly alive to this contradiction, but as he did not wish to doubt his own theory he preferred to doubt the laws of Faraday, and he continually argued against them.

It was through the work of Daniell (T. 1839. i. 97, and 1840. i. 209), which followed up the researches of Faraday, that the electro-chemical theory was fought on the very ground from which it had sprung. The results of the elec-trolysis of neutral salts, such as potassium sulphate or sodium chloride, when the two elec-

trodes of the electrolytic cell were separated by a porous diaphragm, showed that detonating gas (or hydrogen) was formed in the same quantity as in a voltameter with dilute sulphuric acid inserted in the circuit, and besides this an equivalent quantity of the salt was decomposed into acid and base. This phenomenon cannot be brought into accordance with the law of Faraday otherwise than by assuming that electrolysis does not split up the salts into base and acid, but rather into the metals and into the elements (such as Cl) or radicles (such as SO₄) combined with these. Thus, when potassium sulphate is electrolysed, hydrogen and oxygen are only secondary products, just as the free acid and the base are only secondary products; the salt K2SO4 splits up rather into free potassium K2, which acts on the water, forming potash $(K_2+2H_2O-H_2+2KOH)$, and into the radicle SO₄, which with water gives sulphuric acid and oxygen $(SO_4 + H_2O = H_2SO_4 + O)$. Correspondingly, when copper sulphate is electrolysed we obtain, not copper oxide and sulphuric acid, but on the one side metallic copper, and on the other side SO. which gives sulphuric acid and oxygen. Thus Daniell arrived at the conclusion that the conception of salts as consisting of base and acid would have to be given up, and that they rather consist of a metal and a simple radicle (Cl, S, &c.), or a compound radicle (SO, NO, &c.), and he further pointed out that Davy had already expressed similar ideas.

The same result to which the application of Faraday's law led in this case had meanwhile been arrived at by chemical methods. After Graham (T. 1833. ii. 253) had reduced the differences exhibited by the salts of phosphoric acid to differences in the amounts of water held by the acid, and had thus founded the theory of the polybasic acids, Liebig (1838), in his celebrated paper (A. 26, 113), enunciated the theorem that all acids must be looked on as hydrogen compounds, and that the formation of salts consists in the replacement of this hydrogen by metals or metallic radicles. By means of this conception the unnatural division between the salts of oxyacids on the one hand, and the halogen compounds of metals on the other, was again put aside. The necessity for such a division had been felt by Berzelius to be a great trouble, and to do away with it he had for long clung tena-ciously to the assumption that the halogens were

oxygen compounds. The first attempt to utilise the facts of galvanism for chemistry had failed. In 1850 Berzelius's electrochemical theory had but few supporters left. Still the consciousness survived that the mistake had lain in the form in which the relation between chemical and electrical properties had been represented; and it continued to be recognised that this relation actually did exist, and that it was of the greatest importance. A sign of this recognition is found in the fact that although Berzelius's theory had been rejected, the elements which form bases were still called positive and those which form acids were called negative. In fact, the electrochemical contrast of the two constituents of salts is a fact which becomes obvious with each electrolysis. and which, therefore, cannot be doubted. It was only necessary to modify the view of Berzelius

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and to say that it is not the acids and bases (or more correctly their anhydrides) which are the constituents of salts, but rather it is the metal and the acid radicle. This modification of the electrochemical theory has, however, scarcely left any traces in the development of the chemistry of that time. The reason of this is that compounds of the nature of salts receded quite out of the foreground of interest. Organic chemistry, which at that time developed brilliantly, dealt with substances very few of which could be decomposed by the electric current, which were not electrolytes, and for which, therefore, the electrochemical contrast did not exist. The unitary conception was consequently involuntarily extended to the salts, and the important distinction between electrolytes and non-electrolytes was not at all taken into account by chemists. It is true that Faraday attempted to account for these two classes of bodies by supposing that in the one case the number of positive and negative atoms was the same, but in the other case these numbers were different; but this rule proved to be incorrect, and for a long time these relations were neglected because they could not be grasped scientifically. After the refutation of the erroneous views of Berzelius, people thought themselves justified in altogether ignoring the electrochemical relations.

This was the condition of electrochemistry till quite recently. The only attention it received was from the hands of some physicists, and it is only natural that purely chemical problems did not fare particularly well under these circumstances, the more so as even up to the present day it often happens that from this side the electrolysis of dilute sulphuric acid is represented as the electrolysis of water 'which has been made conductive by an addition of sulphuric acid. Although the direct furtherance thus given to chemistry was insignificant, yet the indirect effects were important. If it is possible to speak to-day of the development of a new electrochemical theory, it is almost exclusively to these physicists that we owe the means for so doing.

The first investigator to be mentioned at this point is Hittorf. He connected his work with the experiments of Daniell, and took up the consideration of a phenomenon which had remained incomprehensible to the latter. When Daniell experimented with sulphuric acid in his apparatus, he found that besides the electrolysis of the acid a change in the concentration had occurred; at the negative pole the acid solution had become more dilute, at the positive pole it had become more concentrated. It occurred to Hittorf that this phenomenon must be due to the different velocities with which the two constituents or ions of sulphuric acid—that is, 2H and SO,—travel through the liquid. If, for instance, the hydrogen remained at rest and the group SO, alone moved, it would follow that after the electrolysis of one molecular weight of sulphuric acid, there must be an increase, by that amount, in the concentration at the positive pole towards which SO, had travelled. If, on the other hand, the hydrogen alone travelled, the concentration would remain unchanged. Now, it had been found by Daniell that the increase in concentration at the positive pole was equal to less than a quarter of the quantity of acid electrolysed; the necessary

conclusion was, therefore, that both ions travelled, but that SO₄ moved much more slowly than 2H. Hittorf proved by a great number of careful experiments (P. 89, 177; 98, 1; 103, 1; 106, 337) that this conclusion always agreed with observed facts. He then used the knowledge thus gained to answer chemical questions, for the solution of which no other means existed at the time. But in spite of their great importance, Hittorf's results have been all but completely ignored by chemists.

These experiments afforded verification of the conclusion that the more immediate components of salts - or what is the same, the ions of saltsreally are the metal and the acid radicle. At the same time electrolysis furnished Hittorf with the means of solving certain old problems. Thus, for instance, opinion was divided as to whether potassium platinichloride and similar salts should be considered as double salts—for example, as 2KCl and PtCl₄—or as salts of chloroplatinic acid H₂PtCl₈. Hittorf submitted sodium platinichloride to electrolysis. If it consisted of 2NaCl+PtCl₄, it followed that sodium and platinical states of the sodium and states of the sodium and states of the sodium and states of t num were the positive ions and chlorine the negative; if, on the other hand, it was Na₂PtCl_s, the ions would be 2Na and PtCl. In the first case, therefore, the platinum must travel to the negative pole, in the other case to the positive. Experiment decided for the latter view; the platinum did not go to the negative pole, as the metals generally do, but travelled with the chlorine to the positive pole, thus proving itself to be a constituent of the acid radicle. In a like manner Hittorf decided quite a number of similar questions.

During these investigations, Hittorf drew attention to another point, which at a later time proved to be of the utmost importance. The fact that electrolysis can be started by the weakest currents is in contradiction to the usual chemical views, according to which the constituents of salts, such as KCl, Na, SO,, &c., are held together by very strong affinities. At the same time, Hittorf emphasised the fact that those substances which conduct electrolytically are the most ready to interchange their constituents. This fact also is against the assumption of a specially firm binding together of the constituents of salts. Occasion will be found later to refer to this remark.

There are other parts of the science of electricity, besides the phenomena of electrolysis, in which the chemical nature of substances has to be considered. These are the electrical conductivities of electrolytes, the electromotive force of galvanic cells, and galvanic polarisation. These fields have been cultivated till quite lately only by physicists to whom purely chemical questions were foreign.

The electrical conductivity of an electrolyte is a quantity the determination of which was formerly attended with great difficulties. These difficulties occur because the ions which separate where the electric current enters or leaves the liquid 'polarise' the electrodes, and thus produce new and unknown electromotive forces. Details of the various attempts which have been made to overcome this difficulty will be given later. It was, however, only in 1880 that, after long and varied preliminary investigations, F. Kohl-

rausch (W. 11, 653) indicated a really practical and accurate method. As soon as the values of the electrical conductivities of electrolytes could be determined easily, by Kohlrausch's method, important relations soon came to be recognised between the conductivities and the chemical properties of electrolytes. The most important of these relations, the discovery of which was made by Arrhenius (Bigh. Swensk. Ak. 8, Nos. 18 and 14, 1884), lies on the path opened up by Hittorf. The fact that substances capable of conducting the current and of being electrolysed are also specially capable of entering into chemical reactions, which was emphasised by Hittorf, can now be made definite by saying that both powers are nearly proportional for different electrolytes.

From this discovery there has grown a new electrochemical theory, which, in its entirety,

will be expounded later on.

The other problem concerning the connection between the electromotive force of galvanic cells and the chemical processes within them was solved by Helmholtz (Die Erhaltung der Kraft, 1847), and later on also by Sir William Thomson (P. M. December, 1858), and this was done primarily on the basis of a hypothetical assumption. According to Faraday's law, when equivalent quantities of different substances are used in the galvanic cell, equal quantities of electricity are always put into motion. This being so, the intensity of the motion, or the electromotive force of the galvanic cells, must be proportional to the quantity of heat produced by the chemical processes within the cell. The supposition is made here that all heat is used for electrical work-a supposition which Thomson found verified by an experiment of Joule. Meanwhile it has been proved, however, that the heat produced in chemical changes is not generally completely transformed into electrical work, and Helmholtz (B. B. 1882) himself has worked out the formula representing the general relation between these two quantities.

To these problems are joined those concerning the nature and the value of galvanic polarisation. In this province investigation has, however, made so little way as yet that it suffices to point out that most important problems here await their solution.

The foundations of the science of electricity. General considerations.—If a bar of perfectly pure or amalgamated zinc is placed in dilute sulphuric acid, no chemical action takes place, nor is there any such action if a bar of platinum be introduced into the acid. But as soon as the two metals are made to touch each other an action occurs; the zinc is transformed into zinc sulphate by expelling the hydrogen from the sulphuric acid, which hydrogen, however, is evolved at the surface of the platinum har.

Direct contact of the two metals is not necessary. If, for instance, the zinc is touched with the end of a copper wire and the platinum is touched with the other end, the same effect is produced. On the other hand, there is no effect if the connection is made through glass, wood, or sreh like substances.

The copper wire which connects the two metals has acquired special properties which remain as long as the chemical action of the sulphuric acid on the zinc lasts. If the wire is held parallel to a movable magnetic needle at a short distance above it, the needle will be deflected; further, the wire becomes heated; and finally, if the wire is cut in one place, and the two ends are placed close to each other on a paper moistened with solution of potassium iodide and starch, it can be perceived that a chemical decomposition of the potassium iodide has occurred, since a blue spot of iodide of starch appears under one end of the wire.

Work, or, speaking generally, energy, is gained by the chemical action between the zinc and the sulphuric acid; the energy usually appears in the form of heat. The arrangement described shows that it is possible to conduct this energy away from the place where it is set free—that is, from the point of contact between zinc and sulphuric acid—and to make it effective at any point of the connecting wire we please. And further, we can obtain this energy at will in the form of thermal energy, chemical energy, or mechanical energy.

The sole difference that can be perceived between the metals in their usual state and when dipped into the acid is that they have become electric; the platinum shows itself charged with positive, and the zinc with negative, electricity. By means of a suitable electrometer this electric charge can be measured. It appears to be dependent on the nature of the metals, as well as

on that of the acid.

Hence we conclude that the chemical action between the zinc and the acid does not, as usual, give out its energy in the form of heat, but in that of electrical energy. As to whether the transformation from chemical to electrical energy is complete or partial, and, if the latter, by what this partial amount is determined, these are questions with which we can occupy ourselves only at a later stage.

As is generally the case with all forms of energy, electrical energy allows itself to be resolved into two factors, one of which is a capacity, and the other is an intensity. For these factors the general law holds, that a system can be in a state of rest only if the intensity of the energy is the same throughout the whole. An exception to this occurs only if different kinds of energy are present simultaneously in one system; then there can exist a difference of one intensity, if a compensation is produced by a corresponding difference of the other intensity.

The two factors of electrical energy are called quantity of electricity and electromotive force or potential; the first is the capacity, the second the intensity. In accordance with the above-mentioned law, we shall therefore say that in a system in which electrical forces alone act there is equilibrium if the electromotive force or the potential is the same throughout. This is the well-known law for conductors of electricity.

tricity.

With substances which do not allow of any motion of the electricity—that is, with non-conductors—the potential can, it is true, be different in different places. But then, owing to the striving for equalisation on the part of the potential, small displacements are produced in the body, and the reacting forces of elasticity which accompany these form the compensation

for the inequality of the electric intensity or potential.

When contact between two different substances occurs, a mutual action at the surface of contact is generally set up, and a displacement of energy is thus brought about. The changes in the surface-energy which thus occur seem to be generally compensated by corresponding differences in the electric intensity, i.e. the surfaces of different substances in contact attain different potentials. If the substances are conductors of electricity, each of them must be at a uniform potential at every point within it; hence a difference of potential exists at the surfaces of contact only.

Electrical measurements. In order to examine systematically the nature of electrical phenomena we are at the very beginning obliged to find a measure for them. We generally start from a definition of quantity of electricity, since the historical development of the science has led to the notion that this is one of the most important factors in the phenomena. And yet we have no proof that such things as the hypothetical electric fluids actually exist. What we do know of electrical phenomena are the mechanical, thermal, and chemical effects, i.e. the manifestations of the electrical energy, and it is this which is the real thing underlying the electrical phenomena. Quantity of electricity is a magnitude of the same order as perhaps a volume, and one which can be increased or decreased at will. But electrical energy cannot be destroyed, nor can it be produced; it can only be changed into, or obtained from, other forms of energy.

Electrical energy is then to be measured by the same units as other forms of energy, mechanical energy especially. The unit used for kinetic energy, which is represented by the for-

mula $\frac{m}{2}v^2$ (where m stands for the mass, v for

the velocity), is double that energy which the mass of one gram possesses when moving with the velocity of one centimetre in one second. This being so, the unit accepted for electrical energy is that quantity of energy which is obtained by the transformation of the above-defined quantity of kinetic energy into electrical energy. And further, since, as has been already mentioned, electrical energy has to be looked upon as the product of two factors, the quantity of electricity Q and the potential E, the unit by which the product QE must be measured is thus also fixed.

The units of the two factors Q and E are still arbitrary, in so far as we can choose one of them at will, that for the other being then fixed. In fact, different units have been chosen for different purposes, and accordingly different systems of electrical magnitudes have been obtained. Faraday's law, that equal quantities of electricity travel with equivalent quantities of separate ions, supplies the chemist with a natural unit for quantity of electricity. Hence that quantity of electricity which is combined with one gram of hydrogen as ion will serve as unit of quantity of electricity. Then the unit of potential would have to be the potential at which the above-defined quantity of electricity must be in order to produce unit of work.

This system of units has not become general;

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it has been displaced by a system which is derived in a somewhat complicated manner from the action of current electricity on magnets. This system also has not been retained unchanged, but another one has been deduced from it, in which instead of the unit of length of 1 cm. a 10°-fold value has been introduced, while instead of the unit of mass of 1 g. a value 10⁻¹¹ times as small is accepted; the second has been retained as the unit of time. Hereby the unit of electrical energy has been changed also; it no longer coincides with the mechanical unit, but is 10' times greater.

In this system the unit of potential is called a volt; in order to get a conception of its magnitude it is to be remembered that the difference of potential at the ends of a Daniell cell is about 1-1 volt.

The unit of quantity of electricity has been called a *coulomb*. When a coulomb is forced through a volt a quantity of work equal to 10⁷ mechanical units is done.

In order to fix the relations of these units to other units of energy, we must first remember that the work necessary to move 1 g. through 1 cm. against gravity is equal to 980 mechanical units, approximately. The above-defined unit of electrical energy would therefore be equal to

 $\frac{10'}{980}$, or approximately to 10200 gravitation-units.

Further, the unit of thermal energy is the quantity of heat which raises the temperature of 1 g. of water by 1°C., and which, according to the measurements of Joule, is equal to 42350 gravitation-units, i.e. 42350 g. would on falling through 1 cm. give up 1 calorie of heat, or would heat 1 g. of water by 1°C. Hence this magnitude is equal to 4.15×10^7 mechanical units. Since the unit of electrical energy amounts to 107 mechanical units, it follows that 1 calorie is equal to 4.15 electric energy units, or $1 \text{ volt} \times \text{coulomb}$ is equal to 0.241 calories. A number obtained more recently by Dieterici by direct measurements is probably more accurate. According to him, the energy of 1 volt × coulomb is equal to 0.2356 cal.; the unit of quantity of heat here used is $\frac{1}{100}$ of the quantity of heat given out by 1 g. of water in cooling from 100° to 0°. This factor is important for the relations between the heat produced in chemical changes and the corresponding electrical phenomena. Finally, we know from measurements by F. Kohlrausch and Lord Rayleigh that 1 g. of hydrogen as ion carries with it 96540 coulombs. The same quantity of electricity is carried by each equivalent of any other ion, for instance, by 107 g. of silver, $\frac{1}{3} \times 63.4$ g. of copper, $\frac{1}{8} \times 27$ g. of aluminium, &c.

Electric currents. If the potential of electricity is different at different places of a conductor, a movement of the electricity is produced in the same way as motion is produced in a mass capable of moving freely when it is at a higher level than its surroundings. If the difference of potential is maintained, the movement is maintained also.

The compensation of a difference in electric potential necessitates a decrease in the electric

energy, which in this process changes into another form. The form most easily produced is thermal energy, but, as has been mentioned

above, it is possible also to obtain mechanical or chemical work.

The measure of the quantity of energy changed into another form is given by the product of the quantity of electricity moved into the decrease of the potential. If the differences of potential are distributed over measurable distances the process is called an electric current The term has been taken from the analogy which in fact does exist between currents of water and currents of electricity. The level of the water corresponds to the potential of the electricity, and the quantity of water corresponds to the quantity of electricity. In both cases the transformable energy, or the available work, is equal to the product of the quantity (of electricity or of water) into the difference of level or of potential, and a current sets in when a difference of level exists. It must only be borne in mind that the phenomena which are brought about by the kinetic energy of the moving masses in water currents have no analogy in the domain of electricity. The electricity behaves as if it possessed only an inappreciably small velocity or mass. Therefore, when the electricity has sustained a definite decrease in potential, it has lost the whole corresponding quantity of energy, while the water can retain a part of it in the form of velocityenergy.

Electric currents are measured by their intensity. By this is understood the quantity of electricity which flows in the unit of time through a cross-section of the current-path; hence the intensity I has to be put as $\frac{Q}{t}$, where t stands for the time. Since electricity moves only when using energy, a loss of electrical energy corresponds to every current, the lost energy generally reappearing as heat. When heat is the only form into which electric energy is transformed, the following equation must hold; W = EQ, and $\frac{W}{t} = EI$, where E stands for the loss of potential between the two ends of the

loss of potential between the two ends of the path of the current considered.

Experience has further shown that different conductors when introduced into a circuit cause different losses of potential along their lengths. This property has been ascribed to a resistance of the conductor to the motion of the electricity, in a way similar to that wherein tubes of different bore offer different resistances to the motion of water within them. In accordance with this, the resistance R is defined as the ratio between the difference of potential E and the quantity of electricity forced by means of it, in unit time, through the conductor, or, what is the same, the intensity I. Here we have $R = \frac{E}{T}$ or $I = \frac{E}{R}$. This is the

celebrated law of Ohm, that the intensity or the strength of the current is equal to the ratio between electromotive force and resistance.

Combining this law with the preceding one, it follows that $\frac{W}{t} = I^2R$. This expression is the law arrived at experimentally by Joule, according to which, for currents of equal strength, the heat evolved in unit of time is proportional to the resistance, and for equal resistances it is proportional to the square of the current strength.

It is easy to deduce from the units already given the values for the units of the two new terms introduced when defining electric currents. namely, the strength of the current S and the resistance R. The unit of intensity I is that strength of current at which, in every second, one coulomb flows through the cross-section of the conductor; the unit of intensity is called an ampère. The unit of resistance is that resistance by means of which a current of unit intensity (1 ampère) produces in unit of time the unit of heat. As we have seen above, this last is equal to 107 mechanical units. or to 0.2356 calories. An attempt has been made to represent this resistance with the utmost possible accuracy, and it has been found that it equals the resistance of a column of mercury of 1 sq. mm. section and 106.3 cm. in length, at the temperature of melting ice. This unit of resistance has the name ohm. Slightly different to this is the legal ohm, which has been fixed arbitrarily as equal to the resistance of a column of mercury 106 cm.long and 1 sq.mm. section, the exact determination of the true ohm being a very difficult operation. The commercial resistances are graduated according to the legal ohm, and in scientific works in which the absolute value of the ohm comes into consideration, this difference of 3 per thousand must be taken into account. In most cases this is not necessary, just as in most cases it is immaterial whether or not a set of weights is made according to exact grams, as

long as it is only made exact in its proportions. The Law of Faraday. Those substances which allow a movement of electricity through them, i.e. conductors of electricity, are divided into two classes. In the cases considered so far, the assumption has been made that the only form of energy into which the electric energy of the current is transformed is heat. Conductors which undergo no change other than that they get heated when the current passes through them are called conductors of the first class or metallic conductors. To this class belong the metals, their alloys, carbon, and certain compounds.

Now, there are many substances which permit a movement of electricity through them, but in so doing themselves suffer chemical change. To these belong aqueous solutions of acids, bases, and salts, as also salts in the molten state, and a few other compounds. Such substances are called conductors of the second class or electrolytes.

The movement of electricity in electrolytes takes place in such a manner that the components of the electrolyte move independently of each other. Hydrogen, the metals, and the metallic radicles move with the positive electricity, or, according to the usual designation, from places of higher to places of lower potential. The halogens, the acid radicles, and hydroxyl, on the other hand, travel with the negative electricity, or from places of lower to places of higher potential.

According to Faraday's nomenclature the components of electrolytes—that is, on the one hand the metals, hydrogen, &c., on the other hand the halogens, the acid radicles, &c.—are called ions. The first named, which travel down with the current, are called cations, those which travel up anions.

In a circuit consisting exclusively of an electrolyte, a motion of electricity can take place without the corresponding chemical action being apparent. But if a conductor of the first class directly touches an electrolyte, whenever there is a movement of electricity there is also a production of the ions at the surface of contact, and the chemical effects become apparent. As proposed by Faraday, the surfaces of metallic conductors, where they touch the electrolytes, are called electrodes. The surface at which the anions appear is called the anode, and that whereat the cations appear is called the cathode.

Faraday also formulated the general law to which all movement of electricity in electrolytes is subject. In every electrolyte the quantity of ion separated out is proportional to the quantity of electricity which has passed through, and the same quantity of electricity passing through different electrolytes separates quantities of the different ions that are in the ratio of the chemical equivalents of those ions. In these propositions it has been assumed for simplicity's sake that the ions are actually separated out at the electrodes, but this is not essential for the truth of Faraday's law. The following wording completely expresses the actual meaning of Faraday's law: Electricity can move in electrolytes only simultaneously with the ions, and so that chemically equivalent quantities of the different ions carry with them equal quantities of positive or of negative electricity.

We arrive at a perfectly appropriate conception of the nature of electrolytic conduction if we assume that each equivalent of the different ions possesses an equally large capacity for electricity, or, to use the words of Helmholtz, 'the electricity in electrolytes behaves as if if were divided into equal atoms.'

In applying Faraday's law, it is important that we should keep in mind the difference between the equivalent and the atom; the equivalent weight is equal to the atomic weight divided by the valency. Thus the same quantity of electricity travels with 107 g. of silver, $\frac{1}{3} \times 65.5$ g. of zinc, and $\frac{3}{3} \times 27$ g. of aluminium; also with 35.5 g. of chlorine there travels only half as much electricity as with 96 g. of SO₄. As an amplification of the illustration used above, we must assign to each atom as many equal capacities for electricity as the number of valencies it possesses. Perhaps this is the path by which in the future we may arrive at an elucidation of the nature of chemical valency.

The truth of Faraday's law was tested by its discoverer himself in many ways (Exp. Res. vii. ser.). He convinced himself that the same quantity of dilute sulphuric acid was always decomposed by the same current, whether the electrodes, or the E.M.F., were large or small. Nor did the strength of the acid exert any influ-Finally, he introduced into the same circuit dilute sulphuric acid and tin chloride, lead chloride, or lead borate, or he led the same current through different beakers with dilute sulphuric acid, using electrodes of different metals: in all cases he found his law verified. Later on, other investigators have tested whether part of the electricity is not perhaps conducted through electrolytes in the same way as through metals, but no trace of this has been discovered; in every case the electricity passed through has been strictly proportional to the quantity of ion separated out. And the law of equivalency has also proved itself so correct that it has been used lately as a means for determining equivalent

weights, and hence atomic weights.

The cases in which the same element exhibits different valencies according to circumstances are of special interest. Many such cases have been examined by Regnauld (A. Ch. [4] 11), who formed cells by taking platinum and different metals, along with the corresponding electrolytes, and then determined the quantities of metal dissolved when equal quantities of electricity had been allowed to pass. Thus it was found that mercury in dilute nitric acid has the equivalent 200, since it is changed into HgNO,, a compound of mono-valent mercury; in a solution of potassium cyanide it has the equivalent 100, because it becomes Hg(CN)2, in which the mercury is divalent. Similarly, copper in hydrochloric acid forms CuCl and has the equivalent 63.4; in nitric acid it forms Cu(NO₈)₂, and has the equivalent 31.7. Tin scarcely ever passes into solution except as a di-valent element; it is only in alkaline polysulphides and in a solution of caustic potash containing nitre that it acts as a tetravalent element. Tellurium also can be made to act as a di-valent and a tetra-valent element. It is known that the elements mentioned show quite different reactions in their solutions, according as they are present as mercurous or mercuric, as stannous or stannic compounds, &c. Hence in these cases the nature of the chemical reactions depends on the quantity of electricity with which the ions are combined.

This is the place to refer back to a misunderstanding concerning the law of Faraday into which Berzelius first fell, and which has recurred many times since. As has been mentioned, Berzelius saw in this law a contradiction to his theory, and tried to refute it by the following argument: if equal quantities of electricity were required for the decomposition of equivalent quantities of the most diverse compounds, it would follow that these could be decomposed by equal forces; this, however, is not possible, since the different substances are evidently held together by quite different forces of affinity. From the illustration given above it can be seen that, contrary to the assumption of Berzelius (which, however, was very excusable, considering the state of electrical science at that time), Faraday's law does not in any way deal with forces, but rather with quantities of electricity. Comparing electrical decomposition to the raising of water out of wells by means of buckets, Faraday's law asserts that in spite of the differences in the depths of the wells, the size of the buckets is always the same. We would fall into an error similar to that of Berzelius if we concluded from the equal size of the buckets that the work necessary for raising the water out of all the wells of different depths was the same. The illustration also shows directly wherein the differences in chemical work, corresponding to the different depths of the wells, assert themselves, namely not in the quantities of electricity required, but in the differences of potential.

Electrolytes and ions. It follows from the nature of an electrolyte that it must be a compound substance, since otherwise its parts or ions could not transport the electricity in both directions. Yet all compounds do not possess the power of conducting electricity. In particular, we do not know of any substance liquid at the ordinary temperature which is an electrolyte; amongst pure substances it is only the salts melted at a higher temperature which conduct at all appreciably.

On the other hand, all substances having the character of salts, acids, and bases possess electric conductivity when in aqueous solution. In these cases the water exerts a specific influence, since solutions of the same substances in alcohol conduct much less if at all; and solutions in ether, carbon disulphide, and similar solvents are

scarcely conductive.

There are two factors, therefore, which are necessary for the production of electric conductivity, i.e. the nature of the substance and the condition in which it is.

In electrolysis the molecule of the conducting compound appears to be split into two parts. There can be no doubt as to what these parts are with substances such as chloride of silver or potassium iodide; since they consist of two elements only, the metal must be one ion and the halogen the other. The question gets more doubtful with ternary compounds, such as nitric acid, sodium acetate NaCO2CH2, or ammonium chloride NH,Cl. Since, however, no important distinction manifests itself in the behaviour of these substances and those mentioned above, it is simplest and most obvious to assume that their ions correspond to those of the substances first named. Hence the ions of nitric acid are H and NO₂, those of sodium acetate Na and CO₂CH₃, and those of ammonium chloride NH, and Cl.

It is possible to examine this assumption on the basis of Faraday's law. In the electrolysis of potassium sulphate as much oxygen and hydrogen are obtained as from the electrolysis of dilute sulphuric acid by means of the same current, but in addition to this there is found at the anode an equivalent quantity of free sulphuric acid, at the cathode a corresponding quantity of free caustic potash. If we assume with Berzelius that the current has decomposed the potassium sulphate into potassium oxide and sulphur trioxide, K2O and SO3, which combine with the water to form 2KOH and H,SO,, we should have to assume further that the same current had in the same electrolyte simultaneously decomposed an equivalent quantity of water. But this is in contradiction to the law of Faraday. If we assume, however, that the ions are K2 and SO4, it is comprehensible that the potassium separated out should act on the water of solution, evolving hydrogen, according to the equation $2K + 2H_2O$ = 2KOH + H₂; in like manner the ion SO₄ acts on the water and forms sulphuric acid and free oxygen, according to the equation H2O + SO4 $= H_2 SO_4 + O$. With those metals which do not decompose water, such as zinc, copper, and silver, we actually obtain the metal only at the cathode; and similarly, solutions of chlorides, bromides, and iodides do not generally give free oxygen, but free halogen.

It is necessary, therefore, to distinguish carefully between the primary and the secondary effects of electrolysis. The primary effects consist in the separation of the ions at the electrodes. But since the ions do not continue to exist as such after having given up their electricity, they undergo further changes according to circumstances. The simplest case is that of the metallic ions, when these cannot act on the water; they simply become solid or liquid metals. So also the separated atoms of the halogens pass on, combining into gaseous chlorine, Cl₂, or into the molecules of the dissolved bromine or iodine.

The metals which decompose water can still be obtained in the metallic state when water is excluded or reduced to as small a quantity as is possible. A classical example of this is furnished by the electrolysis of slightly moistened potash, by means of which H. Davy discovered potassium. The electrolysis of molten salts, that of the chlorides especially, furnishes a means for preventing the secondary reaction of the metals; such electrolyses, which had first been used by Bunsen for scientific investigations, have obtained at the present time great technical importance in the production of magnesium and aluminium.

Complex cations of the type of ammonium, NH₄, are very unstable, and until recently it was doubtful whether the so-called ammonium amalgam, which is obtained on using a mercury cathode in the electrolysis of ammonium chloride, really contained the compound radicle NH₄. By measurements of another kind (viz. of the electromotive force) the existence of ammonium in the amalgam has now been placed beyond a doubt.

The compound anions as such are all of them very unstable. Many of them, such as the anions of the nitrates, phosphates, sulphates, &c., simply act on the water by taking up from it the hydrogen required to produce acids, and liberate the oxygen. The latter, which appears in the nascent state, is capable of bringing about powerful oxidising effects. If, for instance, lead or manganese salts are electrolysed, the oxygen acts on the dissolved metal and separates it at the anode as peroxide. In other cases the anion splits up into more stable compounds. Thus in the electrolysis of formates we obtain carbon dioxide and hydrogen from the ion HCOO, according to the equation 2HCOO = H2 + 2CO2. The acetates give carbon dioxide and ethane $2CH_3CO_2 = C_2H_6 + 2CO_2$, and similarly the salts of the higher fatty acids give the corresponding paraffins.

In some cases the secondary action can take place in another direction; in this manner the formates can produce—instead of hydrogen and carbonie acid—water, carbon dioxide, and carbon monoxide, as is shown in the following equation, $2HCO_2 = H_2O + CO + CO_2$. In most cases such secondary processes take place along with the primary actions, and the proportion between the two depends on external circumstances, temperature, concentration, current-density, &c.

In a few cases it is somewhat difficult to recognise the primary or secondary character of a product of electrolysis. On electrolysing potassium silver cyanide or sodium platinichloride, silvor or platinum separates at the cathode, so that it seems as if the silver cyanide or the platinichloride were decomposed primarily. This, however, would be an erroneous view, since the

ions of the salts are K and Ag(CN), and Na, and PtOl, respectively, and the heavy metals are only secondarily reduced from the solutions by the alkali metals. We can convince ourselves of this by using small electrodes and strong currents, when the salt near the cathode is soon used up, and potassium (or sodium), or, more correctly, hydrogen which has been formed secondarily, separates. More distinct proof of the nature of the ions is, however, obtained by studying the migration of the ions, to be discussed later.

So far the assumption has been made that the ions exert no action on the metal of the electrode; this, however, is not always correct. Only a few metals when used as anodes offer resistance to the chemical attack of the anions. If the metal can combine with the anion it forms the corresponding salt, which either dissolves in the water of solution, or, if insoluble, remains attached to the electrode and often interrupts the current. But also when used as cathodes the metals often combine with the cation; thus mercury forms amalgams, and the hydrogen separated at the cathode is occluded in greater or smaller quantities, especially by palladium and platinum, and also by iron. In other cases the cathode simply becomes coated with a layer of the corresponding metal, which separates more or less uniformly according to its nature and that of the electrolyte. These processes are used for coating substances which are conductors, or if non-conductors, whose surfaces at least have been made conductive; such processes have also been employed for galvanoplastic coating with gold, silver, copper, &c.

The constitution of electrolytes. It has already been remarked that all compounds are not electrolytes, and that some substances which do not conduct electrolytically by themselves obtain this power when dissolved in certain solvents, especially in water. From this it follows that electrolytes have a special constitution with which their special property is connected. The question as to what this constitution is has scarcely been raised yet, much less has it been answered. This is because most of the compounds known in the days of the electrochemical system of Berzelius were electrolytes, and later on, when many organic compounds, which are mostly non-electrolytes, were discovered, the electrochemical theory had been given up, and the interest in it had disappeared. The first investigator who proposed the question clearly and who, as far as he could, answered it—was Hittorf. He it was who established the proposition 'electrolytes are salts.' Under the name of 'salt' Hittorf comprised all those compounds which are capable of exchanging their consti-tuents within the shortest time. Hence along with the salts proper must be classed acids and

By enunciating this law, Hittorf was the first to draw attention to a most important relation which has become the starting-point for the due comprehension of the constitution of electrolytes. The power to conduct electricity and the power to exert chemical reactions are so closely parallel that both must be considered as the outcome of the same cause. This cause, however, has to be looked for in the constitution of the electrolytes themselves.

Electrolytic conduction is brought about by the positive and negative electricities moving through the conductor bound to their ponderable carriers, the ions; it is necessary, therefore, to assume a considerable mobility of the ions.

And, further, electrolytic conduction occurs in consequence of any difference in potential, however small. It is necessary, therefore, as was already pointed out by Clausius (P. 101, 338), that in every electrolyte there should be present a certain number of ions capable of moving freely, i.e. in every electrolyte a portion of the salt-like compound which produces the conduction must be partly split up into ions. Clausius left undecided how great this portion is, as he possessed no means to determine it. To the chemists who, in consequence of a peculiar shifting of judgment, considered electrolytes generally as specially stable compounds, he made the concession that this portion might be very small. In the imagery of the kinetic hypothesis he conceived that, owing to the collision of the molecules of the electrolyte with each other and with those of the solvent, one or other occasionally splits into its constituents, and so gives the necessary free ions.

It is now possible to urge similar considerations from the chemical side. While gaseous substances at the ordinary temperature react but seldom, acids, bases, and salts dissolved in water do so instantaneously. A mixture of oxygen and hydrogen does not form water until it has been heated to between 400° and 500°; a mixture of hydrochloric acid and potash, however, when in aqueous solution, passes so quickly into potassium chloride that it has been found impossible as yet to determine the time required for the change, and yet in the first case 68,000 cal. of heat are produced, while in the second the heat produced, or the energy liberated, is only 13,700 cal.—that is, five times less. In general, non-electrolytic solutions also react with each other extremely slowly. An example of this is afforded by the means which must be taken to hasten the reactions of non-electrolytes, and especially of organic compounds; this has to be done by heating the reacting mixtures, often under increased pressure. For the formation of acetic ether from a mixture of alcohol and acetic acid—that is, for a process intimately connected with that of the formation of salts—at least ten years are needed, at the temperature of the room, before it has approximately attained the end which under these conditions it can reach. From the chemical point of view, therefore, it is necessary to suppose that compounds which react instantaneously—that is, electrolytes—possess a special mobility of their parts or ions; and that it is the constituents of 'salts'-the term being used in the widest sense - which are the ultimate causes of electrolyses as well as of chemical reactions. These constituents of salts are, on the one hand, the metals, the metallic radicles, and hydrogen; on the other hand, the halogens, the acid radicles, and hydroxyl. It is possible, therefore, to apply to substances which react rapidly that is, to salts, as this term was used by Hittorf-the same kinetic considerations as were made use of by Clausius to show the presence of ions in electrolytes.

We are now confronted on both sides by the

question of how great is the proportion of free ions, or of decomposed molecules, in solution of electrolytes. The fact that a maximum reacting power exists for a given class of substances is in contradiction to the assumption that this proportion is inappreciably small. It has already been explained (compare the article Affinity) how the substances which have been most carefully examined from this point of view—the acids—exert their chemical reactions according to a definite co-efficient which is characteristic of each acid. This co-efficient—which can be determined, for instance, by means of the velocity of inversion of cane sugar-cannot be increased at will, and does not exceed a definite maximum value, which maximum is possessed by hydrochloric acid, nitric acid, benzenesulphonic acid, &c. All the conditions-such, for instance, as the introduction of oxygen, sulphur, or halogen into the molecule --which increase the velocity of reaction of weak acids are without any measurable effect on the strongest acids. From this it follows that the circumstance which is the cause of the velocity of reaction—that is, the unimpeded mobility of the ions—has in these substances already reached its highest degree, or is at least very near to it, since, in opposition to Clausius and to the views current till now, we must assume that in aqueous solutions of these strong acids it is not, perhaps, a few hundredths or thousandths of the molecules present which are split into their ions, but rather that by far the greater part of the electrolyte is split into ions.

A similar conclusion can be drawn from the degree in which electrolytes conduct the current. The more detailed consideration of this subject will be taken up presently. Here it will only be mentioned that the results of these investigations agree completely with those on the velocities of reactions.

A third entirely independent verification of the unexpected conclusion that aqueous solutions of the strong acids are nearly completely dissociated can be gained from a study of the properties of these solutions.

Only so much of van't Hoff's theory of solutions (Z. P. C. 1, 481) as is necessary for understanding the constitution of electrolytes will be explained here.

The fundamental idea of van't Hoff's theory is as follows. The physical properties of gases, and especially the relations between pressure. volume, and temperature, are practically independent of the special nature of the substance, while such an independence does not exist with liquid and solid substances. The cause of this phenomenon may be sought in the fact that the smallest parts or molecules of gases are always at such a distance apart that they are not capable of exerting an action on each other. In liquid and solid substances, however, the molecules are in close proximity, and exert, therefore, their specific reciprocal actions. But there is another condition in which the molecules of a substance are relatively distant from each other, namely, when the substance is present in a solvent in the state of a dilute solution.

We should expect, therefore, that in this state matter would be subject to laws analogous to those which hold for gases.

The first property characteristic of gases is

their power of extending uniformly through any given space. Solutions exhibit the same pecu-When the pure solvent is placed over the solution of any substance, the substance begins directly to enter into the solvent (the pure solvent playing the part of a vacuum), and the movement does not cease until, as with gases, the substance is uniformly distributed through the whole space. The only apparent difference is that in the case of gases this process is completed in a very short time, while with solutions months, and even years, are needed in order to bring it, practically speaking, to an end. This movement can be impeded by introducing between the solution and the pure solvent a partition which allows the passage of the latter but not of the former. Such a 'semi-permeable' partition can be produced by saturating a porous clay cell, for instance, with a solution of copper sulphate, washing the cell, and filling it with a solution of potassium ferrocyanide. A pp. of amorphous copper ferrocyanide is thus formed in the pores of the clay, and this pp. allows the passage of water through it, but does not allow various other substances to pass. In order to arrive at a conception of the cause of such a property we can imagine that the pp. of copper ferrocyanide acts as a filter which allows the passage of the smaller water molecules, while it retains the larger molecules of the dissolved substance.

W. Pfeffer, who has studied such cells very thoroughly, found that if the cells are closed a very considerable amount of pressure is produced in their interior. A 1 p.c. solution of sugar produces a pressure of more than 50 cm. of mercury; a solution of nitre of equal strength gives pressures of between three and four atmospheres. Pfeffer also found that the pressures increased, at constant temperature, to maximum values, which remained constant as long as the partition remained intact. These pressures proved to be proportional to the concentrations, and further, they increased with a rise of temperature.

According to van't Hoff, this pressure, the 'osmotic pressure,' is to be looked on as analogous to the gaseous pressure. It is possible to form a similar conception of its cause as is formed of the cause of gaseous pressure; the molecules of liquids, like the molecules of gases, possess a quantity of kinetic energy which is proportional to the temperature. The solvent-for instance, the water—can pass through the semi-permeable wall, and since it is present on both sides its pressure is the same on both. The substance dissolved, however, bombards the wall from one side only, and thus a pressure is set up. This view-which, however, is only hypothetical, and which can be accepted or rejected without the theory itself being called in question—has been objected to on the ground that if it were true a weak-walled vessel must needs be broken by a solution placed in it. But this assertion overlooks the important point that the osmotic pressure can exert itself only within the solution. If we imagine for simplicity's sake a drop of a solution floating in space, it is true that a pressure due to the substance dissolved will be exerted on the internal surface of the drop, which pressure may easily amount to 100 atmospheres. But to this pressure there is opposed the normal capillary pressure of the liquid, the magnitude of

which we do not yet know very exactly, but of which we can assert that it has to be measured by thousands of atmospheres (in the case of water it is approximately 20,000 atmospheres). Hence this pressure is under all circumstances quite sufficient to maintain equilibrium with the osmotic pressure, and the only effect of the latter is to slightly increase the volume of the solution.

It has been shown by van't Hoff that the laws of osmotic pressure exactly coincide with those of gases. Boyle's law enunciates that pressure and volume are inversely proportional. Pfeffer's law says that pressure and concentration (or density) are proportional; this agrees with Boyle's law. Further, the law of Gay-Lussac tells us that at constant volume (or at constant density) the pressure of a gas increases with the temperature, and that for each degree it does so by $\frac{1}{278}$ of the value at 0°. A similar thing has been proved by van't Hoff from the numbers obtained by Pfeffer. And finally for gases we have the law of Avogadro, that, temperature and volume being the same, equimolecular quantities of different gases exert the same pressure. Taking the molecular weights in grams, and taking for the common volume 1 litre, then at 0° this pressure is equal to 22:37 atmospheres. Now, it has been observed by Pfeffer that the osmotic pressure of a one per cent. solution of cane sugar, which therefore contains 10 g. in one litre, is 0.649 atmospheres at 0°. The molecular weight of sugar $C_{12}H_{22}O_{11}$ is 842; a solution containing this quantity in grams in one litre would exert a pressure $\frac{342}{10}$ times as great—that is, a pressure of 22.2 atmospheres. This that is, a pressure of 22.2 atmospheres. number agrees within the limits of experimental error with that which holds good for gases; and hence Avogadro's law holds also for solutions, or, more correctly, so far, for solutions of cane sugar.

The question whether the law holds for other dissolved substances must be answered in the affirmative. It is true that only very few measurements of osmotic pressures have been made, but these phenomena are so closely connected with others to be considered shortly, that it has been possible to place the general validity of the law beyond doubt.

Let us imagine a solution poured into a vessel of the form of an inverted funnel, the opening of which is closed by a semi-permeable wall in contact with which is a quantity of the pure solvent. Then the solution will rise in the tube, because of the greater internal pressure, and equilibrium will be established only when the pressure of the liquid column in the tube has become equal to the osmotic pressure. Let this arrangement be covered by a bell-jar. from which the air has been completely expelled; the solution will then be surrounded by the vapour of the solvent only. But the pressure of the vapour is slightly smaller in the upper part of the bell-jar, at the level of the solution, than below at the level of the pure solvent, and it is smaller by an amount which is equal to the pressure of the weight of the vapour present between the two levels. Moreover, it is by this amount that the vapour-pressure of the solution must fall short of that of the solvent, because, if this were not the case, liquid would either continually evaporate, or condense, at the surface of the solution; in consequence of the osmotic pressure, the changes in level produced would always equalise themselves again, and we should have a perpetuum mobile, which is impossible.

It follows, therefore, that in general the vapour-pressure of a liquid must decrease when a substance is dissolved in it, and that this takes place according to laws which run parallel with those of osmotic pressure.

This theoretical conclusion is in accordance It has been found by with observation. Wüllner (P. 103, 529) that, at a constant temperature, the decrease of the vapour-pressure of an aqueous solution is proportional to the concentration of the solution. Babo on his part had established that the ratio between the vapourpressures of the pure solvent and the solution is independent of the temperature. Both results have been confirmed, on the whole, by later investigations. Raoult especially has occupied himself with this question; he has shown that this behaviour is general, and he has also found that equimolecular weights of different substances dissolved in the same solvent produce equal diminutions of the vapour-pressure, and finally that on using equimolecular quantities of different solvents the relative diminution in the vapour-pressures is the same. If f stands for the vapour-pressure of the solvent, f' for that of the solution, and if further N stands for the number of molecules of the solvent (in gram-units), n for the number of molecules of the substance dissolved, the general expression holds

$$\frac{f-f'}{f} = \frac{n}{N+n}.$$

While Raoult discovered this law empirically, van't Hoff has deduced it theoretically from the laws of osmotic pressure (Z. P. C. 1, 494).

This formula can be used for determining the molecular weight of a substance in solution. Let p grams of a substance with unknown molecular weight m be contained in 100 g. of the solvent, the molecular weight of which is M; then $\frac{p}{100}$ M = grams of substance dissolved in one molecular weight of the solvent; let this = s; then putting $n = \frac{s}{m}$, and N = 1, and solving the

equation for m, it follows that $m = \frac{sf'}{f - f''}$

Finally, a third set of phenomena is known, governed by similar laws. It has long been known that salt water freezes at a lower temperature than pure water. As far back as 1788 Blagden (T. 1788) established the fact that the lowering of the freezing-point of a solution of salt in water is proportional to the amount of salt dissolved. Later on, Rudorff (P. 114, 63) rediscovered this fact, which had been forgotten, and de Coppet found that equimolecular solutions of salts having analogous constitutions exhibited equal lowerings of the freezing-points (A. Ch. [4] 23, 866). Raoult discovered that the same law held for the most diverse solvents. This law states that equimolecular quantities of whatever substances we take, added to a constant quantity of the solvent, lower the freezing-point by the same amount (C.R. 1882). Raoult thought for a time that the lowering of freezing-point was the same for equimolecular quantities of different

solvents, but this proved to be erroneous. Van't Hoff has deduced this important relation from the laws of osmotic pressure, and his formula agrees well with experiment (Z. P. C. 1, 481).

We can convince ourselves of the necessity of these relations by an argument similar to that used in order to prove the connection between osmotic pressure and change of vapourpressure. As the mechanical theory of heat tells us, below 0° the vapour-pressure of ice decreases more rapidly than that of over-cooled water. A temperature must therefore exist whereat a saltsolution, which has a vapour-pressure less than that of water, exhibits the same vapour-pressure as ice. This is the only temperature at which the solution can exist side by side with ice, as otherwise a perpetuum mobile would be possible. Since, therefore, this temperature, which is the freezing-point of the solution, is in inseparable connection with its vapour-pressure, the laws holding for the one must also hold for the other, and from this the laws already stated follow directly. Formulating these laws, we get $\Delta = r - \frac{p}{nl}$

where Δ is the lowering of the freezing-point, lthe weight of the solvent, p that of the substance dissolved, and m the molecular weight of the substance, while r is a constant dependent on the nature of the liquid. From the formula it

follows: $m = \frac{rp}{\Delta l}$. Having, therefore, once for

all determined the constant for a solvent, it is possible to find the molecular weight of a substance by determining the lowering of the freezing-point of its solution in that solvent. Van't Hoff has shown that the constant can be deduced thermo-dynamically from the heat of fusion of the substance.

If we now apply these methods for determining the molecular weights of dissolved substances to solutions of electrolytes, we find that the molecular weights appear smaller, and hence the number of molecules in solution are greater, than correspond with the formulæ. In many cases the amount of this deviation is very considerable, and its existence at first proved a great obstacle in the way of van't Hoff's theory of solutions. To Arrhenius (Z. P. C. 1) we owe the means of accounting for the deviation. The views propounded by this investigator have shed such a flood of light over difficult questions in chemistry and physics that we must consider the work of Arrhenius to be one of the most important advances ever made in this and in allied domains.

Arrhenius' idea consists in ascribing the deviations of the observed from the calculated molecular weights, to a dissociation of the electrolytes into their ions. From the magnitude of this deviation a conclusion can be drawn as to the number of molecules which are dissociated; the problem which was left unsolved by Clausius thus receives its solution.

Reserving the numerical proofs of the truth of this view till later on, it is proposed to deal first with some possible objections and diffi-

culties of a general nature.

The application of the hypothesis leads to the conclusion that in moderately dilute solutions of strong acids and bases, and also of normal salts, as much as 80 to 90 per cent. of the substance in solution is dissociated into its ions. Hence a solution of potassium chloride contains little else than ions of chlorine and potassium. How is it that the chlorine does not escape into the air as a greenish yellow gas, and the potassium does not act on the water?

The answer lies in the consideration that the chlorine which we know as a greenish yellow gas is the electrically neutral molecule Cl₂, and does not consist of the separate atoms of Cl which are charged with a large amount of negative electricity. The potassium again is not present as the compact metal, but in the form of strongly positively charged ions. Considering the great differences exhibited by allotropic forms of the same element (oxygen and ozone, red and yellow phosphorus, &c.), we cannot wonder that in these cases also considerable differences appear.

And it is further possible to produce solutions in which such ions—as, for instance, potassium-are present in excess, and therefore certainly in the free state. For this purpose, let us imagine two vessels filled with potassium chloride solution and placed on an insulating stand, and then connected conductively by means of a syphon filled with the solution. Let us now bring near to the one vessel a body charged with negative electricity. The vessel becomes charged positively by induction, and an equivalent quantity of negative electricity moves through the syphon into the second vessel. If we now remove the syphon, and then the body charged, the first vessel remains charged positively, the second vessel negatively. So far the experiment contains nothing that is new; it is the elementary experiment used for proving the inductive effect of electricity. If, however, we pay attention to the fact that, according to the law of Faraday, electricity can move in electrolytes only simultaneously with the ions, the conclusion is inevitable that an excess of potassium ions must be present in the vessel charged positively, and the same excess of chlorine ions must be present in the vessel charged negatively. These excesses of ions are maintained as long as the electric charge of the vessels exists. On discharging the vessels the ions give up their electricity, and the substances appear at the point of discharge endowed with their usual properties.

Owing to the magnitude of the quantities of electricity contained in the ions, the question may be raised whether the experiment which has just been indicated really can be carried out. Ostwald and Nernst have shown (Z. P. C. 3) that this is actually possible. By means of a mercury electrode contained in a capillary tube, such small quantities of hydrogen can be observed that the experiment becomes quite feasible. Since under ordinary conditions one gram of hydrogen occupies a space of about 12,000 c.c., a small bubble of 0·1 mm. diameter, which one can still see with the naked eye, has a weight of only 10⁻¹⁰ g., and one of 0·01 mm. diameter, which can be seen

¹ From the number, already given, of 96,540 coulombs, which are combined with 1 g. of hydrogen and which are also present in one litre of normal potassium chloride solution—as positive electricity on the potassium, or negative electricity on the chlorine—it is possible to calculate that this quantity of electricity would suffice to charge a sphere of an approximate diameter of 10¹⁴ cm.—which, therefore, would be larger than the whole solar system—to a potential of one volt.

under a microscope, has a weight of only 10⁻¹⁸ g. This is as small a quantity of matter as can be recognised in any way, and the electricity combined with it can be detected in an apparatus of very moderate dimensions.

Thus the experiment described above is not only conceivable, but feasible; and thus it is proved that free ions, such as those of potassium, can be present in aqueous solutions without acting on the water.

The assumption that some of the molecules of electrolytes are split into ions in solutions explains many properties of these solutions. In the following paragraph we will assume for simplicity's sake that the splitting up is complete—an assumption which is nearly correct in the cases of salts, strong acids, and bases; the modifications which must be introduced into these considerations by taking into account the portions not split up, will be considered afterwards.

Let us consider, first, the chemical reactions of electrolytes. As is well known, certain reactions are used in analytical chemistry for indicating definite substances. These reactions must take place quickly, as otherwise they would be of no practical use; hence they must be reactions between ions. If asked what it is that the ordinary analytical reactions allow us to recognise, we are inclined to answer, the different chemical elements. But this is not at all appro priate. A solution of a salt of silver is called a reagent for chlorine, and in fact by its help chlorine can be detected in metallic chlorides. But the chlorine in potassium chlorate or in monochloracetic acid cannot be detected by means of silver solution. In the same way chlorides indicate silver; but the silver in potassium silver cyanide cannot be recognised by means of chlorides. The different oxyacids of sulphur also give quite different reactions, though they all contain the same elements; similarly, the reactions of iron, copper, mercury, tin, &c., are quite different, according to the degree of oxidation of the metal.

These examples prove that it is not the elements which are indicated by analytical reactions. A more careful consideration shows that since it is the ions which react, the ions are indicated by the analytical tests.

This simple proposition removes all the difficulties which confronted us. If a solution of silver is a reagent for chlorine ions, naturally it cannot indicate the chlorine in potassium chlorate, whose ions are K and ClO,; nor in monochloracetic acid, whose ions are H and CH2ClCO2. And, conversely, silver is indicated by chlorides only where it appears as the ion; but the ions of potassium silver cyanide are K and Ag(CN)₂. The behaviour of chloroplatinic acid H₂PtCl₆ towards silver solutions is specially remarkable, a behaviour about which an incredible confusion has reigned in chemistry, because it has always been believed that its chlorine must be precipi-The tated by silver solution as silver chloride. matter was first cleared up by Jörgensen (J. pr. [2] 16), who found that no silver chloride is formed at all, but rather the silver salt of chloroplatinic acid Ag2PtCle; sodium platinichloride is completely precipitated by two equivalents of silver solution only, instead of by six, which correspond to the proportion of chlorine.

Thus the so-called anomalies of analytical reactions are fully explained. It is further explained why, for instance, all the different sulphates give the same reaction with barium chloride. The nature of the metal is of no account, because the reacting ion SO₄ is not combined with the metal at all, but exists by itself.

It is a remarkable fact that similarly constituted ions give, under certain conditions, different reactions. According to this view, the iron in a solution of FeCl2, and also in a solution of FeCl₃, is contained as an ion; yet the two act differently. In like manner, solutions of K₄Fe(CN)₆, as well as those of K₅Fe(CN)₆, contain the same negative ion Fe(CN), and these two substances give quite different reactions. The explanation lies in recognising that these similarly composed ions are endowed with different quantities of electricity according to their valency. Thus iron appears divalent and trivalent, mercury and copper monovalent and divalent, the atomic group Fe(CN), tetravalent and trivalent, in that they contain corresponding numbers of positive or negative electric units or 'atoms.'

If the theory developed so far is correct, then all properties of salt solutions must be of an additive nature with regard to the two ionsi.e. their properties must be made up of two parts, one of which depends only on the positive ion, and the other only on the negative ion. This fact was recognised before it found an explanation in terms of the hypothesis of the independent existence of ions in solutions. Arrhenius drew attention to this fact (Z. P. C. 1)when he was establishing his theory. examples will make the point clearer.

In 1874 Valson enunciated a law concerning the specific gravities of salt solutions (C. R. 73), which he called the law of the moduli. By means of it the specific gravity of a 'normal solution—that is, of a solution containing one equivalent of the salt in grams in one litre-of any salt can be found by adding to the specific gravity of a solution of ammonium chloride (chosen as the standard) two values, one of which depends on the metal only, and the other on the acid radicle only. According to Valson, these moduliare (NH₄ClAq of specific gravity 1015 being taken as the standard):— K = 30, Na = 25, $\frac{1}{2}$ Ca = 26, $\frac{1}{2}$ Mg = 20, $\frac{1}{2}$ Sr = 55, $\frac{1}{2}$ Ba = 73, $\frac{1}{2}$ Mn = 37, $\frac{1}{2}$ Fe = 37, $\frac{1}{2}$ Cn = 41, $\frac{1}{2}$ Cu = 42, $\frac{1}{2}$ Cr = 61, $\frac{1}{2}$ Pb = 108, Ag = 105; Br = 84, I = 64, $\frac{1}{2}$ SO₄ = 20, NO₃ = 15, $\frac{1}{2}$ CO₅ = 14, HCO₅ = 16. To these are added the value zero for NH and Cl. for NH, and Cl. These values allow us to calculate the specific gravities of solutions of $14 \times 7 = 98$ salts, and the calculations agree very well with experiment. Later on (1883), the law was confirmed by C. Bender (W. 20, 560), and by Nicol (P. M. [5] 18, 179), and was also extended by the first named.

A similar law was established by Gladstone (Pr. 16, 439) for the refraction-equivalents of salts. Gladstone expressed his generalisation in the form that the difference between the refraction-equivalents of the salts of two metals with the same acid is always the same, and is independent of the composition of the acid. The following table shows how the values agree in the cases of salts of potassium and sodium with acids of different compositions:-

		Potassium	Sodium	Difference
Chloride		18.44	15.11	3.3
Bromide		25.34	21.70	3.6
Iodide .		35.33	31.59	3.7
Nitrate .		21.80	18.66	3⋅1
Hydrate .		12.82	9.21	3.6
Alcoholate .		27.68	24.28	3.4
Formate .		19.93	16.03	8.9
Acetate		27.65	24.03	3.6
Tartrate .		57.60	50.39	8.6
Carbonate .		34.93	28.55	2×3.2
Bichromate		79.9	72.9	2 × 3·5
Hypophosp	hite .	26.94	20.93	2 × 3·0

A similar law exists for the salts of strong acids, but with the weak acids the differences are much greater. We are now in a position to account for this discrepancy, which was left unexplained by Gladstone. Since the weak acids are only very partially split into their ions, it cannot be expected that hydrogen as an ion should have the same refraction-equivalent as it has in the undecomposed compound.

Gladstone also enunciated the general law that the colour of the solutions of salts whose acid or whose metal form coloured compounds is independent of that of the other constituent (P. M. [4] 14, 418). If we consider how dependent colour is on constitution in other cases, this fact must be considered as a specially weighty proof of the mutual independence of the ions.

The knowledge of the fact that the characteristic colours of salts are dependent on the nature of the coloured ion only, and are independent of that of the other ion, has existed for a very long time, almost, as one might say, unconsciously. In green liquids we should always expect nickel, in red ones cobalt, in pale blue ones copper, &c. Some exceptions to this can easily be explained. One such exception is that most copper salts show the blue colour of the copper ions, while concentrated solutions of copper chloride appear green. The cause of this is that in concentrated solutions a considerable number of undecomposed molecules CuCl₂ are present, which molecules are coloured a deep yellow, as can be seen in dry copper chloride, which looks almost reddish brown. This yellow colour mixes with the blue colour of the copper ions, and produces the green colouration. From the fact that as the temperature is raised the green colour becomes more intense, it must be concluded that with rise of temperature more molecules CuCl, form at the cost of the ions (v. Ostwald, Z. P. C. 9, 579).

Another additive property has been found by Marignac (A. Ch. [5] 8, 410) in the specific heats of salt solutions. This property is, however, partly hidden by small deviations which. though capable of explanation, require the knowledge of some quantities the measurement of

which has not yet been undertaken.

And, further, Ostwald (J. pr. [2] 18, 853) has proved such relations to hold in a fairly extensive manner for the volume-changes accompanying neutralisation. On saturating solutions of potash, soda, and ammonia by the same acid the total volume changes in a very different manner. These volume-changes are, however, governed by the law that on saturating two bases by the same acid the difference of the volume-changes is independent of the nature of the acid; and in like manner the difference in the volume-change on saturating two different acids by the same base is uninfluenced by the nature of the base. Hence the volume-change always depends on two constants, one of which is determined by the acid only, and the other by the base only; the specific nature of the salt formed has no influence. The following table shows these relations:—

	Potassium	Sodium	Ammonia
Nitric acid	20.0	19.8	- 6.4
Hydrochloric acid .	19.5	19.2	- 6.6
Hydrobromic acid .	19.6	19.3	- 6.6
Hydriodic acid .	19.8	19.6	- 6· 4
Formic acid	12.4	12.1	-13.6
Acetic acid	9.5	9.3	-16.3
Monochloracetic acid	10.9	10.6	-15.1
Dichloracetic acid.	13.0	12.7	- 13.0
Trichloracetic acid.	17.4	17.1	- 8.7
Isopropionic acid .	7.8	7.7	-17.8
Butyric acid	7.0	6.8	-18.6
Isobutyric acid .	6.8	6.1	-19.3
Glycollic acid	9.6	9.8	- 16.5
Lactic acid	8.3	8.1	- 17.7
Sulphuric acid .	11.9	11.5	-14.4
Oxalic acid	9.9	9.7	-16.1
Succinic acid	8.2	7.9	-17.6
Malic acid	8.6	8.5	-17.4
Tartaric acid	9.4	9.2	-17.0

The differences between the corresponding members of two horizontal or of two vertical columns in this table are seen to be constant within the limits of error. The all but complete equality of the values for the four first-named acids, which are almost completely split up into their ions, is remarkable.

The most striking case of the additive properties of salt solutions is, however, given by their thermal relations. Hess long ago established the law of the thermo-neutrality of salts; this law declares that there is no thermal effect on mixing the solutions of two salts provided that everything remains in solution. This fact is difficult to understand as long as we assume the existence in an undecomposed state of normal salts in solutions, since the mutual changes of the salts would have to proceed without any energy-change, and this is a behaviour without analogy. If we assume, however, that the solutions of the salts contain chiefly the ions of the salts, the fact becomes explicable, because on mixing the solutions, the ions remain unchanged, they do not influence each other, and hence there is nothing to cause thermal effects.

More remarkable still is the explanation which the theory gives concerning the heats of neutralisation of acids and bases. At first sight it seems as if no action could take place between such compounds, if we consider the strong acids and bases to be almost wholly separated into ions. Because if the solution of the acid, of the base, and of the salt contains ions only, no change ought to occur when the acid and the base are mixed. If, however, we consider the reaction more carefully, we arrive at different results. For instance, let there be given soda

and hydrochloric acid; the reaction will be

$$N_{a} + OH + H + Cl = N_{a} + Cl + H_{2}O$$

The ions sodium and chlorine remain unchanged; but since water conducts very badly, only a very slight separation of H_2O into the ions OH and H can occur, hence when the ions OH and H meet they must combine to form H_2O .

Hence the process of neutralisation in aqueous solutions is not a combination of the constituents of the salt, but only a combination of the constituents of water.

It is true that this remarkable result is somewhat contradictory of the usual views on the subject, but it is in complete agreement with the facts. If the process of neutralisation between acids and bases decomposed into their ions really consists only in a formation of water from hydrogen and hydroxyl, then its concomitant phenomena must be independent of the nature of the acids and the bases.

The most important of these concomitant phenomena is the heat of neutralisation. This value has been determined by different observers, with special exactness by Thomsen (Th. 1). A number of measurements are collected in the following table, calculated for one equivalent; the unit used is the rational calorie, K=100 gram-units of heat:—

	NaOH	кон	Lion	½BaO₂H,	CaO,H,
HCl	137	137	137	139	139
HBr .	137	137		_	
HI	137	136			
HNO	137	138		140	139
HClO.	138	138		140	
HBrO.	138	138		_	_
HIO	138	138			
HClÖ, .	141	143	_	_	
H2S2O6 .	135	_	_	139	
H2PtCl6.	136	-	_	_	-

And, similarly, tetramethylammonium hydroxide, platinodiamine hydroxide, and triethylsulphine hydroxide give with hydrochloric acid 138, 137, and 137 K respectively.

From this table, which could easily be enlarged, it follows that the heat of neutralisation of the strong acids and bases is a constant, and is approximately equal to 13,700 cal. This number must be looked on as very nearly equal to the heat of formation of water from the ions

H and OH. (A more exact calculation gives 13,500 cal.) The small deviations which are still present originate in the fact that the splitting up of the acids and bases referred to, though very considerable, is yet not quite complete; it is in this undecomposed remnant that the individual nature of the two substances still asserts itself a little.

The conditions become quite changed when we consider the weak acids and bases. Here the constancy of the heat of neutralisation ceases completely, and values are observed which fluctuate between 163K and 10K. In these cases the quantities of heat which are concerned in the splitting up of the acid and the base into their ions must be taken into account, besides the heat of formation of water from its ions, and the beat.

of neutralisation may be represented by an expression of the form $N=13,500+\Delta+B$, where A + B is the quantity of heat necessary for decomposing the acid and the base into ions. As experiment has shown that the salts when in dilute solution are uniformly and nearly completely decomposed, their heats of decomposition need not be taken into account meanwhile. This formula also expresses a law which, discovered some time back by Favre and Silbermann (A. Ch. [3] 37, 486), has been confirmed by the younger school of thermo-chemists. This law is, that the difference between the heats of neutralisation of any two bases is always the same whatever acid is taken, and that likewise the difference between the heats of neutralisation of any two acids is independent of the base. This is only another expression for the proposition that the acid and the base each contributes towards the heat of neutralisation a definite portion, which is independent of the nature of the substance with which the acid or the base reacts.

Very similar facts may be noticed concerning the other phenomena which accompany neutralisation. A table has already been given (p 189) showing that the same relations hold good for the changes of volume that accompany neutralisation; the strong acids and bases give, on neutralisation, very nearly the same change of volume, i.e. 20 c.c. per litre of normal solutions of acid and base. The same holds good for the changes in the refractive indices.

The electrical conductivity of elec-

trolytes.—Ohm's law $I = \frac{E}{R}$ gives a definition

of R, the resistance of a conductor. It has been shown that the resistance is dependent on the nature of the conductor, and on its dimensions, in that it varies directly as the length and inversely as the cross-section. It has been agreed to call the resistance of a conductor of crosssection 1 sq. millim. and length 100 centim. its specific resistance; this depends on the nature of the material of which the conductor consists, and on its temperature.

If we define a quantity $C = \frac{1}{R}$, and call it the

conductivity, Ohm's formula becomes I = EC. We may best regard C as the power of the conductor to allow the electricity to pass through, or the power to transport the electricity. In the illustration of a current of water in a tube, C would be comparable to the cross-section of the tube. The conductivity is the reciprocal of the resistance. In calculating the strength of currents it is more convenient to work with resistances; if these are connected end to end, the total resistance is the sum of the individual resistances; but if the resistances are connected side by side, the conductivities must be summed up. electro-chemical purposes the idea of conductivity has always proved more useful than that of resistance.

Measurements of resistances or of conductivities are made according to different methods, all of which rest on the application of the formula above given. For instance, we can measure the current strength, I, using the same electromotive force E (from a constant cell, as, for in stance, a Daniell), there being included in the liquid taken out. The method suffers from the

circuit at one time the resistance R, which is to be determined, and at another time a resistance of known magnitude Ro. We then get the equations

$$I_{_{1}}=\frac{E}{R_{_{1}}}\text{ and }I_{_{0}}=\frac{E}{R_{_{0}}}\text{, and from these }R_{_{1}}=R_{_{0}}\frac{I_{_{1}}}{I_{_{0}}}\text{.}\quad\text{It}$$

is not necessary to discuss here the various methods of measuring resistances.

As mentioned before, the unit of resistance used is the ohm-i.e. the resistance of a mercury column of 1 sq. mm. section, and 106 or 106.3 cm. length at 0°. The unit of conductivity is given by the same quantity; since this represents the reciprocal value of the resistance, it has been proposed to designate it by Mho, which is the name Ohm reversed, and for which Mo might be put shortly.

While so far the different powers for conducting electricity possessed by the metals have remained more a question of practical than of theoretical importance, the study of the electrical conductivity of electrolytes has led to most important and suggestive views as to their nature. These have only been arrived at quite lately, chiefly because until recently there existed no convenient and good method for determining the conductivities of electrolytes in solution.

The difficulty to be surmounted consists in the fact that the current must be conducted through the electrolyte by means of electrodes, and that these get coated with the ions which separate-i.e. they become polarised. In consequence of this the electrodes become the seat of new electromotive forces, the magnitude of which is variable, and therefore not capable of being determined exactly. One method only has been brought forward as yet in which this difficulty is completely obviated; it is due to Guthrie and Boys (P. M. 1880. 328), and consists in allowing a system of strong magnets to rotate rapidly round the axis of a cylindrical vessel filled with the electrolyte and suspended by a long thin wire. By this means currents are produced in the electrolyte (as would be produced in every conductor), the electrodynamic effect of which tends to rotate the vessel in the same direction. The strength of these currents is proportional to the conductivity of the liquid, and the vessel is rotated until the torsion of the suspending wire becomes sufficient to maintain equilibrium against this rotatory effect. We are therefore justified in putting the deviation as proportional to the conductivity, the velocity of the magnets remaining the same. As the currents produced in the electrolyte occur exclusively within it, no polarisation is possible. The method has not been applied, because the apparatus required is expensive and difficult to handle.

Becquerel and Horsford tried to bring up polarisation to its highest value by using strong currents. In this method the current is led through the electrolyte, then through an adjustable resistance, and finally through an instrument for measuring currents (galvanometer, &c.), and its strength is determined. The distance between the electrodes is next decreased by an accurately measured amount, and resistances are inserted until the strength of the current has regained its value; then the resistance inserted is equal to that possessed by the portion of the lisadvantage that it is difficult to keep the electromotive force of polarisation constant as presupposed; and, further, the liquid is changed at the electrodes by the use of the stronger currents required, so that what we measure is the resistance of the changed, and not that of the original, liquid.

By using electrodes of zinc, which in solutions of zinc salts give no polarisation, Beetz (P. 117, 1) was able to determine the conductivities of such solutions by the methods used for metals. Paalzoff showed (P. 136, 489) that the method can be applied to other solutions also if we bring the two zinc electrodes into two separate vessels, and establish conduction through a syphon filled with the liquid to be investigated. The process is based on the fact that no polarisation is produced when a current is passed through the surface of contact of different liquids.

But this method is again far surpassed in convenience by one given by F. Kohlrausch (W. 11, 653), in which polarisation is made harmless by using alternating currents. Such currents, in which equal quantities of electricity flow alternately in opposite directions, are produced most easily by means of an ordinary induction apparatus. It is true that by such currents the electrodes are also polarised, but the polarisations occur rapidly in opposite directions, so that the polarisation produced by the first current-impact increases the strength of the second current-impact which has the opposite direction, and the polarisation resulting from the latter adds itself on to the third current-impact, and Kohlrausch has shown that by using platinum electrodes of ten sq. cm. surface coated with platinum black, the effect of polarisation may be reduced to less than 0.1 p.c. of the resistance. Kohlrausch uses for his method Wheatstone's bridge in Kirchhoff's arrangement. For details reference is made to Kohlrausch's memoir.1

The definition of specific conductivity as given above, and as employed in physics, is not of practical use in considering electrical conductivity from the point of view of the chemist. Since, according to the law of Faraday, the ions alone conduct, and since each ion carries the same quantity of electricity, it is evident that the conductivities of different solutions must be referred to the same number of ions in order to obtain comparable numbers. That we may form a conception of equivalent conductivity following from this, let us imagine two electrodes of indefinite extension placed at a distance of 1 cm. apart, and let there be brought between them so much of a solution as contains the equivalent weight in grams of the dissolved electrolyte, the conductivity of such a system is the equivalent conductivity of the electrolyte. In order to find the relation between the specific and the equivalent conductivity, we bear in mind that the length of the solution of the electrolyte is 106 times less than 106 cm., which was the length of the conductor assumed in the definition of specific conductivity. When the solution is normal-i.e. contains one g. equivalent per litre—the cross-section is 1,000 sq. cm.; when the same weight is dissolved in v litres it is 1,000v

¹ A detailed description of the method for carrying out such measurements is to be found in Z. P. C. 2, 565.

sq. cm.; this is 100,000v times greater than the cross-section for the specific conductivity, which is 01 sq. cm. Hence from the specific conductivity l we obtain the equivalent conductivity l by the equation l=1.06 × 10.1lv. It should be remembered that the specific conductivity of an electrolyte is not usually referred to ohms, but to the unit of Siemens, formerly commonly employed, which is equal to the resistance of a column of mercury 100 cm. in length; hence instead of 106 we must put only 100, and the equivalent conductivity becomes l=10.1lv.

In addition to the equivalent conductivity, we can calculate the molecular conductivity, which is as many times greater than the equivalent conductivity as the molecular weight contains the equivalent weight. So the molecular conductivity of sulphuric acid is twice as great as the equivalent conductivity, that of aluminium chloride three times as great.

For the general facts which have been established regarding equivalent conductivities we are specially indebted to Kohlrausch, and to several investigators who succeeded him. The most simple relations are exhibited by dilute solutions in which v is greater than 1 lit., and these will be considered first.

1. The equivalent conductivities of normal salts are of the same order of magnitude, but are not the same (Kohlrausch).

2. The conductivities of all salts increase slowly with increase of dilution, and generally reach a maximum value, which is not surpassed after dilutions of 20,000 to 50,000 litres per gram-equivalent (Kohlrausch).

3. The increase of conductivity is least for salts which consist of two monovalent ions; it is about twice as great for salts which contain one divalent and one monovalent ion; and about four times as great for salts whose ions are both divalent (Ostwald).

4. The equivalent conductivities of equally concentrated solutions of the most different salts can be represented as the sum of two constants, one of which is determined solely by the positive ion, the other solely by the negative ion (Kohlrausch). This law agrees the better with experience the more dilute the solutions are.

In order to give an illustration of these laws, I append the equivalent conductivities of several normal salts as measured by Kohlrausch. The values are for 18°.

Dilution	KCl	NaCl	LiCl	∄BaCl_	ŁK₂SO.	½MgSO.
12	91 9	69 5	59 1	65 8	67·2	27·0
102	104 7	86 5	77.5	86 1	89 7	47·4
1002	114 7	96 2	87 5	100 6	109 8	71·5
1,0002	119 3	100 8	92.1	109 2	120 7	93·5
10,0002	120 9	102 9	94 3	112 6	124 9	103·4
50,0007	121 7	102 8	93 5	114 4	126·6	105·2
100,0002	121 6	102 4	96 5	114 2	127·5	105·6

If we attempt to form a picture of the processes occurring in electrolytic conduction, we must first bear in mind that, according to the statements of Kirchhoff (P. 78, 506), free electricity must be present along the surface of each current-path; and the fall of this potential causes the motion of the electricity inside the conductor. Hittorf pointed out that the theories of electrolysis hitherto held could not satisfy this requirement. Assuming the presence of free ions,

it is evident at once that the surface charge is formed by these. Owing to the fall of potential which exists, the positive electricity is now driven to the one side, the negative to the other. Both are bound to the ions; which are, therefore, moved in the same way by forces that are of equal magnitudes on both sides.

The velocities acquired by the ions under the mfluence of these equal and opposite forces will not generally be the same, since it cannot be assumed that the resistances to motion encountered by the ions in the solution will be the same. But in dilute solutions, anyhow, the nature of the second ion, travelling in the opposite direction, will have no influence on the velocity of any definite ion—such as, for instance, Cl. Hence the conductivities of salts, when referred to equal quantities of ions, can be represented as the sum of the migration-velocities of the positive and negative ions. Calling these migration-velocities u and v, we get

 $\lambda = u + v$.

But this is exactly the relation discovered by Kohlrausch (p. 191), to whom also the argument just given is due.

Viewed thus, the phenomena of electrical conductivity are brought into connection with another group of facts, the comprehension of which had presented great difficulties to the older naturalists. These are the phenomena of the 'migration of the ions.' It has been mentioned that Daniell observed the corresponding manifestations, but they were not explained till Hittorf did so.

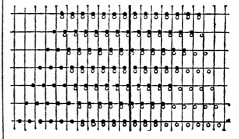
From the table already given it follows that the different ions must possess very different migration-velocities. On comparing, for instance, KCl and LiCl, we find differences of 27 to 30 units. Since the chlorine must possess the same velocity in both cases, and hence participate to the same amount in the conductivity, the difference is due solely to the ions K and Li. In the same manner we can convince ourselves that different negative ions migrate with different velocities.

On electrolysing a substance whose ions travel with different velocities, different quantities of the ions will collect at the electrodes, and, in addition to decomposition, the liquid will undergo an unequal change of concentration at the electrodes. In order to get as good a representation as is possible for these relations, let the ions be represented in the figure in the next column by black and white circles. The different horizontal rows represent different phases of the electrolysis, these being such that between them there is a distance equal to the mean distance of the ions. It is assumed that the black ions migrate twice as quickly to the left as the white ones do to the right. At first there are seven ions on each side of the central line. At the end of the process six ions have been liberated at each end, and, in addition, five out of the seven pairs have remained behind at the left side, and two have disappeared; while of the equal quantity at the right side three have remained behind and four have disappeared. Hence the proportion of salt has decreased at the right side twice as much as at the left—that is, in the same ratio as that in which the two ions migrate.

Therefore, in order to determine the ratio of

the migration-velocities of the two ions of a salt, it is only necessary to determine the proportion of undecomposed salt at each electrode before and after electrolysis; the ratio of the decrease of this proportion at each side is equal to the ratio of the migration-velocities, or is the 'migration-measure.'

This representation holds good on the supposition that the ions which separate at the electrodes do not again go into the solution. If, however, we electrolyse a solution of copper sulphate between copper electrodes, the copper, it is true, will leave the solution at the cathode, but the ion SO₄ will not separate out at the anode, but will combine with the copper of the anode to form copper sulphate, which will go into solution. The solution will, therefore, become, not more dilute, but more concentrated. It is, however, easy to take this circumstance into account. It is only necessary to determine the total quantity of electricity passed through the solution (for instance, by means of a silver voltameter); from this we can calculate how much copper has gone into solution, and we need then only deduct this quantity from the total quantity of copper at the anode, as determined by analysis. It is simpler still to weigh the cathode before and after the experiment: the



increase of weight is equal to the weight of copper which has dissolved at the anode.

Hittorf examined (P. 89 to 106) a great many salts and acids with respect to their 'migration-measures,' and established the fact that the current-strength has no influence on the ratio of the migration-velocities, the temperature has an inappreciably small influence, but the concentration exerts a changeable effect—in some cases it is inappreciably small, while in others it is fairly important.

Hittorf found, for instance, that the concentration of KClAq at the electrodes scarcely changes at all; hence chlorine and potassium must migrate with very nearly equal velocities. If we consider the solution v=1,000l in the table given on p. 191, it follows that of the 119·3 units of conductivity of potassium chloride, 59·65 belong to the chlorine and the same number to the potassium. Knowing these numbers, we can calculate the velocities of migration of all the other ions in the table. NaCl has $100\cdot8$; on subtracting from this $59\cdot7$ for the chlorine, the velocity of the sodium ion becomes $41\cdot1$. Similarly, Li = $32\cdot4$, $18a=49\cdot5$. As $K=59\cdot7$, and as $\frac{1}{4}K_2SO_4=120\cdot7$, it follows that $\frac{1}{4}SO_4=61\cdot0$; hence $\frac{1}{4}Mg=32\cdot5$.

From these numbers we can again calculate the migration-measures, or the changes of concentration at the electrodes, of other salts. Thus $\frac{u}{v}$ is for NaCl = $\frac{41 \cdot 1}{59 \cdot 7} = 0.69$; for LiCl $\frac{32 \cdot 4}{59 \cdot 7}$ = 0.54; for BaCl₂ $\frac{49 \cdot 5}{59 \cdot 7} = 0.83$, &c. It is evident

that when one migration-measure is known, the migration-measures of all corresponding salts can be calculated from the measured conductivities. And since, on the other hand, the conductivities of salts represent the sums of the velocities of their ions, we can realise the very great simplification which these considerations, due to F. Kohlrausch, have produced in the relations of electrical conductivity which formerly appeared so complicated. If, for instance, we know the conductivities of ten salts with the negative ion A and the positive ions B₁ to B₁₀, and of other ten salts with the ion B, and the ions A, to A₁₀, and in addition to these the migration-measure of one salt, we can calculate from these 21 measurements the conductivities and the migration-measures of 100 salts -- that is, we can deduce 200 data.

An extensive examination of these relations, undertaken by Kohlrausch (W. 6, 164), and later by Ostwald (Z. P. C. 1, 74), as well as by Loeb and Nernst (ib. 2, 948), has proved that they hold good with very sufficient accuracy for very dilute solutions. But deviations from these relations become apparent in concentrated solutions, and these deviations are greater the more concentrated the solutions. And, further, different salts behave differently, inasmuch as salts consisting of two monovalent ions show the smallest deviations, salts with one monovalent and one divalent ion show greater deviations, and salts with two divalent ions show the greatest deviations. Now these are the classes of salts which exhibit differences in the changes of conductivity accompanying dilution (p. 191), and we are, therefore, led to the conclusion that the same cause underlies both phenomena.

The free acids and bases only partially conform to Kohlrausch's law. The following data, taken from Kohlrausch's determinations, exemplify this statement. Temp. = 18°:—

preserving very dilute solutions of alkalis perfectly pure.

From these measurements it follows primarily that hydrogen and hydroxyl possess much greater migration velocities than the other ions - namely, 284 and about 150, respectively. It follows, there fore, that if the law of Kohlrausch were generally valid, the conductivities of all acids would be > 284, and the conductivities of all bases would be > 150. On looking at the numbers in the table, we find that phosphoric acid and acetic acid, as well as ammonia, are at a quite hopeless distance from these values. Sulphuric acid, also, does not obey the law. For 280, the value 61.0 was found above; on deducting this number from ½H2SO4=331.6, there remains H=270.6, and the difference of this value from the one found before, 284, exceeds the limits of experimental error.

On carefully studying the table we now see that the acids and bases which form exceptions to the law exhibit the same peculiarity as the exceptional salts; they change their conductivities very markedly upon dilution. While the substances which follow the law have nearly reached the maximum of the equivalent conductivity at 1000l, the numbers show that this is far from being the case in the exceptional substances. We are thus led to inquire as to the cause which brings about the change in conductivity that accompanies dilution.

The arguments used so far have tacitly assumed that all the molecules of the electrolyte participate in the electrical conductivity, and that the electrolyte is therefore completely split This assumption is evidently arbitrary; and when we remember that aqueous solutions of ammonia and acetic acid exhibit nearly normal lowerings of the freezing-points, and that, therefore, dissociation occurs only to a small extent, if at all, we see that the assumption is completely erroneous in these cases. The former equation $\lambda = u + v$, established without reference to the degree of decomposition, must be changed into $\lambda = x (u + v)$ where x represents the degree of decomposition—i.e. the ratio between the molecules decomposed and the total

Dilution	HCl	HNO,	₹H,SO.	H ₃ PO ₄	CH,.CO,H	кон	NaOH	NH4011
11	278·0	277· 0	188·9	29·0	1·2	171·8	149·0	0·84
101	324·4	322·5	208·4	43·0	4·3	198 6	170·0	3·1
1001	341·6	339·5	285·5	79·0	13·2	212·4	187·0	9·3
10001	345·6	342·7	831·6	96·8	38·0	214·0	188·0	26·0

Among the substances quôted, HCl, HNO, KOH, and NaOH follow the law of Kohlrausch. Thus, if 59.7, which is the migration-velocity of K, is deducted from 118, which represents the conductivity of KNO, as determined by Kohlrausch, the difference, 58.3, is the velocity of NO. On deducting this number from the conductivity of nitric acid at 10001—that is, from 342.7—it follows that the velocity of hydrogen From HCl=345.6 it follows that H = 285.9, since Cl = 59.7; these numbers agree to within less than one per cent. In the same way the difference between KOH and NaOH is equal to 26.0, thus approximating to equality with that between KCl and NaCl, which is 19.2. The larger difference proceeds from the difficulty of

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number of molecules originally present. In doing this we make the most obvious, and probably only possible, assumption, that, cateris paribus, the conductivity is proportional to the number of conducting particles or ions.

We may find a method for determining x in the deviations of electrolytes from the simple laws of solutions with respect to vapour-pressure and freezing-point. We have another and much more accurate method, however, in the determination of the electrical conductivity itself. As was found by Kohlrausch, the equivalent (and also the molecular) conductivity increases with rising dilution, reaching a maximum value in the case of salts within limits which are still practically measurable. We can interpret this

behaviour from the standpoint now gained in the sense that the number of molecules split into ions continually increases with increasing dilution, and that the maximum value of conductivity corresponds to a (practically speaking) complete splitting up. Then x=1, and the formula $\lambda =$ x(u+v) changes into the former, $\lambda = u+v$. Now, for most salts the maximum value is reached at a dilution of 1000l, and the decomposition is nearly complete; it follows from this, that the law of Kohlrausch $\lambda = u + v$ holds good for this maximum dilution. This can be asserted for the strong acids and bases, such as HCl, HNO₃, KOH, and NaOH. On the other hand, as can easily be seen from the course of the numbers. the increase has not nearly reached its limit for acetic acid and ammonia at a dilution = 1000l; hence the value x is still very far removed from unity, and the law of Kohlrausch cannot hold in its simplest form.

On the assumption that the molecular conductivities of dilute solutions depend only on the number of ions present, the values of x can be easily calculated from a knowledge of the values of the conductivities at different dilutions, as well as of their maximum values; designating the molecular conductivity at the dilution of v litres by μ_v , and the maximum value by μ_∞ , then, according to the assumption made, $x = \frac{\mu_v}{r}$.

 $= \mu_{\infty}^{-1}$.

Whether the above-mentioned assumption is strictly correct cannot be asserted a priori. It must be looked upon as probable, since the resistances to motion which the ions experience, and which determine the velocities u and v, depend on the nature of the solvent, which in these cases is water; and considering the small differences which exist between the properties of very dilute solutions and those of water, no appreciable difference in the resistances to motion can be supposed to exist in those differently diluted solutions. But the admissibility of the assumption can also be put to experimental proof in determining x on the one hand by the deviations from the laws of solution, and on the other hand by the equation just given.

If m_0 is the theoretical molecular weight, and m the molecular weight determined from the laws of solution (for instance, from the lowering of the freezing-point), then for electrolytes $m > m_0$; following van't Hoff, we put the ratio of

the two $\frac{m_0}{m} = i$. If, on the other hand, x is the

number of molecules split up, the total number of molecules being put equal to 1, and if n is the number of ions into which each molecule splits (for KCl n=2, for BaCl₂ n=3, &c.), the number of molecules in the solution will have increased in the ratio 1:1-x+nx, that is, in the ratio 1:1+(n-1)x, and we have, therefore, i=1+(n-1)x.

Arrhenius, to whom we owe this calculation, has shown (Z. P. C. 1, 634) that in the greatest number of cases investigated there is an excellent accordance between the values of i as calculated from the electrical conductivities and those deduced from the freezing-points. Individual cases which at first seemed to be exceptions have, on later investigation, also proved to be

subject to the law $(Z.\ P.\ C.\ 2,\ 491)$, and the apparent deviations have been shown to be due to incorrect observations. It is only at greater concentrations, where the suppositions made in determining molecular weights from the freezing-points, and those made in determining x from the electric conductivities, become uncertain, and not applicable, that marked deviations are apparent.

In addition to this confirmation of Arrhenius' assumption that in electrolytes a more or less considerable portion of the molecules is split up, a further confirmation has been found by W. Ostwald, which is based on the following considerations (Z. P. C. 2, 36, and 270). spontaneous decomposition which the electrolytic molecules undergo in solutions is of the nature of dissociation; and since, according to van't Hoff, the substances in solution are subject to the gaseous laws, it must be possible to represent the process by the same formulæ as hold for the dissociation of gases. Considering the simplest case, that each molecule splits into two parts, there holds for gases (vol. ii. p. 406) the relation $\frac{p_2^2}{}$ = constant, where p_1 is the partial pressure of the undecomposed portion, and p that of the decomposed portion.

In order to apply this equation to electrolytes, we notice that the pressures are proportional to the numbers of the molecules, and inversely proportional to the volumes occupied by these. If x is the part dissociated, and v the volume, then $p_1 = \alpha \frac{1-x}{v}$ and $p_2 = \alpha \frac{x}{v}$; α represents a factor dependent on the units chosen. Further, according to the former equation, we have $x = \frac{\mu_v}{\mu_\infty}$; introducing these values into the dissociation equation, we get $\frac{1-x}{x^2}v = k$, where k is a constant,

and from this $\frac{\mu_v^2}{(\mu_\infty - \mu_v)\mu_\infty} = kv$.

This equation represents a relation between

This equation represents a relation between the conductivity and the volume, or the dilution, which must be followed by all binary electrolytes, if the assumptions made are correct.

The equation cannot well be tested in the case of salts, as these very nearly reach complete decomposition; the value $\mu_{\infty} - \mu_{v}$ thereby becomes very small, and the experimental errors exert a preponderating influence. Those electrolytes which are only partially dissociated, such as acetic acid &c., lend themselves much better to applying the test. Here also it seems as if a difficulty met us, the quantity μ_{∞} not lending itself to direct measurement because, with the very great dilutions necessary, the inevitable impurities of the purest water make accurate measurements impossible. But the law of Kohlrausch provides us with a means for making the measurements. It appears that the salts of weak acids are dissociated to just the same extent as those of strong acids; it is possible, therefore, as was shown at pp. 192-3, to calculate the migration-velocity of the negative ion. On adding to this the migration-velocity of hydrogen, which can also be determined (p. 193), the sum represents the conductivity of the completely

discociated acid—that is, μ_{∞} , the quantity

sought (Ostwald, Z. P. C. 2, 840).

In this manner, then, we obtain all the data for testing the above equation, and it proves itself to be in direct accordance with the facts. The examination is best carried out by calculating the constant k for a number of values of μ at different dilutions, and seeing whether it really is constant. In the following tables a few examples are given. The numbers hold for 25°, and are molecular conductivities referred to the mercury unit :-

Acetic acid, $\mu_{\infty} = 364$.

•	μ	100 μ _{μ∞}	k
8	4.34	1.193	0.000180
16	6.10	1.673	179
32	8.65	2.380	182
64	12.09	3.33	179
128	1 6·99	4.68	179
256	23.82	6.56	180
512	32.20	9.14	180
1024	46.00	12.66	177
			0.000180

The values of k, calculated from eight different measurements, agree within the limits of experimental error.

The esmotic pressure of acetic acid at v=8corresponds to about three atmospheres, that at v = 1024 to about $\frac{1}{45}$ atmosphere, or to 1.7 mm. The dissociation mercury pressure. has, therefore, proved correct within these wide limits; no examination of the law so farreaching as this has ever been possible in the case of gases.
Further examples are:

Cyanacetic acid, $\mu_{\infty} = 862$.

v	μ	$100 \frac{\mu}{\mu_{\infty}}$	Ł
16 82 64 128 256 512 1024	788·0 105·3 139·1 176·4 219·1 260·9 297·8	21·7 29·1 38·4 48·7 60·5 72·0 82·1	0·00376 373 374 361 362 361 868
			0.00370

Here also the individual values of the constant vary irregularly about the mean value, so that the discrepancies have to be ascribed to experimental errors.

In the accompanying tables, which have been taken at random from a great number, the dissociation law proves itself everywhere to be accurate. It has, in fact, been verified in every case in the course of an investigation which, for reasons to be discussed later, has been carried out with more than 400 acids, and it can be looked upon as generally valid.

We must not omit to remark, however, that, as observed above in the cases of the very strong

Oxyisobutyric acid, $\mu_{\infty} = 355$.

٠	μ	100 4	k
32	20.05	5.65	0.000106
64	28.05	7.91	106
128	38.86	10.95	105
256	53.91	15.20	106
512	73.49	20.70	106
1024	99.52	28.05	106
			0.000106

Oxysalicylic acid, $\mu_{\infty} = 356$.

υ	μ	$100 \frac{\mu}{\mu_{\infty}}$	k
64 128 256 512 1024	84·1 112·5 147·0 187·0 230·0	23·6 31·6 41·3 52·6 64·7	0·00114 114 113 114 116
2048	270.4	76.0	0.00115

Orthochlorobenzoic acid, $\mu_{\infty} = 356$.

ν	μ	100 μ μ _∞	k
64 128 256 512 1024	89·2 119·4 156·1 197·0 238·7	25·1 33·5 43·8 55·3 67·1	0-00131 131 133 133 134
			0.00132

acids, and the neutral salts, which are very nearly completely dissociated, the calculation of the constants becomes very uncertain, and that greater deviations exist between the observations and the formula than can be ascribed to experimental errors. Which of the assumptions already made will have to be modified in order to explain these small discrepancies cannot as yet be determined; but anyhow the discrepancies are so inconsiderable, and there are so many cases of agreement, that the deviations must be looked upon as of a secondary nature, and undoubtedly they will receive their explanation later on. It is well not to pass over in silence, nor to attempt to hide, such deviations from a law which is otherwise generally valid, since in most cases theoretical progress is connected with the investigation of the causes of deviations from general laws. Examples of this are found in the deviations of gases from the simple laws, and the theory connected with this by van der Waals, and in the deviations of some substances from the law of Avogadro and the elucidation thereof by the theory of dissociation.

Chemical applications.—The laws of electrical conductivity, as developed in the preceding pages, give us the means for solving many chemical problems. This they do, because they give a deeper insight into the constitution of dissolved substances than is afforded by the usual chemical methods. A series of such applications is already contained in the preceding pages; others are considered in the following

paragraphs.

The first of these applications is connected with the question as to the nature and composition of the current-conducting particles or ions of salts. The conception of Berzelius that these are the 'anhydrous' acid and base has, in the light of Faraday's law, been recognised as erroneous. Afterwards, in analogy with the compounds of organic chemistry, the salts were looked upon as having a unitary composition, and this view is still generally held. But this conception cannot be regarded as quite correct, since it does not take into account the fundamental difference which undoubtedly exists between salt-like compounds and indifferent compounds, a difference which manifests itself chiefly in the capacities for reaction of the two groups. The conception of Berzelius rested on the correct recognition that salts are dual compounds; he had only misconceived their constituents.

The new electro-chemical theory of Arrhenius avoids both errors, and retains what is correct in each conception. The solid salts, and, according to concentration, a greater or smaller portion of the salts dissolved in water, 'salt' being understood to include acids and bases, have, it is true, a unitary composition; but that portion of them which is capable of reacting chemically, and which therefore is of preponderating interest

to the chemist, is split binarily.

'Salts,' in the general meaning, are therefore substances which very readily undergo such a binary splitting, and herein lies their binary character, correctly recognised by Berzelius.

It is a most remarkable fact that in the splitting up of 'salts' their constituents appear in oppositely charged electrical conditions. This indicates that probably both phenomena stand in the closest connection. It may be asked whether the salts split up so easily because their parts readily acquire opposite electric charges, or, conversely, whether the parts are easily charged electrically because they are readily separated. Not less remarkable is the fact that whenever a salt molecule is split up an equal quantity of electricity is produced (or separated), independently of the nature of the parts. It is beyond doubt that this fact is of decisive importance for the recognition of the nature of electricity, as well as of that of chemical affinity; when the one is elucidated, the other will be so also. But which of the two will first be thus advanced cannot as yet be foreseen.

A further remarkable fact is, that pure salts at ordinary temperatures do not conduct appreciably; the ordinary solid salts do so as little as pure sulphuric acid, acetic acid, &c. Even the halogen acids HF, HCl, HBr, and HI, when liquefied by pressure or cold, are all non-conductors. According to the dissociation theory of electrolytes this is quite conprehensible; the pure substances do not conduct because they are not dissociated, and they are not dissociated because their molecules have no space wherein to dissociate, or because they do not possess sufficient mobility.

Whether dissociation occurs or not when salts are dissolved depends in a marked way on the nature of the solvent. Hydrochloric acid, which when liquid is not dissociated, dissociates at once when dissolved in water. But on using as solvent benzene, xylene, hexane, or ether, no dissociation ensues; these solutions are nonconductors (Kablukoff, Z. P. C. 4, 430). On using alcohols as solvent, methyl alcohol is found to behave almost like water; the solutions of hydrochloric acid in it have a molecular conductivity about three times smaller than aqueous solutions. The conductivities of the solutions become less and less as we use ethyl alcohol, isobutyl alcohol, and isoamyl alcohol.

It is still quite unknown what the condition is which determines the property of different solvents to cause dissociation to a different degree, and it is not known whether, as is somewhat probable, these effects remain proportional in the cases of all salts. Here it seems, however, as if the capability of the solvent itself to separate into ions, even if to a small degree only, determined its power to cause the dissociation of substances

dissolved in it.

Although the questions just mentioned still await solution, partly for lack of experimental researches in this domain, yet the new conception of the binary character of salts allows us to distinguish more sharply between the different classes of these compounds than was possible before. This is especially so in the case of the so-called double salts. This name is at present somewhat uncertain in its application. Potassium copper sulphate K₂SO₄.CuSO₄ is called a double salt, but sodium platinichloride is also formulated as such, 2NaCl.PtCl₄; the first formula is correct, but the latter is wrong. This is so because potassium copper sulphate gives all the reactions of potassium sulphate as well as those of copper sulphate, but from sodium platinichloride silver solutions do not precipitate silver chloride as from sodium chloride, but silver platinichloride Ag₂PtCl₆ (Jörgensen, J. pr. [2] 16, 845). It is probable, therefore, that sodium platinichloride is a salt of chloro-platinic acid H₂PtCl₈. If this is correct, the ions of the sodium salt are 2Na and PtCl₈, and in electrolysis the platinum must not, like the other metals, go to the cathode, but as a constituent of the acid radicle to the anode. And, in fact, Hittorf has proved (P. 106, 520) that this is the case.

Similarly with other salts; for instance, that which is obtained from chromium oxide and potassium binoxalate, and which has the empirical formula $3K_*C_2O_*.Cr_2(C_2O_*)_s$. Since this salt gives no pp. with calcium salts it cannot be regarded as an oxalate. In fact, in electrolysis the chromium goes to the anode; the chromium is, therefore, a constituent of the acid radicle, and the salt must be looked upon as the potassium compound of a tribasic chromoxalic acid $(HC_*O_*)_*Cr$ —that is, as $K_*C_*O_{12}Cr$ (Kistiakowsky, Z_* . P_* . C_* . C

Such salts, therefore, are not double salts, but are salts of complex acids; it is possible to test, in the way already indicated, in every case whether a definite salt is a true double salt or the salt of a complex acid. On examination we find that, not a double salt, but a salt of a

complex acid is present whenever the reactions of the salt in question are not those of its constituents. This decision is specially convenient when the solution of the salt is coloured. Because, since the properties of the positive ions, and therefore their colour also, are independent of the negative ions, it will always be possible to conclude that the positive ions are no longer present when their colouration disappears. In the easily decomposable salt 2KCN.Ni(CN)₂ we must not see a double salt, but the potassium salt of cyano-nickelic acid H₂Ni(CN)₄, because its solution is yellow, and not green as solutions of the nickel salts are which contain nickel as an ion.

The line of demarcation between the two groups of salts, however, is not absolute, as many salts exist which in aqueous solutions are both double salts and also complex ones. This is the case with potassium ferri-oxalate, for instance. Since this salt is green, while all true ferric salts are yellow or brown, it may be concluded that it is a complex salt, the compound of a ferri-oxalic acid H_aC_aO₁₂Fe analogous to the above-mentioned chromo-oxalic acid. The solution of this salt gives, however, a precipitate with calcium solutions, which proves it to be an ordinary oxalate, but on the other hand the iron goes to the anode. From this it is seen that the aqueous solution contains chiefly the ions 3K + C₆O₁₂Fe, but that a portion of the salt has split in the manner of a double salt into potassium oxalate and ferric oxalate, which on their part form the corresponding ions.

The study of such cases has as yet scarcely been begun, because until now we possessed neither theoretical nor experimental means for answering the questions which arise. The electrical methods, in conjunction with the methods for the determination of molecular weights arising out of the theory of solutions, now afford easy access to these hitherto closed and uncultivated domains.

The dissociation-constants of organic acids. The only province of chemistry which has been studied somewhat thoroughly in the light of the new electro-chemical theory is that of the organic acids. We have seen above that the molecular conductivity of these can be

represented by the formula
$$\frac{\mu^2_v}{(\mu_{\infty} - \mu_v)\mu_{\infty}} = kv$$
.

This formula contains only the single constant k, which constant depends on the nature of the acid, the temperature, and the solvent, but is independent of the dilution. At a given temperature, and in the same solvent, the quantity k is therefore a measure of the electrical conductivity, and hence it is the capacity for reaction of the acid; for the substance considered, k therefore represents the long sought for numerical value of the chemical affinity.

It is remarkable that the two conflicting views concerning the nature of chemical affinity, as represented by Bergmann and Berthollet (cf. vol. i. 68) here coincide. Berthollet, who represented the affinity as dependent on the mass, was right, and so was Bergmann, who desired to represent it by a constant. By accurately formulating the idea of the influence of mass as given by Berthollet, we obtain in the equation for the

chemical effects a co-efficient independent of the quantity—that is, one referred to the chemical unit of quantity, which is Bergmann's measure of chemical affinity.

It has already been shown (vol. i. 81) that the electrical conductivities bear a definite relation to the composition and constitution of the acids. But the law of dissociation of electrolytes had not been discovered at the time when the article referred to was written, and the observed relations could not be represented numerically. This has now become possible, by means of the values of the constant k, which have been measured for about 400 acids (Ostwald, Z. P. C. 3, 170; Bethmann, ib. 5, 385; Bader, ib. 6, 289).

Before expounding these relations, it will first be shown how the laws which have been found empirically for the electrical conductivities follow directly from the above dissociation-formula. In order to make the consideration easier, the formula will first be somewhat simplified by substituting for the molecular conductivity μ_v its value referred to the maximum $\frac{\mu_v}{\mu_\infty} = m$.

The equation then assumes the simple form $\frac{m^2}{1-m} = kv.$

In the first place, we see that as the dilution v increases infinitely, the quantity 1-m must approach zero. Hence m must approach unity; hence the electrical conductivity increases with dilution up to a maximum, as has been shown by experience.

Further, it was found that when at any dilutions, v_1 and v_2 , two different acids have the same (relative) conductivity, they also have the same at other dilutions v_1' and v_2' , as long as the new dilutions are in the same ratio as the old ones—that is, when $v_1: v_2=v_1': v_2'$. This follows from the formula in this way: on the supposition that the relative conductivities are the same,

we have first of all
$$\frac{m^2}{1-m} = k_1 v_1$$
; $\frac{m^2}{1-m} = k_2 v_2$;

therefore $k_1v_1 = k_2v_2$. And in the same way for the other dilutions, $k_1'v_1' = k_2'v_2'$, from which it follows directly that $v_1 : v_2 = v_1' : v_2'$. In addition, F. Kohlrausch had already pointed

In addition, F. Kohlrausch had already pointed out that in the cases of acids which conduct badly (which, therefore, are but little dissociated), the molecular conductivity increases nearly as the square root of the dilution. If in the equation $\frac{m^2}{1-m} = kv, m \text{ is very small, } 1-m \text{ differs little}$

from unity, and can be looked upon as constant; then approximately, $m^2 = kv$, or $m = \sqrt{kv}$.

Finally, on taking the logarithms of the dilutions as abscisse and the values of m as ordinates, we also get from the formula a curve resembling a tangent-function (vol. i. page 82); the interpolation-formula given in vol. i. must be replaced by the present rational formula.

As regards the meaning of the constant k, we recognise what this is on putting m = 0.5, when we get $2k = \frac{1}{2}$. 2k is therefore the reciprocal

value of the volume, or the concentration at which the electrolyte is just half dissociated. This number is extremely characteristic for dif-

ferent acids, since its value may vary according to the nature of the substance between the limits 1 and 1,000,000.

The measurement of the quantity k for different acids has led to the following general law:—
The constant k is approximately the product of a number of factors, each of which depends on the nature of the constituents of the acid and on their position relatively to the carboxyl group.

In order to realise what this law implies, and in what manner it is applied, we will proceed to discuss in the following pages the most important groups of the organic acids. The relations, the main features only of which could formerly (vol. i. page 82) be investigated, can now be represented numerically with perfect exactitude. The numerical values for k have been multiplied by 100 in order to get rid of ciphers; they all hold for aqueous solutions at 25°.

The fatty acids. The following constants were measured:—

Formic acid H.CO $_{2}$ H . . . 0.00214 Acetic acid CH $_{3}$.CO $_{2}$ H . . . 0.00180 Propionic acid C $_{2}$ H $_{3}$.CO $_{2}$ H . . 0.00134 Butyric acid C $_{2}$ H $_{3}$.CO $_{2}$ H . 0.00149 Isobutyric acid (CH $_{3}$) $_{2}$ CH.CO $_{2}$ H . 0.00144 Isovaleric acid (CH $_{3}$) $_{2}$ C $_{4}$ H $_{3}$.CO $_{2}$ H . 0.00161 Caproic acid C $_{3}$ H $_{11}$.CO $_{2}$ H . 0.00145 Formic acid has the highest constant; the re-

Pormic acid has the nignest constant; the replacement of its hydrogen by methyl reduces the constant to half its value. But on again replacing one hydrogen atom in acetic acid by methyl the constant only decreases to $\frac{1}{1\cdot 4}$ of

its value, and further similar substitutions no longer produce a weakening of the acid, but some of them even bring about a slight strengthening. These changes are, however, but small, so that the constants of acids of this series having more than three carbon atoms vary irregularly about the mean value 0.0014.

It is evident, then, that the substitution of hydrogen by methyl acts quite differently, according as it occurs next to the carboxyl or at a greater distance from it. This is a result which will afterwards be shown to hold good in all cases; on the whole, the effect of each substituent is the smaller the more distant it is from the carboxyl. The inference that the interposition of even two carbon atoms nearly counteracts the effect is correct for methyl, for which it is relatively small. Other constituents of greater energy extend their action further, but yet not over more than three or four atoms of the 'open chains'; in the case of 'closed chains' other relations hold good.

Halogen derivatives of the fatty acids. The entrance of chlorine in the place of hydrogen in the fatty acids exerts very considerable influence.

The constants are—

1:86:2840:67200. The ratio of the values of the constants for the different acids, referred to the entrance of each separate chlorine atom, is 84, 33, and 24. It is only the ratios of the constants, and not their differences, which are

important; the ratios, though not equal, are yet of the same order of magnitude, while the differences between 0.155, 5, and 121 are quite incomparable. Further, it follows that the relative change of the constants for the entrance of each chlorine atom is not of the same value; the first chlorine atom acts more intensely than the second, and the second more intensely than the third. We shall again meet with a similar relation when dealing with substitution in connection with the same carbon atom. From this it may be concluded that the first chlorine atom which enters into acetic acid exerts its influence under more favourable conditions, therefore probably from a less distance, than the second and third, and it becomes evident how well such measurements lend themselves to investigations and examinations of the relations in space of the atoms within the molecules. The action of other elements and radicles is very similar to that of chlorine.

Cyanacetic acid CH₂.CN.ČO_H . 0.370
Thiocyanacetic acid CH₂.SCN.CO₂H . 0.265
So-called 'carbamine thioglycollic acid' CH₂(SCONH₂)CO₂H . 0.0246
Isothiocyanacetic acid C₃H₃O₂S.N . 0.000246
Thiacetic acid CH₂.COSH . . 0.0469

. 0.138

Monobromacetic acid CH₂Br.CO₂H

The constant of monobromacetic acid does not differ much from 0.155, that of monochloracetic acid, hence both halogens exert about the same influence. Cyanogen acts much more intensely. The ratio of cyanacetic acid to acetic acid has risen to 205 (from 86). Thiocyanacetic acid also is stronger than monochloracetic acid, but not so strong as cyanacetic acid, though sulphur is generally a negative substituent—that is, one which augments the acid properties. The idea of relations in space at once suggests itself; by the introduction of sulphur there has, in fact, taken place, on the one hand, an increase in the acid properties, but on the other hand, since the cyanogen is removed to a greater distance from the carboxyl, there has been a weakening, and the latter effect preponderates over the former. By taking up water, thio-cyanacetic acid easily passes into the acid CH2.SCONH2.CO2H, the group CN changing into This transformation is accompanied CO.NH2. by a marked decrease in the conductivity, the constant falling to less than a tenth of its former value. The change can be conveniently studied in an aqueous solution of thiocyanacetic acid, by determining its electrical conductivity; the conductivity decreases continuously, and after even a few days the greater part of the thiocyanacetic acid is found to be changed. Finally, there is a compound isomeric with thio. cyanacetic acid, which Volhard (J. pr. [2] 9, 6) obtained by the action of hydrochloric acid on sulphohydantoin, and which is isothiocyanacetic acid. The constant for this compound is 10,000 times smaller than that for thiocyanacetic acid; it is also smaller than that of any other carbon acid, so that the inference must be drawn that the substance is not a carbon acid at all. This confirms the view expressed by Liebermann, and the reasons for which were given by Hantzsch (B. 20, 3129), that the compound is a dioxythiazole-

The feebly acid properties are ascribed to the hydrogen of the imide group. The thiacetic acid mentioned at the end of the last table contains sulphur in the place of oxygen in the hydroxyl of the carboxyl. In accordance with the negative nature of sulphur, this acid proves to be twenty-six times stronger than acetic acid, and this factor may be looked upon as the greatest to which sulphur can give rise when replacing oxygen, because in this case the sulphur atom acts from the most favourable position which is ever possible.

Oxyacetic acid and its derivatives.

 $\begin{array}{lll} \mbox{Glycollic acid } \mbox{CH}_2\mbox{OH.CO}_2\mbox{H} &. & 0.0152 \\ \mbox{Methoxy-acetic acid } \mbox{CH}_2\mbox{(OCH}_3\mbox{)CO}_2\mbox{H} &. & 0.0335 \\ \mbox{Ethoxy-acetic acid } \mbox{CH}_2\mbox{(OC}_2\mbox{H}_3\mbox{)CO}_2\mbox{H} &. & 0.0234 \\ \mbox{Phenoxy-acetic acid } \mbox{CH}_2\mbox{(OC}_8\mbox{H}_3\mbox{)CO}_2\mbox{H} &. & 0.0474 \\ \mbox{Glyoxylic acid } \mbox{CH}(\mbox{OH}_2\mbox{OC}_2\mbox{H} &. & 0.0474 \\ \end{array}$

The replacement of hydrogen by hydroxyl in acetic acid produces a rise in the constant of 9 times its former value; hydroxyl, therefore, acts much less intensely than the halogens. By introducing a second hydroxyl the constant again becomes greater, but only 3.1 times greater. Here we find quite the same effects as in the cases of mono- and di-chloracetic acid, the second substitution of the same substance acting less strongly than the first. In the numerical values, even, we cannot overlook a certain correspondence; in the case of chlorine the ratios are 1:86 and 1:33; in the case of hydroxyl they are 1:9, and 1:3.1. It may therefore be said that, using round numbers, in the derivatives of acetic acid chlorine acts ten times as intensely as hydroxyl. When the hydroxylic hydrogen of glycollic acid is replaced by radicles, the constant changes, and, strange to say, methyl acts in this case in a strengthening manner, the constant rising to a little more than twice its former The substitution of hydrogen in this value. methyl by methyl again has a weakening effect, as is the case with the fatty acids; ethoxy-acetic acid is 1.4 times weaker than methoxy-acetic acid. Finally, by the introduction of phenyl, the constant becomes appreciably greater; it is about five times as great as that of glycollic acid, and is 2.3 times greater than that of methoxy-acetic acid. The ratio is smaller than that between acetic acid and benzoic acid, which is 1:3.3, an indication that the exchange of methyl for phenyl has in the latter case taken place in greater proximity to the COOH group, a conclusion which follows also directly from the formulæ.

A number of other substances related to glycollic acid have given the following numerical

Thioglycollic acid CH₂(SH)CO₂H . 0.0225
Thiodiglycollic acid S(CH₂CO₂H)₂ . 0.048
Ditriodiglycollic acid S₂(CH₂CO₂H)₂ . 0.065
Digi ycollic acid O(CH₂CO₂H)₂ . 0.11

As the constant of glycollic acid is '0152, we see that the substitution of S for O in hydroxyl has caused the constant to increase 1.5 times; now, as substitution of S for O in COOH of acetic

acid increased k from 1 to 26, we see here also the great influence of the position occupied by the replacing atom.

The three other acids quoted in the last table are dibasic, and the question at once arises how their constants must be calculated, since the dissociation-formula has been developed for binary electrolytes—that is, for monobasic acids only, and not for ternary ones, to which class the dibasic acids belong. It can, however, easily be shown that, as long as the dissociation is not great, the weaker dibasic acids are not electrolysed according to the formula $2H+R^{n}$, but according to the formula H+HR. In other words, at first one hydrogen atom only is split off, and the dissociation takes place according to the binary scheme of the monobasic acids. This follows from the fact that the change in the molecular conductivity of such acids can be represented by the same formula $\frac{m^2}{1-m} = vk$, so

that k remains constant. But the formula calculated on the assumption of a dissociation according to the scheme $H_{*}R^{1} = 2H + R$ does not represent the change in the conductivity of these acids as determined by experiment.

Malonic acid CH₂(COOH)₂ may serve as an example of dibasic acids. It gives:—

-	•	μ	m	100k
	16	53·1	0.149	0.159
	32	72.3	0.202	0.159
	64	97.1	0.272	0.158
	128	128.5	0.359	0.157
	256	165.9	0.464	0.157
	512	208 8	0.586	0.162
	1024	253.2	0.708	0.168
	2048	294.2	0.823	0.187
			10	

The constant does not change till the value m=0.586 is reached—that is, till about half the acid has been dissociated according to the binary scheme; then it becomes greater—a sign that henceforward the formula loses its validity, and that the decomposition according to the ternary scheme 2H+R'' begins to assume an appreciable value.

If, on the other hand, we attempt to calculate the conductivity according to the formula corresponding to ternary dissociation $\frac{m_s}{m_s} = k$.

sponding to ternary dissociation $\frac{m_s}{(1-m)v^2} = k$, the values of k decrease very rapidly and are far from being constant.

The values for the constants of dibasic acids given in the table preceding the last one have been calculated on this principle, and refer, therefore, to the first stage of binary dissociation $H_2R = HR + H$.

Comparing thioglycollic acid with thiodiglycollic acid and dithiodiglycollic acid, we find that the constants do increase distinctly, though only slightly; they are 0.0255, 0.048, and 0.065, and are in the ratio of 1:21:29. When, therefore, the residue of acetic acid, CH_CO_H, enters thioglycollic acid, this process has no appreciable influence on the constant. On the other hand, the constant of diglycollic acid is 7.2 times greater than that of glycollic acid. From the fact that the same substituent produces such different effects in glycollic acid and in thioglycollic acid, it may be presumed that in the first case the oxygen brings the negative substituent nearer to the carboxyl than the sulphur does in the second case, and this presumption may be further utilised for hypotheses concerning the configuration of the sulphur atom, and the arrangement of the affinity points on it. It has, however, not yet been established with certainty that the greater or less distance of the groups is the only cause of their different actions, and there are some circumstances which allow us to conclude that other conditions are effective also; therefore, until these have been cleared up, conclusions such as the above hold only hypothetically.

Derivatives of amido-acetic acid.

Phenylamido-acetic

acid CH₂(NHC₈H₄)CO₂H . 0.0039 Hippuric acid CH₂(NH.CO.C₈H₃)CO₂H . 0.0222 Aceturic acid CH₂(NH.CO.CH₃)CO₂H . 0.0230 Phthalylamido-acetic acid

 $CH_2(NC_2O_2C_6H_4)CO_2H$. . 0.100

Amido-acetic acid has not the character of an acid; the basic NH₂ group completely removes the acid properties of acetic acid. But if negative radicles are introduced into NH₂, well defined acids are again obtained. Phenyl has this effect; the constant of phenylglycocoll is 2·2 times as great as that of acetic acid. The constants of the acids obtained by introducing CO.C₈H₃ and CO.CH₃ for H in NH₂ are 12 to 13 times greater than that of acetic acid. The substitution of the divalent residue of phthalic acid for the two hydrogen atoms in the NH₂ group raises the constant of acetic acid 55 times.

Derivatives of propionic acid.

The constant of a-oxypropionic or lactic acid is 10 times greater than that of propionic acid, but the constant of β -oxypropionic acid is only 2.3 times greater than that of propionic acid. The difference in the action of the same substituent, according to its nearer or more distant position relatively to COOH, asserts itself most clearly. The ratio between propionic acid and lactic acid is slightly greater than that between the analogous substances acetic acid and glycollic acid (1:9); it is, therefore, not quite immaterial whether the substitution does or does not take place in the group CH_s. The influence of the more distant position of the substituent can be recognised in β -iodopropionic acid and trichlorolactic acid. The first is only 6.5 times stronger than the parent substance. No measurement of a-iodopropionic acid has yet been made, but there is little doubt that iodine in the a- position must act very similarly to chlorine or bromine, and must, therefore, increase the constant by 70 or 80 times. The effect in the B- position is 10 to 12 times less than this. In the same way the substitution of 3Cl for 3H in CH_s.COOH increases k about 67,000 times; but the introduction of 30l into CH3.CHOH.COOH in the \$- position increases k only about 33.7

times. The ratio of these effects is about 1:2000; calculated for each Ol atom the ratio is $2\sqrt[3]{2000} = 12.6$; this ratio comes very near that estimated for a- and β -iodopropionic acids.

Derivatives of higher fatty acids.

Oxyisobutyric acid $(CH_3)_2CH.OH.CO_2H$ 0.0106 Mononitrocaproic acid $O_4H_{10}(NO_2).CO_2H$ 0.0123 Dinitrocaproic acid $C_4H_6(NO_2)_2CO_2H$. 0.069 Lævulinic acid $CH_4.CO.CH_2.CH_2.CO_2H$ 0.00255

Since the constant of isobutyric acid is 0.00144, the hydroxyl in the α - position here causes the factor 7.4, which is somewhat smaller than 9, the corresponding ratio between acetic acid and

glycollic acid.

The ratio between caproic acid and its mononitro- derivative is 1:8.5. Since, as will soon be shown, the nitro- group has a more strongly negative action than chlorine, NO_2 must, in the present case, be in the β - position. The same holds for the dinitro- compound; it is 5.6 times stronger than the mononitro- compound; it is again evident that the second negative group has less effect than the first.

Lævulinic acid, or β -acetyl-propionic acid, allows us to recognise the smaller influence to be expected of the acetyl in the β - position; the constant is only 1.9 times larger than that of

propionic acid.

Benzoic acid and its derivatives. The largest and most varied group of allied substances that has been investigated so far, is that of benzoic acid and its derivatives, and consequently it has been possible to trace here many and striking regularities.

Benzoic acid $C_0H_0CO_2H$. . . 0.0060 Salicylic acid $C_0H_1(OH)CO_2H$. . . 0.102 m-Oxybenzoic acid $C_0H_4(OH)CO_2H$. 0.0087 p-Oxybenzoic acid $C_0H_4(OH)CO_2H$. 0.00886 Oxysalicylic acid [1:2:3] $C_0H_4(OH)_2CO_2H$ 0.114 Oxysalicylic acid [1:2:5] $C_0H_4(OH)_2CO_2H$ 0.108 β -Resorcylic acid [1:2:4] $C_0H_1(OH)_2CO_2H$ 0.052 a-Resorcylic acid [1:2:6] $C_0H_3(OH)_2CO_2H$ 5.0 Protocatechuic

acid [1:3:4] C₆H₃(OH)₂CO₂H . . 0.0088 m-Dioxybenzoic acid C₆H₃(OH)₂CO₂H . 0.0091 Gallic acid [1:3:4:5] C₆H₂(OH)₃CO₂H . 0.0040 Pyrogallic acid [1:2:3:4] C₆H₂(OH)₃CO₂H 0.055 Phloroglucin carboxylic

acid [1:2:4:6] $C_6H_2(OH)_3CO_2H$. 2·1

The numbers indicate the positions of the OH groups relatively to COOH supposed to be

in position 1.

This table contains the constants of all hydroxyl-benzoic acids known up to the present time. These show such close connections with each other that, knowing the constants of benzoic acid and of the three monoxybenzoic acids, those of all the others can be calculated approximately. It is true the calculation is not quite exact, but it is sufficiently so to enable us to deduce the constitutions of the corresponding substances directly from a knowledge of the constants.

Between the constants of benzoic acid and its ortho-oxy-derivative the ratio is 1:17. This value is larger than the ratio when OH is introduced in the α - position into a fatty acid, in which cases the ratio did not exceed 10; in the β - position, which corresponds to the constitution of salicylic acid, the ratio was not larger than 2.3. From this

it appears that the ortho-position involves a much closer relation in the benzene nucleus than even the a- position in the open chain, a fact which later on will be confirmed in many ways, and which seems to be of great importance in dealing with the question concerning the constitution of benzene.

The constant is increased but slightly by OH in the meta-position. But in the para-position the hydroxyl even produces a diminution of the constant to rather less than half the value. We deal in this case, therefore, not only as usual with a greater or smaller value of the influence exerted by a substituent according to its position, but with a reversal of the direction in which the influence is exerted. This fact, which has already become evident in the case of methyl, recurs with other feebly negative substituents in the paraposition of the benzene nucleus. This promises to become very important in the construction of a theory to explain these phenomena.

We have then the following factors for the influence of the hydroxyl:—

Ortho- 1:2 or 1:6=17

Meta- 1:3 ,, 1:5 = 1.4Para- 1:4 = 0.5

By their help we are able to calculate the constants of the di- and tri-oxybenzoic acids by multiplying the constant of benzoic acid by the corresponding factor for every hydroxyl introduced. It must be remembered that such a calculation can only give approximate results; for we have seen that substituents already present somewhat influence the magnitude of the factor of a newly entering substituent, and that, generally, by decreasing it. It is, thorefore, only the order of magnitude of the constants, and not their exact value, that we shall be able to find in the manner indicated.

Four dioxybenzoic acids can be derived from salicylic acid; the constants of these four acids are calculated thus:—

1:2:3 gives $.0060 \times 17 \times 1.4 = 0.143$ 1:2:4 ,, $.0060 \times 17 \times 0.5 = 0.051$ 1:2:5 ,, $.0060 \times 17 \times 1.4 = 0.143$ 1:2:6 ,, $.0060 \times 17 \times 1.7 = 1.73$

The numbers observed are 0.114, 0.052, 0.108, and 5.0. In the case of the first and the third acid which both contain a hydroxyl in the m-position, a diminution of the combined effect therefore makes itself felt as usual. In the second acid 1:2:4, calculation and experiment completely agree; while the fourth acid 1:2:6, which contains two ortho- hydroxyls, shows a value about three times greater than is furnished by the estimation. We shall see later that this behaviour is common; two substituents in the ortho-position do not, as is otherwise usual, weaken one another, but reinforce each other's effect.

For the two dioxybenzoic acids still remaining we must expect the following constants:

1:3:4 gives $.006 \times 1.4 \times 0.5 = 0.0042$ 1:3:5 , $.006 \times 1.4 \times 1.4 = 0.012$

Measurement gives '0033 and '009, both constants being therefore as usual somewhat smaller, a consequence of the combined effect, yet being quite of the expected order of magnitude.

The six possible trioxybenzoic acids allow us to estimate the following constants:

Of the six possible trioxybenzoic acids three are known and measured. The comparison between the observed and estimated constants gives 0.055 instead of 0.07; 0.004 instead of 0.006, where as usual the observed value is smaller than the estimated value; and in the case of the acid with two hydroxyls in the ortho-position we have 2.1 observed instead of 0.9 estimated, which therefore, as before, is about twice as great. In the same way we may expect in the cases of the three trioxybenzoic acids still unknown, that the real constants of the acids 1:2:3:5 and 1:2:4:5 will be somewhat smaller than the estimated values, and therefore about 0.15 and 0.05, while the third acid 1:2:3:6 with two ortho-hydroxyls will probably have a constant from 5 to 5.5. In spite of the uncertainty of the estimation, the differences are so considerable that when in future one of these acids is prepared the determination of the conductivity of an aqueous solution will suffice to decide its constitution.

The constitutional relations of the oxybenzoic acids, as indicated by the dissociation-constants, are therefore in complete agreement with those

deduced from chemical relations.

The methods just expounded may be directly utilised for elucidating constitutional relations. On heating orcin $C_6H_3(OH)_2CH_1$ with KHCO₁ there is formed 'paraorsellic acid,' whose constitution must correspond either to that of α - or β -resorcylic acid. On measurement, the constant appeared to be 4·1; the acid has, therefore, a constitution analogous to that of α -resorcylic acid, corresponding to the arrangement CO₂H:OH:OH:CH₁=1:2:6:4.

Further substitution products of benzoic acid gave the following numbers:—

```
o-Chlorobenzoic acid C_eH,Cl.CO.H m-Chlorobenzoic acid C_eH,Cl.CO.H
                                                                 0.132
                                                                 0.0155
p-Chlorobenzoic acid C<sub>6</sub>H<sub>4</sub>Cl.CO<sub>2</sub>H
o-Bromobenzoic acid C<sub>6</sub>H<sub>4</sub>Br.CO<sub>2</sub>H
m-Bromobenzoic acid C<sub>6</sub>H<sub>4</sub>Br.CO<sub>2</sub>H
                                                                 0.0093
                                                                 0.145
                                                                 0.0137
Monobromgallic
   acid C<sub>6</sub>HBr(OH),CO<sub>2</sub>H
                                                                0.059
Dibromgallic acid C<sub>6</sub>Br<sub>2</sub>(OH)<sub>3</sub>CO<sub>2</sub>H
                                                                1.21
m-Fluobenzoic acid C,H,F.CO,H.
                                                                0.0136
m-Cyanobenzoic acid CaH, CN.CO.H
                                                                0.020
o-Nitrobenzoic acid C<sub>6</sub>H<sub>4</sub>NO<sub>2</sub>.CO<sub>2</sub>H
                                                                0.616
m-Nitrobenzoic acid C,H,NO,CO,H
                                                                0.0345
p-Nitrobenzoic acid CaHANO, CO2H
                                                                 0.040
Bromonitrobenzoic
    acid [1:2:5] CoH3BrNO2CO2H
                                                                 1.4
o-Amidobenzoic acid C,H,NH,CO,H
                                                                 0.0010
p-Amidobenzoic acid C<sub>6</sub>H<sub>4</sub>NH<sub>2</sub>CO<sub>2</sub>H
                                                                 0.0012
o-Acetamidobenzoic
   acid C.H.(NHCOCH,)CO.H
                                                                 0.024
m-Acetamidobenzoic
   acid C<sub>e</sub>H<sub>4</sub>(NHCOCH<sub>8</sub>)CO<sub>2</sub>H
                                                                 0.0085
p-Acetamidobenzoic
   acid C<sub>6</sub>H<sub>4</sub>(NHCOCH<sub>3</sub>)CO<sub>2</sub>H
                                                                 0.0052
o-Acetoxybenzoic
    acid C<sub>5</sub>H<sub>4</sub>(OCOCH<sub>2</sub>)CO<sub>2</sub>H
                                                                 0.038
p-Acetoxybenzoic
```

0.0042

acid C.H. (OCOCH,)CO2H

m-Acetoxybenzoic
acid C₂H₄(OCOCH₃)CO₂H . 0.0099
o-Methoxy-benzoic
acid C₃H₄(OCH₃)CO₂H . 0.0082
p-Methoxy-benzoic
acid C₃H₄(OCH₃)CO₂H . . 0.0032

The same conclusion may be drawn from these numbers as to the influence of the halogens as had been deduced for hydroxyl-namely, that the ortho-position is that of greatest influence. The constant of benzoic acid rises to 22 times its value on the introduction of chlorine in the ortho-position, to 2.6 times its value when Cl is put in the meta-position, and to 1.5 times its value when Cl is put in the para-position. But along with this there is a remarkable difference between the behaviour of OH and that of Cl; hydroxyl acts much more strongly in the orthoposition of the benzene nucleus than in the a- position of the open chain, while chlorine, which in the a-position of the open chain increases the constant to more than 80 times its value, does so only 22 times when in the orthoposition in the benzene nucleus. For this discrepancy the explanation offers itself, on the one hand, that the system of carbon atoms which constitutes the benzene nucleus is not rigid, but alters its form according to the atoms present in it, and that if so, the difference in form must be especially great according as hydroxyl or chlorine enters near the carboxyl. On the other hand, the following possibility strikes us. According to the hypothesis of Kekulé, there are two sorts of ortho-positions, the one being represented by a double, the other by a single, linkage. Now it is quite possible that in benzene itself the change of the double or single linking to the right or the left takes place easily, but that when two adjoining carbon atoms have fixed different atoms or groups of atoms, only one sort of linkage, either a single one or a double one, is stable. Assuming that -in salicylic acid, for instance-double linkage exists between the carbon atoms, combined with

while in o-chlorobenzoic acid the single linkage COOH

sised above can be understood, since in the first case the ortho-relation is a much more intimate one than in the second.

The small changes which the constant of benzoic acid undergoes in m- and p-chlorobenzoic acid correspond to the greater distance of the chlorine from the carboxyl. According to the usual representation of benzene, the influence of the para-position proves to be the smallest. But this behaviour is by no means general.

Bromine behaves very similarly to chlorine. The effect for the ortho-position is somewhat larger than for chlorine—i.e. 24 instead of 22; that for the meta-position somewhat smaller—2·3 instead of 2·6.

When hydrogen in gallic acid is replaced by bromine in the ortho-position, the constant increases to 15 times its value, therefore less than in the case of benzoic acid. This corresponds to the general rule that several substituents simultaneously present diminish each other's action. Only when two substituents, which are the same, occupy the ortho-position is an increase in the simultaneous effect to be observed; in fact, a second bromine atom which occupies the ortho-position brings about a greater rise of the constant than does the first —21 instead of 15.

Hydrofluoric acid being much weaker than hydrochloric acid, it was to be expected that the fluobenzoic acids would be weaker than the chlorobenzoic acids. In case of the meta-compound this occurs, however, only to a small extent; the constants differ but little, and are nearly the same for the bromo- and fluo- compounds. It would be of interest to determine whether in the ortho- position also fluorine acts as strongly as chlorine, but o-fluobenzoic acid has not yet m-cyanobenzoic been examined; markedly stronger than m-chlorobenzoic acid. This completely corresponds to the fact that cyanacetic acid is much stronger than chloracetic acid.

From measurements of the nitrobenzoic acids, the group NO₂ appears to be the most effective negative substituent. The ortho- compound exhibits a 103-fold increase of the constant, the meta- compound an increase equal to 5.8-fold, and the para- compound a 6.7-fold increase. With regard to the influence of position, the numbers agree with those found for the orthor- and bromo- compounds, in so far as the ortho- position is again that of greatest influence; but, on the other hand, the influence of the nitro- group makes itself more felt from the para- position than from the meta- position, in opposition to the relations found in the case of chlorine.

Bromonitrobenzoic acid contains the substituents in the order CO₂H:NO₂:Br = 1:2:5. The constant is made up of that of benzoic acid, and of the factor of the o-nitro- group 103, and that of the m-bromine 2:3, and is therefore calculated to be 1:38, which agrees well with 1:4, the value found by experiment. In this case, therefore, the mutual influence of the simultaneously present substituents is practically zero.

The constants of o- and p- amidobenzoic acid show that the amido- group considerably reduces the strength of the acid. But the values quoted can only be looked upon as approximations, since peculiarities appear in the determinations which are not yet elucidated.

But the numbers become regular when the basic properties of the amido- residue are compensated by the introduction of acetyl. Of the three acetamidobenzoic acids the ortho- compound is the strongest, it surpasses benzoic acid by four times its value. The meta- compound is only 1.4 times as strong as benzoic acid, and in the para-position the atomic group NHCOCH, acts similarly to hydroxyl—that is, it exerts a weakening effect—the constant is only 0.86 of that of benzoic acid. The acetyl-oxybenzoic acids behave quite similarly; their factors re-

latively to benzoic acid are 5.5, 1.6, and 0.76. On comparing with these and the former numbers the factors of the oxybenzoic acids-namely, 17, 1.4, and 0.5—it follows that the substituent which in the ortho-position acts most strongly also exerts the greatest weakening influence in the para- position.

But this relation again is not universal. Replacement of hydroxylic hydrogen by methyl in salicylic acid produces a quite extraordinary weakening of the acid; instead of 17 times, the constant is only 1.4 times, as great as that of benzoic acid. In the para-position, however, the same change produces scarcely any effect; the constants of p-oxybenzoic acid and of p-methoxy-benzoic acid or anisic acid are nearly the same, and the decreasing effect of the group OCH, is much greater in the para-than in the ortho-position.

It is further remarkable that acetosalicylic acid is weaker than salicylic acid, in spite of acetyl being a negative radicle. It may be that the cause of this curious behaviour is to be ound in a change of linkage similar to that mentioned above.

Homologues of benzoic acid.

o-Toluic acid CaH4(CH4)CO2H 0.012 m-Toluic acid C_sH₄(CH₃)CO₂H 0.0051 p-Toluic acid C,H,(CH,)CO,H 0.0052Phenylacetic acid C₈H₈.CH₂.CO₂H 0.0056Phenylpropionic acid CH3.CH(C6H8)CO2H . 0.0043p-Phenylpropionic acid CH₂(C₆H₅)CH₂.CO₂H . 0.0023Cuminic acid C₈H₄(C₈H₇)CO₂H 0.0050

A comparison of the constant of benzoic acid with those of the toluic acids shows that a doubling of value has occurred by the introduction of methyl into the ortho-position, but that, as usual, the methyl group in the meta- or paraposition has a slightly weakening effect. isomeric phenylacetic acid also has a very similar constant. This constant is 3.1 times greater than that of acetic acid—an indication that phenyl is a negative radicle—but it does not attain to the value of benzoic acid.

The constants for the two phenylpropionic acids show that, as in the case of α - and β - oxypropionic acids, the acid in which the phenyl is at a greater distance from the carboxyl is the weaker. The other relations to be expected are also found; between propionic acid and α-phenylpropionic acid there is the same ratio of the constants, 3.1, as between acetic and phenylacetic acid

Cuminic acid differs from p-toluic acid only in that it contains isopropyl instead of methyl. To this small difference in constitution corresponds as small a difference in the constants.

 $\label{eq:Hydroxylised homologues of benzoic acid.} \textbf{Mandelic acid } \textbf{C}_{\textbf{e}\textbf{H}_{\textbf{5}}}.\textbf{CH}(\textbf{OH})\textbf{CO}_{\textbf{2}\textbf{H}} \qquad . \quad 0.0$ 0.0417 Phenoxy-acetic acid C_eH₅O.CH₂.CO₂H . 0.0756 o-Nitrophenylglycollic acid C₀H₄(NO₂)O.CH₂.CO₂H 0.158p-Nitrophenylglycollic acid C.H. (NO.)O.CH. CO.H.
Tropic acid CH. (OH).CH(C.H.)CO.H. 0.153 0.0075 Phloretic scid CH,CH(C,H,OH)CO2H 0.0020Hydroparacoumaric acid CH2(C.H4.OH)CH2.CO2H. . 0.0017

Mandelic acid is phenylated oxyacetic acid; the ratio of its constant to that of glycollic acid is 2.7, while that between acetic acid and phenylacetic acid is 3.1; both values approximate closely to each other.

On introducing into the phenyl of phenoxyacetic acid a nitro-group in the ortho-position, the constant rises to double its value. The introduction of NO2 in the ortho-position into benzoic acid raises the constant to 103 times its former value. This enormous difference exemplifies most prominently the influence exerted by the varying distance of the substituent from the carboxyl. The same thing is made evident by the constant of p-nitrophenylglycollic acid, which is scarcely different from that of the ortho- compound; the change from the ortho- to paraposition, which is generally so influential, is of scarcely any account at this distance from the carboxyl.

Tropic acid is α-phenyl-β-oxypropionic acid. It is $2.\overline{b}$ times stronger than oxypropionic acid. For the influence of phenyl in the a- position we have before found the factors 2.7 and 3.1.

Phloretic acid is phenylpropionic acid whose phenyl contains a hydroxyl in the para-position; its constant is half as great as that of phenylpropionic acid; hence the para- hydroxyl here exerts its characteristic weakening effect.

In the same way hydroparacoumaric is hydrocinnamic acid which contains a parahydroxyl in the phenyl. This also has a decreasing action on the constant, but owing to the greater distance of the phenyl this effect is

not so great as in the preceding case.

Unsaturated acids. The affinity co-efficients of these substances present a special interest owing to their isomeric relations, which cannot be represented by the ordinary chemical formulæ. Following up an idea of van't Hoff, an explanation of these phenomena has lately been given by Wislicenus (Abh. König. Sächs. Ges. der Wiss. 1887). In this explanation the cause of the chemical differences is supposed to lie in differences in the relative positions in space of the radicles linked to the carbon. Since the results so far obtained from a consideration of the affinity-constants invariably point to relations in space, it becomes possible to subject this hypothesis to an experimental examination. Acrylic acid CH2:CH.CO2H 0.0056

Crotonic acid CH(CH₂):CH.CO₂H 0.00204 Isocrotonic acid CH(CH,):CH.CO2H 0.0036 Tiglic acid CH(CH,):C(CH,)CO,H 0.00098Angelic acid CH(CH₂):C(CH₂)CO₂H 0.0050a-Chlorocrotonic acid CH(CH₂):CCl.CO₂H . 0.0720allo-a-Chlorocrotonic acid CH(CH2):CCl.CO2H. 0.158

B-Chlorocrotonic acid CCl.(CH₂):CH.CO,H 0.0144allo-\$-Chlorocrotonic acid CCl.(CH₃):CH.CO₂H.

0.0095

A consideration of these numbers shows that the unsaturated acids have in general greater constants than the saturated ones. We must, therefore, ascribe to hydrogen a strongly basic character. And, further, as in the case of the saturated fatty acids, the constants on the whole decrease as the carbon increases. And, finally, we see that the isomeric acids for which, according to the views hitherto held, no difference of constitution could be formulated are in fact distinctly different.

Now Wislicenus represents the difference between crotonic acid and isocrotonic acid by the following formulæ:--

$$\begin{array}{c} \text{Crotonic acid} \\ H > C = C < \stackrel{CH_3}{<} \\ \text{CO}_2H \end{array} \quad \text{and} \quad \begin{array}{c} \text{Isocrotonic acid} \\ CH_3 > C = C < \stackrel{H}{<} \\ CO_2H \end{array}$$

In crotonic acid the methyl group stands nearer to the carboxyl, in isocrotonic acid the hydrogen. Since, in accordance with present experience, a methyl group linked to carbon which is also directly linked to CO2H decreases the constant, a smaller constant must be expected for crotonic acid than for isocrotonic, this difference being more pronounced owing to greater proximity in space. Experiment confirms this conclusion, and thus proves the appropriateness of Wislicenus' assumption.

It may be asked whether we are not here dealing with one of the cases of which some were mentioned before, in which, as an exceptional case, the methyl acts by increasing the strength. But this view is excluded, since both acids are weaker than acrylic acid. Methyl has, therefore, in both cases reduced the strength, but to a greater degree in crotonic than in the

iso- acid.

Wislicenus assumes similar differences in constitution for angelic and tiglic acids. We get

Angelic acid
$$CH_3$$
 $C=C$ CO_2H CH_3 $C=C$ CO_2H CH_3 $C=C$ CO_3H CH_3 $C=C$ CO_3H CH_3

Hence angelic acid again would have to be stronger than tiglic, as has been verified by experience. But it is remarkable that in this case the difference has been found to be much greater than in the first one. Angelic acid has a greater constant than isocrotonic acid, which contains CH2 less than angelic acid, a fact as unusual as that tiglic acid exhibits a smaller constant than the corresponding saturated acid, valeric, which has for constant 0.00161.

In the same way the values found for the a-chlorocrotonic acids agree with experience. The constants are very nearly in the same ratio as that of those of the two crotonic acids themselves, and the rise effected by the chlorine introduced into the a-position is represented by the factors 36 and 44, which, though markedly smaller, are of the same order of magnitude as

the factor for monochloracetic acid.

In opposition to the agreement holding hitherto between theory and experiment, in the case of the β-chlorocrotonic acids, the allo- compound proves distinctly weaker than the deriva-tive of crotonic acid. The cause of this discrepancy cannot yet be established. The ratio of the constants for crotonic acid and β-chlorocrotonic acid is 1:7, which is nearly the same as the corresponding factor for propionic acid and β-iodopropionic acid, viz. 6.6. On the other hand, the ratio for the factors of isocrotonic acid and the allo-β-chloro- derivative is 1:2.6, and is, therefore, much smaller.

Tetrolic acid is formed from both \$\beta\$-chlorocrotonic acids by splitting off hydrochloric acid; the constant is

Tetrolic acid CH, C:C.CO.H .

Though hydrochloric acid is one of the strongest acids, yet the splitting off of its elements from the β -chlorocrotonic acids not only has not lowered the constant, but has raised it This result makes evident the considerably. strongly acid nature of carbon and the basic nature of hydrogen; the removal of four hydrogen atoms from butyric acid, whose constant is 0.00149, has produced a rise in the constant of 160 times its value.

Of other unsaturated acids the following have been measured-

Hydrosorbic acid C_5H_9 . CO_2H 0.00241Sorbic acid C₅H₇.CO₂H

Parasorbic acid C₅H₇.CO₂H 0.001730.00173Ethyl-methacrylic acid C,H,.CO,H 0.00111 Atropic acid CH₂:C(C₆H₅)CO₂H . Cinnamic acid CH(C₆H₅):CH.CO₂H 0.01430.00355Isocinnamic acid CH(C,H,):CH.CO,H . 0.0156p-Coumaric

acid CH(C₆H₄.OH):CH.CO₂H 0.0022

o-Coumaric acid CH(C₆H₄.OH):CH.CO₂H 0.0021Umbellic acid

0.00188

 $CH(C_6H_3(OH)_2):CH.CO_2H$

a-Bromo-cinnamic acid CH(C6H5):CBr.CO2H 1.44

B-Bromo-cinnamic

as chemically identical.

acid CH(C₆H₅):CBr.CO₂H 0.093 Phenylpropiolic acid C.H. C.C.CO.H . 0.59

o-Nitro-phenylpropiolic acid C₆H₄NO₂.C:C.CO₂H 1.06

In accordance with the general rule, hydrosorbic acid is stronger than the corresponding saturated compound, which is caproic acid (k = 0.00145), while sorbic acid, which contains 2H less, is not stronger but weaker than hydrosorbic acid. It is, therefore, probable that sorbic acid has not one treble linkage, but two double ones, or it may have an altogether different constitution, perhaps one with a closed chain arrangement of the carbon atoms.

Sorbic acid is sometimes obtained in the form of an oil which only when treated with acids or alkalis passes into the crystalline acid. Since it did not seem impossible that this was a case of isomerism, such as that of crotonic and isocrotonic acid, the constant of the liquid parasorbic acid was also measured. It proved identical with that of sorbic acid, and hence, with Fittig and Baringer (A. 161, 325 [1871]), these substances must be considered, not as isomeric, but

The constant of methylethylacrylic acid is remarkably small; it is smaller than that of the saturated fatty acid, and approaches that The acids a-phenylacrylic, or of tiglic acid. atropic, and \$\beta\$-phenylacrylic or cinnamic, behave as we should expect; the phenyl group causes a much higher value of the constant when it is in the a- than when it is in the β - position. But it is remarkable that cinnamic acid is not only weaker than atropic acid, but even weaker than acrylic acid; in this case, then, the phenyl group lowers the constant.

In the case of cinnamic acid, two 'geometri. cally isomeric' compounds may be expected, namely-

$$_{C_{u}H_{u}}^{H}$$
C=C $<_{CO_{u}H_{u}}^{CO_{u}H}$ and $_{H}^{H}$ C=C $<_{CO_{u}H_{u}}^{CO_{u}H}$

Quite recently there has been discovered by Liebermann a second cinnamic acid which must be regarded as geometrically isomeric with the ordinary one; it has a much higher constant than the latter, and hence we may conclude that in the new isocinnamic acid the negative phenyl has a closer connexion with the carboxyl than in the old one, and that the new acid may have the second of the above formulæ.

Similarly, there are known two isomeric bromocinnamic acids, which are produced from cinnamic acid dibromide by splitting off hydrobromic acid, and for which the same kind of isomerism holds. Nor can there be any doubt as to which acid corresponds to each formula, the differences in the constants being very great, and in fact it has been made known quite recently by Erlenmeyer that he has obtained Liebermann's isocinnamic acid from the stronger a-bromocinnamic acid.

The phenylpropiolic acid formed from the bromeinnamic acids by splitting off hydrobromic acid is stronger than the β -compound, but weaker than the isomeric α -compound. The constant is also, as was to be expected, greater than that of tetrolic acid. In o-nitro-phenylpropiolic acid the nitro-group in the orthoposition has again been able to exert but a slightly strengthening action, owing to its being at a

great distance from the carboxyl.

On comparing cinnamic acid with p-coumaric acid, we find that the hydroxyl introduced into the p- position again lowers the constant. But, strange to say, a similar effect is produced in the derivatives of cinnamic acid by hydroxyl in the ortho- position; o-coumaric acid is also weaker than cinnamic acid. It is in accordance with this that umbellic acid, which contains one hydroxyl in the ortho- position and a second one in the para- position, is weaker than all these acids. It is to be expected that these relations will prove of importance for the complete elucidation of the constitution of these remarkable compounds.

Dibasic acids. The acids of the oxalic series

gave the following constants:

It has already been explained that the weaker dibasic acids H.R. split electrolytically according to the scheme H + HR, and that therefore constants may be calculated for these acids in a manner analogous to that adopted for the monobasic acids. In the case of oxalic acid, however, this is not possible, since the dissociation has proceeded too far; the constant is only estimated approximately, but beginning from malonic acid, the values of the constants can be very well determined.

We see that at first the constants decrease very rapidly; that of malonic acid is about 60 times smaller than that of oxalic acid. From malonic to succinic acid the step is much smaller, 24: 1, and much smaller from the latter to pyrotartaric acid, 14: 1, and henceforward

the numbers vary only by small amounts. The influence which one carboxyl group exerts on the hydrogen of the other decreases, therefore, very rapidly in consequence of the interposition of the carbon atoms; sebacic acid does not surpass by more than twice the fatty acids richer in carbon. Of the numerous isomerides with side carbon chains, the following have been measured:

β-Dimethylsuccinic

The amido- group entering into one of the carboxyls of oxalic acid causes a weakening of the acid to one-twelfth its value; the much more weakly basic urea residue produces a much smaller weakening effect, viz. to about half the value. The decrease caused by the residue of aniline is smaller also, to about the eighth part; when chlorine enters into the phenyl of the latter, there again occurs an increase in the acid properties, which increase, however, in consequence of the greater distance, is much smaller than in benzoic acid. But a distinct difference between the ortho- and the para- position still asserts itself. Methyl in the para- position exerts a distinctly weakening effect.

 $\begin{array}{lll} \textbf{Tartronic acid CH(OH)(CO,H)}_2 & & \textbf{0.107} \\ \textbf{Phenylmalonic acid (C_6H,)_2C:(CO_2H)_2} & & \textbf{0.408} \\ \textbf{Malonanilic acid CH_2CO(NHC_6H_3)CO_2H} & & \textbf{0.0196}. \end{array}$

Strange to say, in spite of the hydroxyl. tartronic acid is weaker than malonic acid (k=0.158). The replacement of both hydrogen atoms of malonic acid by benzylic radicles raises the constant. Finally, malonanilic acid is abouteight times weaker than the parent substance.

Derivatives of succinic acid.

Malic acid C₂H₃(OH)(CO₂H)₂ . . . 0.0395
Inactive malic acid C₂H₃(OH)(CO₂H)₂ . 0.0399
Dextrotartaric acid C₂H₄(OH)₂(CO₂H)₂ . 0.097
Lawotartaric acid C₂H₂(OH)₂(CO₂H)₂ . 0.097
Racemic acid C₂H₂(OH)₂(CO₂H)₂ . 0.097
Mesotartaric acid C₂H₂(OH)₂(CO₂H)₂ . 0.060
Phenyl-succinamic
acid C₂H₄(CONHC₄H₅)CO₄H . 0.002C

o-Chlorophenylsuccinamic acid C₂H₄(CONHC₆H₄Cl)CO₂H . 0.0021 m-Chlorophenylsuccinamic acid C₂H₄(CONHC₆H₄Cl)CO₂H 0.0021p-Chlorophenylsuccinamic acid C₂H₄(CONHC₆H₄Cl)CO₂H 0.0021o Tolyl-succinamic acid C₂H₄(CONHC₇H₇)CO₂H 0.0021p-Tolyl-succinamic $acid C_2\mathbf{H}_4(CONHC_7\mathbf{H}_7)CO_2\mathbf{H}$ 0.0020 Succinuric acid C2H4(CONHCONH2)CO2H 0.00311Thio-succinuric acid C2H4(CONHCSNH2)CO2H 0.00333

Oxysuccinic, or malic, acid has a constant six times as great as succinic acid. The factor is somewhat smaller than for the monobasic acids, which may be because a negative radicle carboxyl is already present. The natural optically active acid has exactly the same constant as the inactive acid got from monobromsuccinic acid. The same relation recurs in tartaric acid and racemic acid. Both the dextrorotatory and the lavorotatory acid, as also the inactive racemic acid formed by a combination of these two, are all of exactly the same strength. This result is in accordance with the present views concerning the constitution of these compounds. dextrorotatory and levorotatory acids contain their constituents in a perfectly symmetrical arrangement, and must, therefore, have equal co-efficients of affinity. It has been assumed that the optically inactive acids produced from the optically rotating ones consist each of two molecules of the active acids. This is probably correct for the solid crystallised substances, but not for solutions; in these, both compounds are present side by side, uncombined, a conclusion which has also been arrived at in another way by Perkin (C. J. 52, 362). The undecomposable inactive mesotartaric acid has, however, another, and that a smaller, constant; theory also ascribes to it a different constitution.

In the preceding table are further contained a number of derivatives of phenylsuccinamic or succinanilic acid. This acid itself is only 3.3 times weaker than succinic acid, while the introduction of the aniline residue into CO2H of malonic acid decreased the strength 8 times; this indicates that the position of the substituent at a greater distance weakens the effect. The almost complete equality of the constants for all substitution products of succinanilic acid is of great interest; the distance of the benzylic residue from the carboxyl is here too great to allow the substituents to exert their effect over it. The usual assumption, that the molecular forces cease to act at even very small distances, is thereby made more definite, in that such distances are already given by the molecular dimensions.

As already found, the introduction of the urea residue produces a smaller decrease than that due to aniline. Smaller still is the effect of the thio-urea residue.

Dicarboxylic acids of benzene.

o-Phthalic acid C₆H₄(CO₂H)₂.

o-Phthalic acid C₆H₄(CO₂H)₂.

o-25

Oxyterephthalic acid C₆H₄(CONH₂)CO₂H

O-0160

Phthaluric acid C₆H₄(CON₂H₃CO)CO₂H

o-0290

As we were led to expect, the entry of a carboxyl in the ortho-position produces a somewhat considerable rise in the constant—vis. to twenty times the former value. For the metaposition the factor is only 4.8. p.Phthalic acid is too insoluble to be measured. But assuming that in oxy.p-phthalic acid the hydroxyl acts as strongly as in salicylic acid, the constant of p-phthalic acid would be estimated as 0.014 to 0.015, and it would follow that, in accordance with the usual result, carboxyl in the paraposition—i.e. in the distant one—exerts the least action. A weakening effect, such as is exerted by hydroxyl in the paraposition, seems to be excluded.

The amide residue in phthalamic acid has lowered the constant to one-eighth its value, but the acid is yet stronger than benzoic acid, and the group CONH₂ must be looked upon as negative. The urea residue exerts, as it always does, a much less marked weakening effect.

Unsaturated dibasic acids. The dicarboxylic

Unsaturated dibasic acids. The dicarboxylic acids $CH_{2n-2}(COOH)_2$ exhibit very similar isomeric relations to those of the compounds of the acrylic acid series. The following were measured:—

The isomeric compounds are seen to have widely different constants. These results agree very well with the views of van't Hoff and Wislicenus. For fumaric and maleic acids the following formulæ have been adopted:

$$\begin{array}{ccc} & & & & & & & & \\ & & & & & & & & \\ H > C = C < & & & & & & \\ CO_2H & & & & & & \\ & & & & & & \\ CO_2H & & & & & \\ \end{array} \\ \begin{array}{cccc} & & & & & & \\ & & & & & \\ CO_2H > C = C < & \\ H \end{array}$$

According to this, the two carboxyl groups are much nearer to each other in maleic than in fumaric acid, and the first must be stronger, as is actually the case. Similarly, according to Wislicenus, the formulæ

$$\begin{array}{ccc} \text{Citraconic acid} & & \text{Mesaconic acid} \\ \text{H} & \text{C=C} < \begin{array}{c} \text{CO}_2\text{H} \\ \text{CO}_2\text{H} \end{array} & & \begin{array}{c} \text{CH}_s \\ \text{CO}_2\text{H} \end{array} > \begin{array}{c} \text{C=C} \\ \text{CO}_2\text{H} \end{array}$$

hold, and for the same reason citraconic acid must be stronger than mesaconic acid, which conclusion again agrees with experiment.

 $CH_2: C\longrightarrow CO_2H$ Itaconic acid has the formula | $H_2C - CO_2H$ and therefore no double linkage between the carbon atoms combined with carboxyl. Accordingly, its constant is much smaller; it is only

twice as great as that of succinic acid.

It may be mentioned that acetylene dicarboxylic acid C—CO₂H is so strong that a constant cannot be determined for the first stage of dissociation. The acid is but little inferior to sulphuric acid.

Thiophene and pyridine group.

a-Thiophenic acid C ₁ H ₂ S.CO ₂ H			0.030
Pyromucic acid C ₄ H ₃ O.CO ₂ H		•	0.071
Picolinic acid (a) CaHaN.COaH			0.0003
Nicotinic acid (\$) C.H.N.CO.H			0.00137
Isonicotinic acid (7) C.H.N.CO.	H		0.00169
	H		0.00109

0.011

Lutidinic acid (ay) C ₅ H ₃ N.(CO ₂ H) ₂		0.60
Cinchomeronic		
acid $(\beta\gamma)$ C ₅ H ₈ N. $(CO_2H)_2$		0.21
Isocinchomeronic		
acid ($\alpha\beta$) C ₅ H ₃ N.(CO ₂ H) ₂ .		0.43
Chinolinic acid $(\alpha\beta)$ $C_bH_3N.(CO_2H)_2$		0.30
ββ-Pyridine dicarboxylic		
acid CaHaN.(COaH)2		0.15
The replacement of the group C.H.	in.	benz

acid by sulphur raises the constant to five times its value, provided the carboxyl is placed next to the sulphur. In the β -position the carboxyl is without doubt influenced less, but we have yet no measurement of the constant.

It is remarkable that a similar replacement of C2H2 by oxygen, forming pyromucic acid, acts much more strongly than a replacement by sulphur. On account of the magnitude of the constant it is very probable that in pyromucic acid the carboxyl is in a position next to the oxygen.

Differently from sulphur and oxygen, the replacement of C₂H₂ by CNH has a strongly weakening effect on the constant. The values found for the monocarboxylic acids of pyridine correspond fairly with the assumptions which might be made on the basis of the constitutions of these substances; picolinic acid, which contains the carboxyl next to the nitrogen, is by far the weakest; but the β -acid is stronger than the y- compound, so that in pyridine the β- position proves the place of smallest influence.

The behaviour of the dicarboxylic acids of pyridine is very unexpected. While all monocarboxylic acids of pyridine are weaker than benzoic acid, all dicarboxylic acids are stronger than the strongest dicarboxylic acid of benzene -that is, o-phthalic acid, whose constant is 0.12. A further difference is that the strongest dicarboxylic acids are those which contain carboxyl in the a- position—that is, next to the nitrogen. Then follow the acids which contain carboxyl in the γ - position, and lastly those with carboxyl in the β - position. These relations are the exact reverse of those observed for the monocarboxylic acids; where a decrease of the constant had been observed there, an increase is found here. No plausible hypothesis concerning the cause of these peculiar phenomena has yet been brought forward. Oxycinchomeronic

acid C ₅ H ₂ (OH)*N.(CO ₂ H) ₂ .			1.67
Methyl pyridine dicarboxylic			
acid $C_3H_2(CH_3)^2N.(CO_2H)_2^{\beta\beta}$.			0.20
Dimethyl pyridine dicarboxylic			
acid $C_5 H(CH_3)_2$ and $N(CO_2H)_2$ $\beta \beta 1$			0.34
Dimethyl pyridine dicarboxylic			
acid $C_3H(CH_3)_2^{a\gamma}N.(CO_2H)_2^{\beta\beta}$			0.55
Owing to the entry of hydroxyl	in	the	ortho.

position, cinchomeronic acid has had its constant increased to eight times its value. methyl groups increase the strength of the \$\beta\beta\$. dicarboxylic acid, and they do so to a greater extent in the γ - position than in the α - position.

Further, several poly-carboxylic acids of pyridine were examined; the constants cannot, how-ever, be calculated, because the dissociation assumes a somewhat complicated character. But it is evident that the relations found in the case of the dicarboxylic acids concerning the influence of the position of the carboxyl hold good in these cases also. The rules found for

the dicarboxylic acids were also confirmed in the case of methyl derivatives of the tricarb. On the other hand, phenyl on oxylic acids. entering into lutidine dicarboxylic acid, whose constant is 0.34, has a weakening effect, the constant for the compound formed being only 0.012. The action here is the reverse of that observed in the case of the fatty acids.

Quinaldinic acid C.H.N.CO.H 0.0012 Cinchonic acid C,H,N.CO,H. 0.0013 Dipyridyl monocarboxylic acid C₁₀H₇N₂.CO₂H Dipyridyl dicarboxylic 0.002 acid C10H6N2(CO2H)2 0.032 o-Pyridine benzoic acid C₁₁H₈N.CO₂H . 0.0005 Phenyl pyridine dicarboxylic

acid C, H, N. (CO, H), The quinoline carboxylic acids, which have a constitution analogous to that of the pyridine carboxylic acids, are somewhat stronger than these, as the naphthoic acids are stronger than the benzoic acids. Concerning the other acids but little can be generalised.

Amidobenzene sulphonic acids and similar compounds.

7.000		
o-Amidobenzene sulphonic		
acid $C_6H_4.NH_2.SO_7H$.		0.330
m-Amidobenzene sulphonic		
acid C,H,.NH2.SO,H .		0.0185
p-Amidobenzene sulphonic		
acid C, H, NH2.SO, H .		0.0581

The results obtained from measurements of the amidobenzene sulphonic acids are somewhat remarkable. While benzene sulphonic acid itself is so strong an acid that its dissociation constant cannot be determined, its amido-derivatives can be measured with ease; the amido- group has, therefore, as it usually does, exerted a weakening effect. But in the case of the carboxylic acids the ortho- position was always that of greatest influence; here it appears as the position of least influence. The weakest of the three acids is the meta-compound, then follows the para- compound, and then the ortho- compound, which comes nearest to the parent substance. The assumption is thus suggested that, by the agency of the group SO2, the hydrogen is brought to such a distance from the benzene nucleus that it is more influenced by the radicles in the meta-position than by those in the orthoposition.

Diamidobenzene sulphonic 0.0050acid (2,3) $C_0H_3(NH_2)_2SO_3H$ Bromamidobenzene sulphonio acid (2,5) C_sH_sBrNH₂SO_sH 7.9Bromamidobenzene sulphonic acid (3,6) C₆H₈BrNH₂SO₃H 0 072 Dibromamidobenzene sulphonic acid (3,4,6) C₆H₂Br₂NH₂SO₃H 2.5 Toluidine sulphonic acid (3,4) C₈H₃(CH₂)NH₂SO₂H . 0.024Xylidene sulphonic acid (4,2,5) $C_6H_2(CH_3)_2NH_2SO_8H$ 0.044 In indicating the constitution, the sulpho-group is put = 1, then follows amide, and then bromine

or methyl. These acids exhibit the usual relations to be expected from the nature of the substituents and their position, and thus do not call for any detailed discussion.

General considerations. The properties of chemical compounds, which can be expressed numerically, may be divided into three groups—the additive, the cumulative, and the constitutive. The first are of such a nature that in chemical compounds their value is the sum of the corresponding values of the constituents; these values have led to the conception of the atomic theory—i.e. to the assumption that the components continue to exist in the compounds as such.

The cumulative properties exhibit the peculiarity that, independently of the number and nature of the constituents, their numerical value is the same for certain complex systems. These have led to the conception of the molecular theory—i.e. to the assumption that the components of chemical compounds primarily form similar groups or systems, which have an individual existence, and out of which the visible and ponderable substances are built up.

And, finally, there exists a third group of properties, which are neither additive nor cumulative, since even in the case of substances having the same composition they can assume different values. These have led to the assumption that, in addition to the nature and number of the composing atoms, the properties of substances are influenced by still another condition. This condition has, from the time of Berzelius, been considered to lie in the different arrangement of the atoms within the molecule—i.e. in the constitution.

Of all the constitutive properties hitherto known—such as boiling-point, melting-point, colour, crystalline form, &c.—not one affords a clear insight into the determining cause, the chemical constitution. It is possible, when the latter has been found, more or less approximately, by purely chemical methods, to establish empirical relations; but these lie on the surface only, and lack generality. The cause of this is the defective development of our conceptions concerning the kind and the amount of the influence which the arrangement of the elementary atoms in the molecule exerts on the properties mentioned.

I believe I am not deluding myself when I affirm that the affinity constants, which represent a property of eminently constitutive nature, are far superior to all others with respect to their theoretical applicability. The result that one and the same atom exerts quite different effects, according to the 'position' which it occupies in the molecule, and that these effects are the greater the more close is the relation of the atom to that of the acid hydrogen, leads to the general inference that these effects are functions of the distance in space of the atoms in question. But thereby, for the first time, is gained a means of taking measurements in space of molecular structures. There is no doubt that long and laborious work is needed before such measurements will lead to results which will give an allsufficient representation of the form of molecules. But it already seems certain that this goal can be reached.

It has been mentioned, to start with, that k, the affinity co-efficient of acids, appears, generally speaking, as a product whose factors are given by the nature and position of the elementary

atoms composing the acid. This follows from the fact that with analogous changes in analogous substances the constants change in the same ratio-i.e that in consequence of these changes the constants of the original substances must be multiplied by the same factor. But these factors are never exactly the same, just because two analogous changes are never exactly the same. If, for instance, we change acetic acid into glycollic acid, and propionic acid into lactic acid, the two changes are, it is true, very much alike; but in the first case the hydroxyl takes up a position next to two hydrogen atoms, in the second case next to one hydrogen atom and one methyl; the hydroxyl is therefore subject to a different influence, and will in consequence necessarily act differently on the carb-The same holds for all corresponding cases. The approximate character of the general relation follows, therefore, as a necessity; the deviations from the type are caused by the nature of the subject, and themselves afford a means for bringing to light their cause, which is the secondary influence mentioned above. What has been said concerning the constitutive properties generally—that they can never be completely represented by a general scheme, because this is contradictory to their natureholds particularly for the affinity co-efficients. In them the whole variety of nature asserts itself, and within the framework of great and general regularities produces the finest individualisation.

The form of the dissociation-constant as the product of factors $c=c_1.c_2.c_3...$ has a deeper significance, which is recognised on going back to the dissociation-formula itself. This formula is obtained from the mechanical theory of heat in the following form:

$$\log \frac{p}{p_1 p_2} = \frac{\rho}{R \tau} + C$$

where p_1 and p_2 are the partial pressures of the products of decomposition, p the pressure of the undecomposed substance, ρ the heat of dissociation, r the absolute temperature, R the gaseous constant, and C is a value which is a function of the temperature only, and which, therefore, at constant temperature can be looked upon as a constant.

In the case in point $p_1 = p_2$, and since p_1 and p must be put proportional to $\frac{m}{v}$ and $\frac{1-m}{v}$, it

follows that
$$\frac{p}{p_1^2} = \frac{(1-m)v}{m^2}$$
 and thence $\log. \frac{(1-m)v}{m^2} = \frac{\rho}{R\tau} + C$ or $\log. \frac{m^2}{(1-m)v} = -\frac{\rho}{R\tau} + C^1$.

Now the affinity co-efficient is $k=\frac{m^2}{(1-m)v}$; and since, on the other hand, quite generally $k=k_1.k_2.k_3...$ it follows that

$$\log k_1 + \log k_2 + \log k_3 + \dots = -\frac{\rho}{R_{\sigma}} + C^1$$

The constant C¹ depends only on the unit adopted and on the point chosen as zero; if, therefore, there appears on the left side of the equation a sum of terms represented by log.k, there must correspond to each of these on the right side a term of the form $-\frac{\rho_n}{R_T}$, so that $\frac{\rho}{R_T}$ also resolves

itself into a sum of corresponding terms

 $-\frac{1}{R_{\pi}}(\rho_1+\rho_2+\rho_2+\ldots)$ in which one R_n and

one ρ_n always correspond to each other. This means: The natural logarithm of the

dissociation-constant or affinity-constant is (save for a constant) proportional to the heat of dissociation of the acid in splitting into ions. Since the value of the dissociation- or affinity-constant is known from experiment to be the sum of a series of terms which depend on the nature and on the position of the constituent atoms, it follows that the electrolytic heat of dissociation also is the sum of a corresponding number of terms each of which is fixed by the nature and

position of each individual atom.

Now in this case the heats of dissociation are the exact measures of the quantities of work which are done in the separation of the acid hydrogen atom from the negative ion. This is so because no external work has to be taken into account, and because the condition of the substances dealt with very closely approaches that of ideal gases. Hence the heat of dissociation measures the potential or the force-function of the atomic system at the corresponding point, and we see that this force-function is the sum of the values which, according to their nature and position, the individual atoms contribute to the total value. And thus elements which, like chlorine, sulphur, &c., raise the constant k, cause a decrease of the heat of dissociation, since log.k and ρ have different signs. The amide group, on the other hand, which decreases the constant, raises the heat of dissociation-i.e. it increases the amount of work necessary for splitting off the acid hydrogen.

It is by these considerations that the longmooted question concerning the connection between chemical affinity and the thermal values of reactions has been decided in principle. In this result we see the eminently constitutive property of affinity-quantities brought back again to the additive form. This has been effected by introducing the influence of constitution, or of position in space, into the terms themselves. A similar development awaits the theory of all constitutive properties, as, in fact, we are forced to conceive the combined action of different elements as the sum of various effects.

Electrical theory of chemical affinity. Elsewhere (Affinity, vol. i. p. 67) the laws of chemical affinity have already been explained. Nothing definite could be stated concerning their cause; but it followed from the close connection which was shown to exist between the co-efficients of chemical reacting power of electrolytes, and the values of the electrical conductivities, that an intimate relation exists between these two properties. The prospect is thus opened of carrying out in an exact form the electro-chemical theory of affinity which was rather conjectured than worked out by Davy and Berzelius.

The new view which affords such a possi-

bility is the common cause underlying electrolytic conduction and chemical reactions as recognised by Arrhenius, vis. the existence of free ions. On making the assumption, which is justified by i in the affinity-quantities. Vol. IV.

this concordance, that chemical processes between electrolytes take place only by the agency of the free ions, the laws of chemical affinity, which have been established empirically, can be deduced so as to be in accordance with experiment. To do this we only further need van't Hoff's theory of solutions, and the theory of the laws of dissociation as deduced for gases (v. Dissociation in vol. ii.), which laws, thanks to van't Hoff's theory, can also be applied to substances in solution.

The general law of chemical affinity first recognised is that manifestations of this affinity on the part of a definite substance are expressible by a definite co-efficient, dependent on the temperature and dilution, but independent of the nature of the chemical reaction in which

the substance participates.

Now, if the reacting power of a substance in solution depends on the number of dissociated ions which are present, it can be seen at once that the above-mentioned law must of necessity hold good. So, for instance, the state of dissociation of an acid determines the velocity with which it must act on methylacetate, acetamide, marble, &c. The nature of the other substance determines a factor only, which in analogous cases is the same.

Hence we must conclude that all acids, for instance, when completely dissociated are equally strong. The strong inorganic acids HCl, HBr, HI, HNO₂, HClO, must be classed here, as also the sulpho- acids and similar organic acids. No one of these substances exhibits any of the specific differences discussed in former paragraphs for acids of medium strength. They all exert a nearly equal effect when used in equivalent quantities, and approach the maximum conductivity even at moderate dilutions. At a time when the dissociation theory of electrolytes did not exist, the author of this article vainly tried to obtain from benzene-sulphonic acid, by all the means which make weak acids strongersuch as the introduction of halogens, nitrogroups, &c .- acids stronger than the parent substance, which may itself be ranged side by side with hydrochloric acid. This was not possible; all 'strengtheners' remained in this case in-effective. This remarkable fact shows that it is not its special nature, but its electrical condition, which is determinant for the chemical effect of a substance, especially if that substance be an acid.

The greater the number of the molecules which are split into their positively and negatively charged parts, the stronger is the acid; but any ion, when once present in the separated, electrically charged, condition, is as effective as any other. Here, then, we see that Berzelius' conclusion drawn from false premisses, by means of which he attempted to demonstrate the absurdity of Faraday's law—that then the chemical affinity between the ions of all electrolytes would have to be the same—is perfectly correct. The affinity of the most diverse ions is really the same, because it does not depend on their nature but on their electrical charge, and this, according to Faraday's law, is everywhere the same. What does depend on the nature of the ions is the degree to which they dissociate, and herein alone lies the cause of the differences

In the arguments used so far, it has been assumed that the state of dissociation of the reacting substances is determined by the volume and the temperature, and that the presence of other substances does not alter the state of the one considered. This case, however, only occurs when there are present either one dissociated substance and nothing but non-electrolytes besides, or when all the dissociated substances present are for all practical purposes completely split up. But as soon as several partly dissociated substances are present, the relations become much more complicated.

It is known from the theory of dissociation that when two gaseous substances are present, and the quantities of these substances vary, the product of their active quantities, or of their concentrations, must be constant, constant temperature being presupposed. If, for instance, a given space is filled with the gaseous products of decomposition of ammonium hydrosulphide, the addition of sulphuretted hydrogen will cause a decrease in the quantity of hydrosulphide dissociated, because on increasing the one factor the other must decrease in the same ratio. The same must evidently hold for electrolytes; on adding to a solution which contains the ions A + B an excess of ions A, the quantity of ions B must decrease until the product of the two quantities has again become equal to its original value. Now, it is not very well possible to add ions A only, but an electrolyte can be added which is also dissociated, and which contains the ions A + B'; if B' is different from B, the two do not influence each other.

We may now ask how two solutions must be constituted in order that their state of dissociation shall not vary when the solutions are mixed. We have the solutions of the electrolytes A+B and A+B', which contain a common ion A; let their volumes be v and v'. Then, in accordance with the general law of the equilibrium of dissociation, the following equations will hold:

$$\frac{1-\xi}{v} = k\frac{\xi}{v} \cdot \frac{\xi}{v}$$
$$\frac{1-\xi'}{v'} = k'\frac{\xi'}{v'} \cdot \frac{\xi'}{v'}$$

in which the quantities of the electrolytes originally present are put as equal to 1, and the dissociated portions equal to ξ and ξ' respectively. On mixing the solutions, the volume becomes v+v', the dissociation ratios ξ and ξ' remaining unchanged, as supposed. The quantities of the undissociated portions are $1-\xi$ and $1-\xi'$, but the products $\xi.\xi$ and $\xi'.\xi'$ change into $\xi(\xi+\xi')$ and $\xi'(\xi'+\xi)$, since the ion A common to both substances participates in the equilibrium condition of both. We have, therefore,

$$\frac{1-\xi}{v+v'} = k, \quad \frac{\xi}{v+v'}, \quad \frac{\xi+\xi'}{v+v'}$$

$$\frac{1-\xi'}{v+v'} = k', \quad \frac{\xi'}{v+v'}, \quad \frac{\xi+\xi'}{v+v'}$$

On dividing each of these equations into each of the corresponding equations given above, it follows that $\frac{\xi}{\xi'} = \frac{v}{v'}$. Therefore, in order that the state of dissociation should not alter on mixing solutions of two electrolytes which have one ion

in common (for instance, two acids), the portions dissociated must be in the ratio of the volumes.

Writing the equation in the form $\frac{\xi}{v} = \frac{\xi'}{v'}$, the

law can also be expressed in this form: when the concentration of the common ion is the same in both solutions, these do not influence each other.

Arrhenius, to whom we owe these considerations, calls solutions which do not influence each other in the sense just mentioned isohydric solutions. Since, in accordance with the above formula, it is only necessary for this that the concentration of the common ion in the two solutions should be the same, the ratio of the quantities of the two solutions does not enter into consideration; isohydric solutions can be mixed in all proportions without mutually changing their condition.

If, however, two non-isohydric solutions which contain a common ion are mixed, the process can be represented by saying that the one solution withdraws water from the other until both have become isohydric; then they leave each other uninfluenced. This withdrawal of water occurs in the following manner. First, the total concentration of the solution which loses water increases, and thereby the concentration of the ions increases also. But, owing to this, the state of dissociation changes directly, since, in consequence of the smaller volume, the dissociation decreases, and, therefore, the concentration of the ions increases more slowly than the total concentration. Conversely, the concentration becomes less in the portion which takes up water; but, owing to the increase in dissociation, it does so to a smaller degree than corresponds to the volume change.

Since the relation between the volume and the state of dissociation is known for binary electrolytes (p. 194), a formula can easily be deduced which allows us to calculate the characteristic quantities for a mixture of several electrolytes. But it is generally simpler to proceed by way of successive approximation.

Two solutions which are isohydric with a third must be isohydric with each other; because, in order to be isohydric with the third, there must be in both the same concentration of the common ion as is in the third solution.

These laws were all established empirically by Arrhenius (W. 30, 51), and it was only lately that he deduced them theoretically from his theory of dissociation (Z. P. C. 2, 284 [1888]). The examination of these relations is best conducted by making measurements of electrical conductivities. Since the conductivity of a compound in solution is proportional to the number of free ions, it can easily be recognised whether two substances with a common ion—two acids, for instance—are isohydric; if the conductivity of the mixture is equal to the sum of the conductivities of the components, the solutions are isohydric; if this is not the case, the two numbers differ from each other more or less. To quote an example: with a phosphoric acid solution, of specific conductivity l = 225.6, was mixed an equal volume of hydrochloric acid solution l = 225.9; the mixture gave $2l = 469 \cdot 0$, while the sum of the two values is 451.5; the observed conductivity was too great by 17.5. Further, the same phosphoric acid solution mixed with an equal volume of a more dilute hydrochloric acid, $l=185^{\circ}1$, gave for the conductivity of the mixture $2l=407^{\circ}4$; this is smaller by 3.3 than the sum 410.7. Finally, hydrochloric acid of $l=168^{\circ}8$ gave for the mixture $2l=394^{\circ}4$, the sum being $394^{\circ}4$; both solutions were isohydric. By such experiments as this Arrhenius arrived at the laws stated above.

It need scarcely be mentioned that any other property of solutions which depends on the molecular condition of the dissolved substance could likewise have been used for determining the relations of isohydricity. The best of these properties would be measurements of the velocities of reaction—for instance, those of the inversion of sugar—these being proportional to the state of dissociation; but the freezing-points, vapour pressures, &c., could also have been used.

So far the only case considered has been that in which the mixed electrolytes contain a common ion. If this is not the case, new phenomena appear, which we will now consider in greater detail. It has already been pointed out that the theory of free ions leads to a view of the processes of the formation and mutual decomposition of salts which is very different from that generally accepted. It has been shown that, neglecting the small undissociated portions, the process of neutralisation between the strong acids and bases consists exclusively in the formation of water from the ions H and The cause why these ions coalesce to form undissociated water lies in the fact that water is an extremely stable compound, which, under ordinary circumstances, is only very slightly split into ions. If, therefore, such ions anywhere appear simultaneously, they combine directly to form undissociated water.

The phenomena take a quite similar course when a strong acid-hydrochloric, for instanceacts on the salt of a weak acid-sodium acetate. for instance. The former solution is almost exclusively composed of ions H and Cl, the latter of ions Na and CH₃.CO₂. Since the sodium chloride also is almost completely dissociated when in dilute solution, the ions Na and Cl are already present in the same condition as in sodium chloride solution, and there exists no tendency to the formation of this compound. But, in addition, there are present the ions H and CH₂CO₂, and these cannot exist side by side without for the most part combining. Hence a corresponding quantity of undissociated acetic acid will be formed, as is actually shown by experiment. But while formerly this fact was ascribed to a specially great affinity between the chlorine and the sodium, and the acetic acid was allowed to play a passive part, being supposed to be 'expelled' by the hydrochloric acid, we now see that the entire cause of the process lies in the action of the acetic acid. Owing to the great affinity of its ions, it is formed whenever its ions meet, while the components of the 'strong' reagents, chlorine and sodium, remain perfectly indifferent in the process.

The conditions under which chemical equilibrium is established between four substances,

formed by a combination in pairs of two positive and two negative electrolytes, must now be established in a general manner. We assume we have produced four solutions of the substances A₁B₁, A₁B₂, A₂B₁, and A₂B₂, so that those solutions which contain a common ion are isohydric. When this is the case we can, it is true, mix the isohydric solutions in all proportions; but if we mix all four solutions the simple law stated above evidently no more holds, and we must ask in what proportion the mixture must be made in order that no change in the state of dissociation should occur. Let a, b, c, and d be the relative volumes of the four solutions in which this occurs, and let α , β , γ , and δ be the undissociated quantities of the four substances. We can represent the dissociated quantities by ha, hb, hc, and hd, since, in consequence of the assumption made, there must in all cases be equal concentration of the ions. The equations of dissociation of the solutions thus become

$$\frac{a}{a} = k_1 \left(\frac{ha}{a}\right)^2 \qquad \qquad \frac{\beta}{b} = k_2 \left(\frac{hb}{b}\right)^2$$

$$\frac{\gamma}{c} = k_3 \left(\frac{hc}{c}\right)^2 \qquad \qquad \frac{\delta}{d} = k_4 \left(\frac{hd}{d}\right)^3$$

or, $\alpha = k_1 h^2 a$, $\beta = k_2 h^2 b$, $\gamma = k_3 h^2 c$, $\delta = k_4 h^2 d$.

If we suppose the four volumes to be mixed, new equations will make their appearance which

have the form $\frac{a}{a+b+c+d} = k_1 \frac{h(a+b)(a+c)}{(a+b+c+d)^2}$, &c., and which can be brought to the form

$$\mathbf{a} = \frac{k_1 h^2 (a^2 + ab + ac + bc)}{a + b + c + d},$$

$$\mathbf{\beta} = \frac{k_2 h^2 (b^2 + ab + bd + ad)}{a + b + c + d}, &c.$$

The quantities a, β , γ , and δ are to remain unchanged, according to the assumption. Now, by dividing the corresponding equations by each other, the same equation follows from each pair ad = bc,

i.e. the volumes a, b, c, d of the four solutions must be such that the products of the volumes of such pairs of solutions as contain no common ion must be equal to each other.

Now, the volumes a, b, c, and d are proportional to the active or dissociated portions of the four substances present. Calling the total quantities of the latter, p, p_1 , q, and q_1 , the values a, b, c, and d are proportional to m_1p , m_2p_1 , m_3q , and m_4q , and hence we get from ad = bc the equilibrium formula

 $m_1 p \cdot m_1 q = m_2 p_1 \cdot m_4 q_1$ This equation represents not only Guldberg and Waage's formula for chemical equilibrium, but it also contains the extension, made by Ostwald, that every co-efficient of the Guldberg and Waage equation, $k_1 pq = k_2 p_1 q_1$, is made up of two factors each of which depends only on the nature of the ions.

But the equation goes still further than this extension. In the theory of affinity the co-efficients m_1 , m_2 , &c. have been considered as constant to a first approximation. But from what has been said it follows that they are not constant; they depend on the state of dissociation of the four substances, and are rather complicated functions of the dissociation-constants, of the quantities, and of the total volume. The

various deviations from the simple laws which were brought to light by experiment find here their appropriate explanation. Thus it follows that in the case of slightly dissociated acids the degree of dissociation must be considerably lowered by the presence of the normal salts of these acids, which salts are themselves considerably dissociated. The corresponding fact, that such mixtures act much less strongly than the quantity of free acid present in them would do, was known long before theory explained it.

Arrhenius has subjected his investigations

on the saponification of ethyl acetate by means of ammonia in the presence of different salts (Z. P. C. 1, 110) to an examination from the standpoint of these newly-established ideas (ib. 2, 289), and has everywhere found them to be in complete agreement with experiment. has also re-examined a great number of older investigations on conditions of chemical equilibrium in the light of his theory (ib. 5, 1), and here also the discrepancies found when considered from the point of view of the older theory have for the most part disappeared (v. also Z. P. C. 9, 487).

Electromotive forces. chemical theory? Contact theory or

The laws of electromotive force, or of differences in potential produced on the contact of different substances, are not nearly so well known as those of electrical conductivity. It is true the knowledge of the phenomenon itself is as old as that of galvanic electricity, since Volta had already carried out the first approximate measurement of it, but up to the present day no agreement has been arrived at between the different investigators concerning even the chief phenomena.

Volta screwed a disc of zinc coated with isolating varnish on to his straw-electrometer; on the top of this he placed a similar disc of copper, connected the two discs metallically, and after the connection had been broken he took off the copper disc by means of an insulating handle. The electrometer then proved to be charged; examination showed that the straws were charged

with positive electricity.

This phenomenon indicates that the substances used-that is zinc, copper, and the isolating coating—charge each other electrically on contact. Volta assumed that, of the three contacts, only the one between copper and zinc was electromotively active; the contact between the metals and the isolating layer of resin was regarded by him as indifferent. Though neither then, nor up to the present day, has a proof of this assumption been brought forward, it has yet met with fairly universal acceptance, and it often passes as a fact not to be doubted.

On the further assumption, which certainly is incorrect, that liquids also-such as water or salt solutions—show no differences of potential with the metals, Volta constructed his electric pile by superposing alternately two metals and a moist card disc. By this he expected an intensification of the electrical effects, since the electricity of the metal lying underneath was to communicate itself through the moist discs to the one lying above, without any change, and so

 1 A complete discussion of this question by O. Lodge is to be found in P. M. [5] 19.

the effects of the individual pairs of metals were to be added together. Volta did, in fact, obtain the expected intensification - a proof that great discoveries may be made on the basis of quite

false hypotheses.

Fabbroni first refused to accept the assumption that the tension of the voltaic pile has its seat only at the point of contact of the metals, and numerous other investigators have taken the same line. Faraday especially, whose work in electrochemistry ranks with the most important done in this subject, was an opponent of Volta's views. He attributed the production of the electric current solely to the chemical processes present, and transferred the cause of the production of electricity to the places at which these processes occur—i.e. the point of contact between the metals and the electrolytes.

The contention thus aroused between the contact theory and the chemical theory of the galvanic cell has lasted to the present day, and even now it is not completely decided. But most of the discussions raised have certainly originated in the insufficient lucidity with which the questions have been stated.

In so far as a galvanic current can be produced only by using electrical energy, it can be maintained, to begin with, that in order to cause a galvanic current there must be present a corresponding source of energy. If other kinds of energy are excluded, as is the case in an ordinary galvanic cell, it can further be said that chemical energy changes into electrical energy, and that, therefore, a theory of the electric currents produced by the voltaic cell must in any case be a chemical one.

But to an electric current there appertain two factors: a quantity of electricity, and a difference of potential. The first is given by the law of Faraday. Since in electrolytes the motion of electricity takes place only simultaneously with the motion of the ions, equivalent quantities of the different ions carrying equal quantities of electricity, a current in a galvanic cell can again not be produced without a corresponding chemical process. From this point of view, also, the theory must be a chemical one. As regards the other factor of the current energy-i.e. the difference of potential—an unequivocal decision. as in the preceding cases, can certainly not be pronounced easily. The experiments on this subject, taken alone, have proved that all substances, when separated from each other after close contact, show themselves to be electrically charged. The smallest differences suffice for this. It has been long known, for instance, that black and white silk stockings which have been pulled over each other are electrically charged after separation. Even in such homogeneous materials as crystals of mica or calcite, the two parts obtained by cleavage of the layers are found to be in different electrical conditions at the original point of contact.

It is imperative, therefore, to accede to the proposition of the 'contact theory,' in so far as to admit that a process occurs at the surface of contact when two surfaces touch, in consequence of which the surfaces are at different potentials after separation. We do not know what the nature of this process is. Helmholtz supposes the different substances to possess a different

attraction for electricity, in consequence of which there is brought about a separation of the electricity at the planes of contact; then when the bodies are separated they appear electrically charged. A difficulty with respect to this theory is found in the electrical phenomena attending the cleavage of crystals, where a difference in attraction seems excluded.

In order that the equilibrium of electricity may be changed when two bodies charge themselves oppositely on contact, the expenditure of a certain quantity of energy is required. The source of this energy has to be sought in the change which the surfaces of the bodies undergo owing to their mutual contact.

In order that this difference of potential may cause an electric current, two things are needed. Firstly, the bodies touching must be conductors, as otherwise the electricity cannot move; and secondly, a cause must be present which will maintain unchanged the condition of the surfaces in contact, in consequence of which condition the arrangement of the electricity is changed; otherwise a charging motion of the electricity could take place once only, but not a The current can be procontinuous current. duced by mechanical action, and then the phenomena of frictional electricity appear, or the renewal of the surfaces occurs by chemical action, and we deal with galvanic cells.

When considered from this point of view also, the chemical processes appear as the causes of galvanic currents, and a final theory of these currents will doubtless be much more a 'chemical' than a 'contact' theory.

Galvanic cells. The simplest conditions for examining the relations between chemical and electrical energy are found in the so-called constant galvanic cells, of which the Daniell cell is a type. This cell consists of a copper plate in a solution of copper sulphate, and a zinc plate in a solution of zinc sulphate; both solutions are in contact with each other. Such agalvanic cell has an electromotive force of about 1.09 volts, which does not change appreciably when the metals are connected by a conductor, and the galvanic current is thus allowed to pass for a longer or a shorter time.

The chemical process in such an element is also of the simplest kind; zinc dissolves to zinc sulphate, while copper is separated from the copper sulphate. Hence the same process takes place as when a piece of zinc is brought into a copper sulphate solution.

The chemical energy which is available in this process can easily be measured by experiment; it proves to be 50,100 cal. for the reaction

 $Zn + CuSO_Aq = Cu + ZnSO_Aq$.

Now, since (p. 180) the unit of electrical energy, volt × coulomb, is equal to 0.2356 cal., this quantity of heat corresponds to 212,600 electrical units.

Now, of the two factors making up the electrical energy of the voltaic cell, one is given by Faraday's law. One gram-equivalent of a metal carries (p. 180) 96,540 coulombs. Since copper and zinc are divalent, the above equation refers to 2 gram-equivalents; when, therefore, 65 g. of sinc dissolve in the Daniell cell, 193,080 coul. are transferred. Dividing this value into the

number 212,600 (volt × coulomb), we get 1·10 volts for the electromotive force of the Daniell element, a number which agrees well with the observed value.

This calculation was first carried out by W. Thomson with the help of numbers due to Joule. From the agreement between observation and calculation, the inference was drawn that the assumption made in this case—namely, that the chemical energy transforms itself completely into electrical energy—was universally correct. From this was deduced the possibility of calculating, on the basis of thermochemical data, the electromotive force of all constant elements, and further, the electromotive force of all elements in which the chemical process is known. It was only necessary to multiply the heat of the reaction referred to one equivalent (not molecular

weight) by $\frac{0.2356}{96540} = 0.00000244$. It would also

follow that the electromotive force of all cells would be proportional to the heat of the chemical reaction.

This theory could not, however, be maintained. On the one hand, theoretical doubts were raised, and that specially by Braun (W. 5, 188), against the premisses that the chemical energy of the galvanic cell was always completely transformed into electrical energy; and, on the other hand, experience proved that as a rule the assumed proportionality between heat of reaction and electromotive force did not exist. Raoult (A. Ch. [4] 4, 392) discovered a number of cells, some of which produced less, and some produced more, energy than corresponded to the heat of reaction of their chemical changes; and at a later time it was provedchiefly by F. Braun and Alder Wright (P. M. [5] 19, 1), by means of direct measurement-that only in rare cases do the chemical and electrical energy exactly correspond to each other.

The simple theory must, therefore, be abandoned, and without assuming that the chemical energy is wholly transformed into electrical energy, we must seek to discover a relation between the two. Such a formula has been found by W. Gibbs and H. von Helmholtz $(B.\ B.\ 1882)$. The following statement is a simple way of deducing this equation. Let there be given a galvanic cell whose heat of reaction calculated for one equivalent is W. Let its electromotive force at the absolute temperature τ be ϵ . On raising the temperature by $d\tau$ the electromotive force is changed by $d\epsilon$.

Now let us carry out the following cycle. The cell is heated from τ to $\tau + d\tau$. Then let the quantity of electricity q, by means of which one equivalent of the reacting substance undergoes chemical decomposition, pass through the cell, the electromotive force remaining $\epsilon + d\epsilon$, and the temperature being kept constant. Now let the temperature be brought to τ and, the electromotive force being ϵ , and the temperature being constant, let the same quantity of electricity q be passed through the element in the opposite direction. Thereby the element returns exactly to its initial condition.

According to the mechanical theory of heat, the portion of heat transformable into work, in a reversible cycle taking place in the temperatureOE

interval $d\tau$, is $\frac{d\tau}{\tau}$ of the total quantity of heat produced. Now, the work is given by $qd\epsilon$, and the total heat which has been communicated to and taken from the cell is $\epsilon q - W$. If an electric current alone were produced, the quantity of heat would be ϵq . But since a chemical process also takes place, which yields a portion of the heat—namely, the heat of reaction, W—it is only necessary to communicate the difference $\epsilon q - W$. We have then the equation—

 $\frac{d\tau}{\tau} = \frac{qd\epsilon}{\epsilon q - W}$

 $\mathbf{e} = \frac{\mathbf{W}}{\mathbf{q}} + \frac{1}{\tau} \cdot \frac{\mathbf{d}\mathbf{e}}{d\tau}$

The former equation was $\epsilon = \frac{W}{q}$; we see, therefore, that a correcting term must be added, and that this term is dependent on $\frac{d\epsilon}{d\tau}$, that is, on the change of electromotive force with change of temperature. In the case that $\frac{d\epsilon}{d\tau} = 0$, the new equation is the same as the old one; in the Daniell cell the temperature co-efficient is, in fact, nearly equal to zero, and thus it is explained that in this case the chemical energy is equal to the electrical energy.

The equation has been examined in many ways. At first it gave only partially correct results, and in some cases observation and calculation differed from each other. The differences occurred, however, only in cells which contained mercury compounds, and the cause was the use of a wrong number for the heats of formation of these compounds. When this error was corrected by Nernst (Z. P. C. 2, 23) the discrepancies disappeared; and a very careful investigation by H. Jahn (W. 34, 755), in which the thermal and electrical quantities were measured simultaneously, has fully confirmed the correctness of the Gibbs-Helmholtz formula.

The formula has been deduced primarily for constant galvanic cells, and when deducing it the conditions assumed have been expressed; the chemical process resulting from the passage of the quantity of electricity q (which, however, can be as small as we will) must be reversible, by allowing the same quantity of electricity q to pass through the element in the opposite direction. In order that a galvanic cell should fulfil this condition, it is generally requisite that each metal should be surrounded by its corresponding salt. Therefore, when the salt is soluble, the metal must stand in a solution of the salt. the case of insoluble salts it generally suffices to cover the metallic surface with the solid salt; in this case, either the salt of another metal with the same acid (which salt is not precipitated by the first metal), or the free acid itself, serves as the electrolyte.

The action in a Daniell cell is as follows. Individual atoms of the metal pass from the zinc as positively charged ions into the solution of the sulphate. Since no free electricity can exist within the solution, the copper ions of the copper sulphate must go to the copper plate, where they give up their positive electricity, passing into ordinary metallic copper. Owing to this

action, a motion of the positive electricity from the zinc to the copper takes place in the solution; on connecting both metals by a wire the current passes continually through it, corresponding quantities of zinc and copper being dissolved, and precipitated, respectively. If the cell is not closed, the zinc becomes charged negatively, owing to the exit of positive ions, and the copper positively, owing to their entrance; and this goes on until the attraction of the zinc for the positive ions, and their repulsion by the copper, is so great that neither can zinc pass into solution nor can copper separate out. The capacity of the ions being enormous, this is brought about by even extremely small numbers of them.

Nernst, who has studied these changes in detail (Z. P. C. 4, 129), assigns to the metals which are in contact with electrolytes definite solution pressures, just as volatile liquids or solids have definite vapour pressures. This pressure is the cause of zinc tending to pass into solution. The solution pressure of copper is very much smaller than that of zinc. Hence when the Daniell cell is newly set up, both metals will at first send positive ions into the solution. But, since no free electricity can exist in the interior of the solution, that metal which has the greater pressure will maintain the preponderance; its ions will pass into solution, and, in order that no free electricity may accumulate in the liquid, an equal number of copper ions must leave the solution, and be precipitated in the metallic state.

Similar arguments hold for all reversible elements. By their help, and with the help of the theories of electrolytic solutions expounded above, a great number of special cases referring to galvanic cells can be deduced theoretically. The results thus obtained have, on the whole, been confirmed by experiment; but details cannot be entered into here.

Besides the constant galvanic cells of the type just described, which are reversible, there still exists a great number of cells which, though practically constant, are not reversible. The constancy of these cells rests on the fact that the nature of the surfaces in contact is not altered by the chemical processes. When metal dissolves at the surface of the zinc, and when metal deposits at the surface of the copper, the nature of the bounding surfaces is not thereby influenced; this cell is also reversible, because on sending a current through in the opposite direction zinc is precipitated and copper dissolved, but the surface again undergoes no alteration.

Cells of the type of that of Grove, which consists of zinc in sulphuric acid and platinum in nitric acid, are also constant. On working this cell, zinc dissolves, expelling the hydrogen ions from the sulphuric acid; these hydrogen ions go to the nitric acid, whose own hydrogen ions go to the platinum; at the moment when they give up their positive electricity there, and pass into ordinary hydrogen (which, however, is in the nascent state), they are oxidised by the nitric acid to water, and the platinum surface remains unaltered. But such elements are not reversible. When an opposite current is passed through the cell, since the platinum ions cannot pass into solution, the ions NO₂ of the nitric

acid separate out at the platinum, while hydrogen appears at the zinc; both these actions change the condition of the bounding surfaces, and thereby simultaneously change the electromotive force also.

Finally, there still remains the large group of the non-constant cells. These are distinguished by the fact that even when the current passes through them in the direct way, the ions separated out produce an alteration in the bounding surfaces, or in one of them at least. To these belongs Volta's cell, which consists of zinc and copper in salt water, and all the other similar combinations. A type still in use is Smee's cell, which consists of zinc and platinum in dilute sulphuric acid. On closing the circuit, zinc ions enter into the solution, expelling the hydrogen, which gives up its electricity at the platinum, whereby it first combines with or dissolves in the metal, and when that is saturated escapes in the gaseous form. In consequence of this, the surface of contact between the acid and the platinum is altered, and the electromotive force also changes, until saturation with hydrogen has taken place; after this point has been reached the electromotive force of Smee's cell remains practically constant. If the cell is left with the circuit open, the hydrogen gradually disappears, partly through diffusion into the air, partly through combination with the free oxygen of the air, induced by the catalytic action of the platinum. Thus, the cell again gradually resumes its original electromotive force. Such cells are, therefore, quite workable in cases where only small quantities of electricity are required at long intervals.

The seat of the electromotive forces. Volta found that no galvanic pile, giving a current, could be produced from metals only. When considered from the point of view of the principles of energy, such behaviour follows necessarily, because, no continuous source of energy being present in a circuit of metals alone, no current can be produced.

The cause of this may be of a two-fold nature. Either there exist no differences of potential between metals, or these follow a definite law, which involves that in a closed circuit these differences just neutralise each other. With Volta, we will assume the latter alternative, of which the former is only a special case. If, then, we have several metals A, B, C. . . and if we call the differences in potential between A and B, A and C, B and C, &c., in succession (a-b), (a-c), (b-c) . . . the law is that we must always have $(a-b)+(b-c)+(c-d)+\ldots+(y-z)=(a-z)$. The difference of potential of a series of metals depends only on the nature of the first and the last metal, and not on the nature of the intervening metals.

This result necessarily follows if the values $a, b, c \dots$ in the quantities (a-b) &c. are considered as the potentials of the individual metals referred to a zero value which must be fixed arbitrarily. For the zero value of potential we generally use the potential of the earth, or, more correctly speaking, that of a large metallic mass which is in connection with the earth, as are, for instance, iron water-pipes. It can, therefore, be assumed that every metal con-

nected with the 'earth' assumes a definite value of potential which depends only on the nature of the metal. These values of potential are the same for every metal, whether the different metals are in contact with each other or not.

In order to measure these differences of potential, the experiment of Volta mentioned before (p. 212) has been improved upon, in that instead of a dividing layer of resin one of air is used; and, dating from the time that R. Kohlrausch (P. 75, 88) thus carried out the first experiments, a great deal of similar work has been done. But in this case also we are dealing, not with one, but with three separating surfaces—namely, copper-zinc, zinc-air, air-copper, and only when we make the assumption that the differences of potential at the last two surfaces are zero, are we enabled to judge from the measurements of the total difference of potential as to the difference between the metals.

The legitimacy of this assumption has formed the subject of much discussion. Considering the universality of differences in potential between heterogeneous substances, the absence of a difference between metal and air would be very strange, and we are quite entitled to demand a proof of it from the champions of this view. But, on the contrary, it seems as if the following fact proved the existence of such differences of potential. All the investigators who have undertaken such measurements complain of the extraordinary inconstancy in the differences of potential observed. On placing two freshly-cleaned plates opposite to each other and measuring their difference, one is sure to find different values after a little time. If the surfaces are again cleaned, the original values approximately recur. Therefore, it is not the point of contact of the two metals (besides, this can be made unalterable by soldering), but their surfaces with respect to air, which are the seat of the changes. Let us now assume that at a definite moment the difference of potential of both plates towards air is equal to zero; since another value is observed after some time, the difference of potential of the metals towards air cannot now be zero, i.e. between metals and air there exist differences of potential. How great these are cannot as yet be determined. There is nothing against the assumption that these make up the greatest part of the apparent differences of potential between the metals (measured in air). It is true an attempt has been made to eliminate the influence of the air by making the experiment in vacuo. But since we have learned by observation of the electrical phenomena in vacuum tubes, how difficulthow impossible, even—it is to withdraw from a substance the last remnants of air condensed on it and in it, these experiments cannot be looked upon as affording any conclusive proof.

Other methods for solving the difficulties must be sought. Unfortunately, there does not exist a single process quite free from objection. The following consideration still appears to be the best:—

When a given quantity of electricity ϵ is brought from a potential A to a higher potential A + a, a quantity of work aq is required. There-

fore, when we lead a current through two metals which touch in one place, and which there exhibit a difference of potential a, an absorption or a production of energy will occur at the place of contact, according to the direction of the current; and, if there is no other source of energy, this energy-change will be produced from, or will be added to, the heat which is present, as the case may be. In the case of metals such a phenomenon is actually known; it is the effect discovered by Peltier (A. Ch. 56, 371 [1834]). If the strength of the current, and the total heat produced, are measured, the corresponding difference of potential can be calculated. Such experiments have been made by Le Roux, and again quite recently, with great accuracy, by H. Jahn (W. 34, 755), and the results have shown the values to be very small; they are mostly only a few thousandths of a volt.

It is true we must not pass over in silence the following objection made against the validity of this argument. Since we know nothing definite concerning the nature of the electric current, the possibility is not excluded that it may be connected with movements of energy of other kinds. It has been especially assumed as probable by several authors that, simultaneously with the electric current, a heat-current flows through the conductor. If so, the quantity of heat flowing with the unit quantity of electricity might be different in different conductors; and at the point of contact of the metals, in addition to the thermal exchange resulting from the difference of potential, there might occur a further exchange of heat resulting from the difference in the 'thermal capacity of electricity.' Since these two quantities need not have a necessary connection, it would not be permissible to draw any conclusion from the Peltier effect-that is, from the sum of two independent effects—as to the magnitude of the difference of potential.

Without attempting to decide this difficult question here, I should like to draw attention to a circumstance which makes it probable that the second effect, if present at all (which so far has not been proved), must probably be of the same order of magnitude as the difference of potential corresponding to the Peltier effect. According to the second law of the mechanical theory of heat, the carrying of a definite quantity of heat Q from a lower to a higher temperature involves

work which is given by $Q^{\frac{\tau_1-\tau_2}{\tau_1}}$. If, therefore,

we pass an electric current through a wire whose temperature varies, it follows that, if heat is carried with the current, in places of rising temperature a corresponding quantity of work must be expended, i.e. an opposing electromotive force must there assert itself. Indications of such a force appear to manifest themselves in the so-called Thomson effect (supposing that this effect does not find its explanation in differently heated portions of one metal having differences of potential to each other), but in any case the quantities of work, or the differences of potential, indicated by this phenomenon are extremely small (of the same or of a smaller order of magnitude as those corresponding to the Peltier effects), so that the assumption of an appreciable carrying of heat by means of elec-

tricity (for which, at any rate, we lack the power of forming a conception) seems excluded.

From this it is evident that though nothing quite certain is known concerning the differences of potential between different metals, yet preponderating reasons favour the assumption that such a thing does exist, but that its magnitude does not exceed a few thousandths of a volt, and particularly that it does not attain the order of magnitude of 1 volt. But, varying slightly with the nature of the metals, such a value has been observed by means of the air-condenser method.

As regards differences of potential between various electrolytes, such differences certainly do exist. This can be inferred from the fact that there exist 'liquid chains,' i.e. combinations of different liquids, which produce an electric current. If, for instance, we put into a series of beakers water, concentrated hydrochloric acid, concentrated soda solution, and water, and if we connect these by small syphons or by moist threads, then the two quantities of water at the ends exhibit a difference of potential which may attain to 0.5 volt.

In such liquid chains we have always to deal with three distinct differences of potential at least; designating the three liquids named above by A, B, and C, there exist in the chain the differences of potential (a-b), (b-c), and (c-a), whose sum is observed. There is no combination of the liquids by which we could succeed in obtaining as many independent observations as there are differences of potential, so that it is impossible to determine by measurements on liquid chains only the individual differences of potential.

This determination has, however, been achieved in another way. Since in electrolytes the electricity moves only with the ions, differences in potential between two liquids in contact can only be produced if the different positive and negative ions distribute themselves unequally. A cause for such an inequality lies in the different velocities of migration of the ions (p. 92), and, in an extensive investigation, W. Nernst has shown (Z. P. C. 4, 129) how thereby the actually observed differences of potential can be theoretically calculated in a great number of cases. The fundamental idea of these calculations is the following. When any solution is brought into contact with pure water, forces make themselves apparent at the surface of separation, which forces can be calculated by van't Hoff's theory (pp. 184-5); for 'normal' solutions (which contain one gram-molecular weight per litre) they are equal to the pressure of about 22 atmospheres, and they are proportional to the quantity of substance in the solution. Owing to the osmotic pressure, the molecules of the substance are forced into the water. The velocity of their motion is determined by the resistance which they encounter, and this depends on their form as well as on the nature of the solvent.

This statement contains a physical theory of hydro-diffusion (Nernst, Z. P. C. 2, 613), and from it the law of Fick follows directly, that the velocity of diffusion at any point is proportional to the change in concentration at that point. From this we see at once how a separation must occur in the diffusion of dissolved substances,

because those molecules which are endowed with greater mobility or suffer less friction outstrip the slower ones. The same laws hold for each kind of molecules, only the constants are

These arguments can only be applied immediately to electrolytes in solution if both ions have equal rates of motion. This condition is very nearly fulfilled in the case of potassium chloride, and this substance conforms, therefore, to the simple law of diffusion.

It is now possible, in the case of electrolytes, to determine the resistances which the ions encounter, as well as to measure the impelling If the ions are moved past each other by electrical forces of separation, they encounter exactly the same resistance, and since in the electrical conductivity we have a measure of the velocity of the corresponding motion of the ions, it is possible, as first shown by F. Kohlrausch (W. 6, 160), to calculate the magnitude of the resistances from a knowledge of the impelling force and of the velocity attained. On introducing this resistance (which, owing to the smallness of the molecules, is enormously great) into the equation for the velocity of diffusion, indicated above, Nernst has found (l.c.) that the velocities of diffusion agree very well with the results of experiment.

In the case of electrolytes whose ions do not migrate with equal velocities the phenomena become much more complicated. If, for instance, we imagine hydrochloric acid, in which the hydrogen ion travels about six times more quickly than the chlorine ion, to be brought into contact with water, the quicker hydrogen atoms will, to start with, pass into water, and the chlorine atoms will remain behind. But since the hydrogen atoms are charged with positive electricity, the water will become positively electrified, while the acid, owing to the preponderance of chlorine atoms, becomes charged negatively. But the further exit of hydrogen atoms is thus rendered more difficult, owing to the electro-static effect, while from a similar cause the chlorine atoms are subject in addition to the osmotic pressure, and also to an electric impelling force, and are thus driven into the water. The consequence is that both ions influence each other's velocity until they again migrate at equal rates.

At the same time, we see that every electrolytic solution whose ions travel with unequal velocities on contact with water-or, speaking more generally, on contact with a solution of different concentration-at once acts as the cause of difference of potential, and also that the more dilute solution will always assume the sign of the ion which has the greater migration velocity. The exact expression which has been deduced by Nernst (l.c. p. 138) is

$$e_1 - e_2 = 0.0000860 \quad \tau \quad \frac{u - v}{u + v} \log \frac{p_1}{p_2}$$

Where $\epsilon_1 - \epsilon_2$ is the difference of potential (in volts) of two solutions in which the osmotic pressures of the (positive as well as negative) ions are p_1 and p_2 ; u is the velocity of the positive, v that of the negative, ion; τ is the absolute temperature, while log. stands for the natural logarithm.

We see that for u = v the electromotive force becomes zero, and it becomes the greater the greater the difference between the migrationvelocities of the two ions. Further, we see that as regards the electromotive force, only the ratio

 P_1 of the two pressures, and not their absolute

values, enters into consideration. By increasing the concentration n times in both solutions, the difference of potential does not change. If one of the solutions is pure water, we get $p_2 = 0$, and the electromotive force becomes infinitely great. Since 'pure water' in this sense does not exist, such values cannot come under our observation. But yet we see that liquid chains with 'water' can give very different values according to the degree of purity of this liquid.

The preceding formula is the fundamental expression for all kinds of liquid chains, since in the combination of any solutions the corresponding differences of potential are superposed. It would take us too far afield to enter into the different cases; in the memoir of Nernst, a number of cases are discussed theoretically as well as experimentally. But on the whole it may be concluded that the differences in potential apparent on contact of approximately equally concentrated solutions of the most varied kind, vary within fairly small limits, which probably do not surpass 0.01 volt.

Now, since we cannot suppose that differences of potential of the order of magnitude of those observed in galvanic cells exist either between metals alone, or electrolytes alone, we are obliged to regard the surfaces of contact between the metals and the electrolytes as the real seat of the electromotive forces. This result quite corresponds with what was to be expected from the relations between the chemical and electrical energies. This is so because these surfaces of contact are the only seat of the processes by which energy becomes available in the chains, i.e. they are the result of the chemical changes. Since, on the one hand, the cause of the galvanic current of the chains has to be sought in the transformation of chemical into electrical energy, and since, on the other hand, a galvanic current can be produced only when a difference of potential exists, it does not seem very logical to wish to separate in space, and in their nature, the more approximate and the less approximate causes of the same phenomenon, as is done in the assumption that the difference of potential is to be sought at the surface of contact of the metals.

The question now arises again, whether some means does not exist for directly measuring the differences of potential between metals and electrolytes. The answer is that up to the present only one such means is known. A second independent way for arriving at these numbers does not as yet exist, and hence it has hitherto been impossible to put to the proof the results obtained by that method.

The method in question is based on the phenomena exhibited by mercury surfaces when in contact with electrolytes. Lippmann (P. 149, 561; A. Ch. [5] 5,494) has shown that there is a close connection between the condition of the surface of mercury touching an electrolyte and

the electrical state of the system. If, for instance, mercury is covered with dilute sulphuric acid, and the surface of contact between the two is then increased, a galvanic current is produced, because the mercury becomes less positive on expansion. This can be observed most easily by allowing mercury to drop out of a funnel with a small orifice immersed in dilute sulphuric acid. On connecting the mercury in the funnel and that collected below the sulphuric acid by means of a galvanometer, a current passes from the lower mercury to the higher; in the sulphuric acid the reverse occurs.

This phenomenon is reciprocal. On leading a current through the surface of contact between mercury and sulphuric acid the surface tends to diminish or to increase, according to the direction of the current—i.e. the capillary tension at this point changes. As is always the case, both phenomena are in such a relation that they oppose each other—i.e. when the surface is made larger the current has the direction such that if it alone were active it would tend to make the surface smaller, and vice versa.

Helmholtz (W. 16, 35 [1882]) has given a theory of these phenomena which is based on the following considerations. When a metal and an electrolyte are in contact, they generally possess a varying difference of potential. Owing to this circumstance, the opposite electricities must accumulate on both sides of the bounding surfaces, since they are prevented from combining by the cause which produces the difference of potential (and which, without making a definite assumption, can be designated as a different attraction of the substances for the electricity). The result is the formation of an 'electric double layer,' which will exhibit a similar arrangement to the charge of a Franklin plate.

But such a double layer will have the property of tending to increase its surface as much as possible, since the similarly electrified particles present at each side repulse each other, and therefore tend to stretch the surface. Hence the development of such a double layer will act in opposition to the surface tension, which seeks to decrease the surface, and the actually observed surface tension is the difference between the real tension and the electric force.

When a surface so constituted is made larger, the electric layers are thereby extended, and their difference of potential decreases. But since, as is known from experience, a constant difference of potential is always formed at the boundary, which difference of potential does not depend on the size of the surface of contact, electricity will flow from the mercury as well as from the electrolyte to replace the loss, and in this way the current is produced.

If, on the other hand, the difference of potential of the electric double layer is changed, the surface tension must change also. It must get smaller when the difference of potential is made larger, and larger when it is made smaller.

By leading electric currents of varying electromotive force through a system consisting of two masses of mercury in sulphuric acid—one of which had a very small, the other a very large, surface of contact with the acid—Lippmann was able to follow the change of surface tension at the small bounding surface which accom-

panied the change in the difference of potential at that surface. Because, since the change in the difference of potential between a metal and an electrolyte is proportional to the density of the current (current strength divided by the cross-section), the resulting difference of potential practically restricted itself to the small bounding surface, which was surpassed in extension some thousand times by the large one. It was found that when the mercury of the small surface was connected with the positive pole of the external cell, the surface tension decreased. On the other hand, when negative electromotive forces acted, the surface tension increased until the external force was 0.9 to 1.0 volt; beyond that it again decreased.

In accordance with Helmholtz's theory, we must infer from this that the difference of potential between mercury and sulphuric acid is of such a kind that the former becomes charged positively-i.e. that the positive electricity of the double layer is on the side of the mercury, and the negative electricity is on the side of the electrolyte. If the difference of potential is increased by addition of positive elec tricity to the mercury, the electric repulsion increases, and therefore the surface tension decreases. If, however, negative electricity is added, the charge of the double layer decreases and the surface tension increases. On making the successive additions of negative electricity larger and larger, the difference of potential of the double layer eventually becomes zero; the surface tension has reached its greatest value, and, beyond this, a new double layer of opposite sign is produced. When this state is reached, the surface tension must again decrease.

According to the experiments of Lippmann, the maximum of surface tension is reached when the external electromotive force is 0.9 to 1.0 volt. Then no more double layer is present at the small surface of the mercury, and the metal and electrolyte have the same potential. Since this condition is separated from the ordinary one by about 0.9 volt, it follows that the usual difference of potential between dilute sulphuric acid and mercury is 0.9 volt.

This result is confirmed by the following experiment of Pellat (C. R. 104, 1099). When the surface of contact between mercury and sulphuric acid is increased, a motion of electricity is produced, owing, as has been mentioned above, to the stretching of the double layer. But this must disappear when, by the use of a difference of potential of about 10 volt, the double layer has been made to disappear; and, in fact, under these conditions, Pellat, on changing the surface, could observe no current.

Another confirmation can be obtained in the following manner. When mercury is dropped very quickly out of a fine point into an electrolyte, it must, after a short time, as Helmholtz has shown, assume the potential of the latter (W. 16, 35), because if, for instance, the mercury were at the beginning positive, every falling drop would form a double layer at its surface, which would take positive electricity out of the mercury, and which would make its positive potential smaller and smaller till it became equal to that of the liquid.

But it is to be remembered that the mercury

in contact with sulphuric acid spontaneously becomes positively charged; everything will depend, therefore, on allowing the formation of drops to take place as quickly as possible, in order that the discharge may preponderate over the charge. Experiments by W. Ostwald (Z. P. C. 1, 583 [1887]) have shown that the discharge can be achieved to a great extent, but not completely; there still remain over, even with the best dropping electrodes which have yet been made, differences of some hundredths of a volt. But within these limits the experiment confirms the conclusion which Helmholtz had drawnthe difference of potential between the dropping electrode and the mercury at rest in sulphuric acid amounts in 'normal' sulphuric acid solution to 0.81 volt, while the maximum of surface tension (that is, the true difference of potential) lies at 0.86 volt.

Finally, another inference, which must be drawn from the theory of Helmholtz, has been confirmed by the same author. If the maximum of the surface tension of mercury, as obtainable by means of suitable electromotive forces, actually is the real surface tension of mercury in contact with aqueous liquids, free from auxiliary effects, then it must have the same value whatever be the nature of the electrolyte, while experience shows that the natural surface tension changes considerably with the nature of the electrolyte. This also was confirmed; while the natural surface tension in different solutions varied between 485 and 564, the maximum values were 640 to 644, and were, therefore, practically constant.

All these facts speak for the correctness of Helmholtz's theory of the electric double layers, and leave us, therefore, with the final conclusion that the electromotive force required for producing the maximum of surface tension of mercury in contact with an electrolyte is equal to the ordinary-difference of potential at this bounding surface (but with inverted sign). But it must be again emphasised that this is the only group of phenomena from which the values considered can be deduced. An examination of the results by means of another entirely independent method has not yet been feasible.

If, now, the difference of potential between mercury and sulphuric acid is known, the difference of potential between all other metals and sulphuric acid can be measured by experiments which are easily carried out. For this purpose the electromotive force of the combination mercury, sulphuric acid, and metal alone need be measured; this is the sum of the differences of potential of mercury-sulphuric acid and sulphuric acid-metal, and since the first of these is known the other follows by difference. But here it is supposed that no appreciable difference of potential exists between the metals. Again, if the difference of potential between mercury and any other electrolyte is determined, either by the maximum surface tension or by means of the mercury-dropping electrode, the difference of potential of these electrolytes against any other metals can also be determined. In this manner, therefore, the whole subject of the differences of potential between metals and electrolytes has been made amenable to measurement.

But few trustworthy measurements of this

kind have as yet been made; in the following table some numbers are given:—

	HCI	HBr	HI	H ₂ SO ₄
Zn	-0.54 -0.24 +0.02 +0.03 +0.35 +0.41 +0.51 +0.57 +0.57	-0.46 -0.18 +0.12 +0.10 +0.35 +0.47 +0.60 +0.51 +0.50	-0·30 -0·08 +0·28 +0·26 +0·36 +0·60 +0·54 +0·45	-0.62 -0.22 -0.02 -0.04 +0.46 +0.48 +0.73 +0.86

The values refer to 'normal' solutions of the electrolytes mentioned, and indicate the potential of the metal when that of the electrolyte is put equal to zero.

We see that those metals which dissolve in the acids become negative; the others become positive. The values are also in very nearly the same order as the corresponding heats of reaction, but, as a calculation shows, they are not proportional. The nature of the electrolyte has generally a smaller influence than that of the metal, but there are cases where the reverse is true.

It should be mentioned that nearly the same numbers are obtained when salts of the acids, with metals which are not precipitated by the metal under examination, are used instead of the acids themselves. The difference of potential between the metal and the electrolyte depends, therefore, chiefly on the negative ion of the latter. From the numbers in our possession it is seen, for instance, that in the Daniell cell, whose difference of potential is 1.08 to 1.10 volt, the greater part of this (0.62 volt) is present at the bounding surface between zinc and zinc sulphate, and the smaller part at the surface (0.46 volt) between copper and copper sulphate.

No further conclusions of more general interest have as yet been drawn from these num-

Galvanic polarisation. When two metal plates are introduced into an electrolytic liquid and a galvanic current is made to pass through the electrolyte by means of the plates, these plates generally become the seat of a new electromotive force which acts in opposition to the current already active. This phenomenon is called galvanic polarisation, and it may be perceived by connecting the plates with a galvanometer immediately after breaking the primary current.

The cause of this phenomenon is to be found in the change which is produced by the current at the boundary surface between the metal and the electrolyte. There the ions of the electrolyte separate out, and cause changes of the most manifold kind.

The simplest case occurs when the metal plates are surrounded by the solution of a salt which contains the same metal, for such a system scarcely exhibits the phenomenon of polarisation at all; such electrodes are 'unpolarisable.' The reason is that in this case the changes occurring at the electrodes do not change the condition of the bounding surfaces. If, for instance, we have zinc in zinc sulphate.

zinc is separated out at the cathode, and this combines with the metal present without changing it, and at the anode SO₄ separates out, which then dissolves zinc and forms zinc sulphate. The only effect is that the zinc sulphate solution becomes more concentrated at the anode, and accordingly a small concentration-current can be observed, whose electromotive force is, however, very inconsiderable.

Such a system of unpolarisable electrodes differs from all similar combinations in that it allows any, even the smallest, current to pass through. Polarisable electrodes do not behave thus. If, for instance, two platinum plates are placed in dilute sulphuric acid, and a current of small electromotive force—say, 0.1 volt—is allowed to pass, we observe by the galvanometer that at first a motion of electricity takes place, but that this ceases after a very short time. It is a phenomenon such as a condenser exhibits on being charged, with this difference, that the capacity of such electrolytic condensers appears to be very great. On increasing the electromotive force, a corresponding quantity of electricity answering to a greater charge of the condenser always enters, but no permanent current is produced until the electromotive force is somewhat more than 2 volt; then a more copious passage of the current suddenly takes place, and simultaneously gaseous oxygen and hydrogen appear at the electrodes.

We must, therefore, represent the phenomenon to ourselves in the following way. Between the electrolyte and the platinum plates there exists originally some one difference of potential, and in consequence the double layers form there. Now, on allowing the electromotive force to act, on the one side the difference of potential of the double layer is increased, on the other it is diminished. The corresponding process to this is the condensing charge. On increasing the difference of potential, there is a moment at which the condenser no more isolates under the influence of the electro-static effects; a passage of electricity from the metal to the ions (or vice versa) occurs, corresponding to the disruptive discharge of the condenser, and the ions, deprived of their electric charges, separate out. When sulphuric acid is the electrolyte, the ions are H2 and SO4; the latter substance is not stable, and reacts with the water present to form sulphuric acid and oxygen, the latter escaping in the gaseous state.

From this we see that a difference of potential of definite magnitude must exist when the electricity is to leave an ion in order to travel to a metal; until this difference of potential is reached, the system acts at the electrode like an isolator. But why does this not occur with unpolarisable electrodes? The answer to this question follows from the consideration of the processes occurring at these electrodes. Let us again take zinc plates in zinc sulphate, and let us lead positive electricity into one plate. This electricity can at once enter into the solution, by using as its vehicle a corresponding number of zinc atoms, which travel with it as positively charged zinc ions. At the other zinc plate an equal number of zinc ions can also deposit at once, and since there exists no difference of potential between zinc and zinc these can at once deliver up their electricity. Here, therefore, no cause exists for condensing effects. It is not, however, as if no difference of potential existed between the metals and their solutions; such differences, in fact, are present, and hence also corresponding double layers are formed, which at one side are made up of ions. But these do not act as condensers, free communication of their ions between the metal and the electrolyte being possible, and consequently the difference of potential, independent of the current, maintains itself at an unaltered value.

Beside the unpolarisable electrodes and those polarisable at both sides, there still exist combinations in which polarisation occurs only at one side, mostly at the cathode. This takes place, for instance, with copper plates in dilute sulphuric acid. At the cathode, when the electromotive force is sufficient, hydrogen separates out, just as it does at a platinum plate. But at the anode, on the other hand, copper sulphate is formed by the ion SO₄ acting on the copper—i.e. the ion SO₄ need not give up its negative electricity there, since in its stead positive electricity fixed to copper comes out of the plate. Hence, no polarisation occurs here.

From what has been said an explanation follows of the cause of the polarisation current, which occurs after breaking the primary current and connecting both plates in the opposite direction. It is, in brief, the discharge of the electrolytic condenser. Looking at the process more in detail, we see that when the charge in the double layers is formed, the electricity led in has remained in the electrodes, equal quantities of the opposite electricities having collected from the electrolyte on the electrodes—the electricity being, of course, bound to the ions. the primary current being broken, and the electrodes being connected with each other, the electricities present in these neutralise each other, and the ions, which are no longer fixed, form the carriers of the current in the electrolyte. This lasts until the original condition of equilibrium has re-established itself.

In accordance with what has been said, the measurement of galvanic polarisation refers almost exclusively to the maximum value it can assume, or to the value of the difference of potential at which the double layer condenser no more isolates.1 The measurement of this quantity is somewhat difficult; because, when carrying out the simple experiment indicated above-i.e. when increasing the primary current up to decomposition, interrupting it, and afterwards connecting the electrodes with an electrometer—we observe that (in consequence of the secondary effects) the electromotive force of polarisation decreases very rapidly. We are, therefore, not sure whether a considerable portion of the polarisation has not already been lost in the time which elapses before the measurement. But there exists a means of obtaining information concerning the magnitude of the possible error. If the change of connections is carried out more and more quickly, increasing values at first are observed. But these approach

1 It must be observed that complete isolation does not occur; a small amount of percolation of electricity takes place when the maximum value is not reached. But, as Helmholtz has shown, this phenomenon is a consequence of electrolytic convection, and is of a secondary character.

to a maximum, and when the number of current reversals reaches about 100 per second a further rise in the velocity no longer produces an increase, as was found by Raoult (A. Ch. [4] 2, 326), and the value observed under these conditions may be regarded as the correct one.

Other methods are based on making measurements of current strengths and resistances in the primary circuit, and on calculating from this the electromotive force of polarisation. These become uncertain by the fact that, owing to the processes occurring at the electrodes, the resistance of the cell undergoes variations during the passage of the current, which variations cannot be determined. Finally, we can use electrometric methods, by measuring the difference of potential between the electrode to be polarised and an auxiliary electrode through which no current passes, both before closing the circuit and while the current passes; the difference between the two magnitudes is the polarisation required. But difficulties, which have not yet been removed, appear in the practical application of this ingenious method, which was pointed

out by Fuchs (P. 156, 158).

Because of these difficulties only a small number of trustworthy measurements of the electromotive force of the maximum polarisation have as yet been made. Consequently, no certain relations with chemical conditions have been discovered as yet. Though an attempt has been made to calculate the electromotive force of polarisation in a manner similar to that of the galvanic cell, from the thermal effects appertaining to it, the same objections can be raised against it as in the former case (p. 213): the observed electromotive force is generally

different from that calculated.

It is especially remarkable that the maximum forces of polarisation corresponding to one and the same chemical process may be very different according to the nature of the metal. If we form galvanic cells of zinc, dilute sulphuric acid, and different metals—such as copper, silver, gold, platinum-or of charcoal, and if we close the circuit in the cells, the same process occurs in all: zinc sulphate is formed, and, to make up for it, hydrogen is expelled from the sulphuric The latter appears at the metal, and polarises it. It is found that the electromotive forces of these cells, even after they have completely polarised themselves, are distinctly different; according to the nature of the second metal, a greater or less portion of the energy liberated by the reaction $Zn + H_2SO_4 = ZnSO_4 + H_2$ is transformed into electrical energy.

The cause of this can be understood after what has already been explained. The electricity must overcome unequal differences of potential in passing from the H ions to the different metals; hence different portions of the total available energy are used, and only the residue of this energy finds its expression in the electromotive force of the cells. We see, then, that the electromotive force cannot possibly bear the simple relation to the heat of reaction which was formerly supposed to be the case. But it is possible that the equation of Gibbs and Helmholtz (p. 214) is applicable to galvanic

polarisation also; if this is so we shall be able. from a knowledge of the electromotive force and its changes with the temperature, to draw conclusions as to the thermal changes due to the processes which primarily participate in the polarisation. W. O.

VI. FREEZING-POINTS OF SOLUTIONS, METHODS BASED ON; v. MOLECULAR WEIGHTS, vol. iii. p. 417; and ELECTRICAL METHODS, this vol. p. 186.
VII. OPTICAL METHODS.—Section 1:

REFRACTION AND DISPERSION. When a ray of light passes at an angle from a rarer into a denser medium, it is always reflected towards the perpendicular, the sine of the angle of incidence being in a constant ratio to the sine of the angle of refraction, namely the ratio of the velocities of wave propagation in the two media. This constant ratio is called the index of refraction, and is generally represented in England by the Greek letter μ , and on the Continent by $n: \mu = \frac{\sin i}{i}$ The angle of incidence sine r

being in such case greater than the angle of refraction, the index of refraction will be greater than 1. The reverse will be the case when the

ray passes from a denser to a rarer medium.

The refraction of a ray of light is, however, always accompanied by another phenomenondispersion, as the ray is made up of a number of vibrations of different velocities which are affected diversely in passing from one medium to another. This causes the ray of common or white light to be spread out into a number of coloured rays corresponding to vibrations of different rapidity. In all precise determinations, therefore, a definite point of the spectrum thus formed must be measured for refraction, and two such points for dispersion. The work of early observers on the refraction of light is usually wanting in precision in this respect, the records being commonly described as having reference to the bright or yellow part of the spectrum, or to the extreme red; while they furnish no data at all for calculating the dispersion. Moreover, as each colour occupies a certain breadth in the spectrum, and the boundaries of the colours are not well defined, it is by no means certain that two observations made on the same colour will refer to exactly the same part of the spectrum. In more recent observations the most conspicuous lines of the solar spectrum, such as A, D, F, H, are adopted, or, if artificial light be used, the α , β , and γ lines of the hydrogen spectrum; these are recorded as μ_{A} , μ_{D} , &c.; μ_{a} , μ_{β} , μ_{γ} . The greater part of the English data of recent times are given for the solar lines, while foreign observers have generally adopted the hydrogen lines.

Refraction and dispersion equivalents.

The index of refraction, minus unity, when divided by the relative density (d) and multiplied by the atomic weight if an element is examined, or by the molecular weight if a compound is examined (P), is usually represented by the for-

mula P
$$\frac{\mu-1}{d}$$
 in England and P $\frac{n-1}{d}$ on the

Continent, and is termed the refraction-equivalent. It will be expressed in this article by the simple letter R. When Lorenz's formula is used,

^{&#}x27; See, however, a recent investigation by M. Le Blanc (Z. P. C. 8, 299).

 $\left(\frac{\mu^2-1}{\mu^2+2}\right)\frac{\mathbf{P}}{d}$, the refraction-equivalent so calculated will be distinguished as R. The part of the spectrum to which the observation refers will be indicated by adding the sign for infinity, or the letters of the solar or other lines observed, as R_{∞} , R_{Λ} , R_a , &c. The dispersion-equivalent represents the difference between the refraction-equivalents of two selected portions of the spectrum; thus the dispersion-

equivalent for H-A will be expressed as $\frac{(\mu_H-1)}{d}-\frac{(\mu_A-1)}{d}$, or, more briefly $\frac{\mu_H-\mu_A}{d}$.

multiplied by P; or $R_H - R_A$.

The index of refraction of solids is generally ascertained by the method of total reflection, unless they are transparent, and can be cut into the form of a prism; that of gases is best determined by interference. The refraction of liquids is most conveniently arrived at by placing the substance in a hollow prism, with sides of plate glass; on the prism being so adjusted that the incident and emergent rays make equal angles with the refracting surfaces, the minimum deviation will be arrived at. Let D be the deviation, i and r the angles of incidence and refraction, and a the refracting angle of the prism; then

$$\frac{\sin e \ i}{\sin e \ r} = \frac{\sin e \ \frac{1}{2}(a + D)}{\sin e \ \frac{1}{2}a} = \mu.$$

Newton, acting upon the emission theory, proposed $\frac{\mu^2-1}{d}$ as representing the absolute refractive power of any substance, and this formula met with general adoption till recent times. Gladstone and Dale (T. 1863), as the result of a long series of experiments with different substances at various temperatures, found that the refractive index, minus unity, multiplied by the volume, gave nearly a constant. To $\mu-1$ they accordingly gave the name of 'refractive energy' as really representing the influence of the substance itself on the rays of light; and to $\frac{\mu-1}{d}$ that of 'specific refractive This empirical formula has been energy.'

adopted by most subsequent investigators; and Sutherland (P. M. [5] 27, 141) has lately advocated it on mathematical, as well as on physical, grounds. It holds its position side by side with the theoretical formula we have now to consider.

Lorenz's formula. H. A. Lorentz, of Holland (W. 9, 641), and L. Lorenz, of Copenhagen (W. 11, 70), came almost simultaneously to the conclusion, from purely theoretical considerations, based, in the case of the former, on the electro-magnetic theory, that $\frac{\mu^2-1}{(\mu^2+2)d}$ must express the relation between the velocity of the transmission of light and the density of the medium through which it is propagated. Landolt (B. 15, 64) tested the comparative merits of this formula, and that which he had previously adopted—viz. $\frac{\mu-1}{d}$ —and found that

so far as liquids were concerned there was nothing to choose between them, but that Lorenz's formula showed much less difference between the specific refractive energy of a liquid and that of its vapour. This is shown in the following table, in which it will be also noticed that the difference with the rise in temperature is uniformly plus in the one case, and minus in the other. Weegmann (Bonn. 1888) experimenting at 10°, 20°, and 30°, noted the same rise in the one case and fall in the other.

Substance	State	Temp.	$\frac{\mu_{\mathrm{D}}-1}{(\mu_{\mathrm{D}}+2)d}$	$\frac{\mu_{\rm D}-1}{d}$	
Water Ethyl alcohol Ethyl ether.	Liquid Vapour Liquid Vapour Liquid	10°	0-2062 0-2061 0-2068 0-2804 0-2807 0-2825 0-2026 0-2008	0.4582	
Ethyl acetate Ethyl iodide	Vapour Liquid Vapour Liquid	20° 100° 10° 20° 100°	0·3029 0·3068 0·2547 0·2549 0·2683 0·1557	0·4930 0·4599 0·4174 0·4172 0·4024 0·9062	
Chloroform . Carbon disulphide .	Vapour Liquid Vapour Liquid Vapour	20° 100° 20° 100° 100° 20° 100°	0-1557 0-1558 0-1571 0-1790 0-1791 0-1796 0-2805 0-2805 0-2809 0-2898 0-0089	0-2658 0-2356 0-3000 0-2996 0-2694 0-4977 0-4977 0-4977	

In the case of liquid and solid phosphorus, as well as of other highly refractive bodies, the empirical formula appeared to give the best results. Landolt also found the simpler formula best adapted to his process of optical analysis subsequently described. But while he found that the refraction-equivalents (i.e. the specific refractive energy multiplied by the atomic weight in the case of elements, or by the molecular weight in the case of compounds) of substances calculated by Lorenz's formula are about onethird smaller than the numbers calculated by the formula previously adopted by him, the values which he obtained for the elements carbon, oxygen, hydrogen, and chlorine did not bear the same proportion. Adopting Brühl's figures for Ra (as described later on), the values according to Lorenz's formula vary from about \(\frac{1}{2} \) to \(\frac{3}{4} \) the values found by using the other formula. Thus :-

	Old Formula	Lorenz's Formula
Carbon	5.0	2.48
Hydrogen	1.3	1.04
Oxygen, alcoholic	2.8	1.58
" aldehydic	3.4	2.34
Chlorine	9.8	6.02
Increase for double carbon bond	2.4	1.78

Brühl, however, in 1886 (A. 235, 1), maintained that the statement of Landolt, that the two formulæ lead practically to the same conclusions in respect to chemical constitution and the refraction of light, cannot any longer be accepted in its general sense. In this paper Brühl con-tended that the expression of the molecular refraction hitherto used—viz. P $\frac{\mu-1}{d}$ is generally

adapted only to the paraffinoid series of carbon compounds, and for unsaturated compounds of weak dispersive power; and that in these cases even the results are not altogether satisfactory. For the investigation of the relations between the refractive power and the constitution of bodies, Brühl claimed that in future Lorenz's formula must be used. The tables appended to the paper certainly show better proportionate results for \mathbf{R}_a by this formula, especially in the case of the most dispersive substances; but in using these tables it must be borne in mind that the values assigned to the elements under the two formulæ are not strictly comparable. Nasini (Lincei, Rc. 3, 128) stated that he and Bernheimer did not consider that, from a chemical point of view, any great preference should be given to Lorenz's formula over the formula $\frac{\mu-1}{d}$, as the newer formula was insufficient to

represent every relation between the chemical constitutions of the substances and their refractive powers. Ketteler does not accept the Lorenz formula as altogether satisfactory; in

place of it he has proposed $\frac{\mu^2-1}{(\mu^2+x)d}$, where x is

a variable quantity depending on the nature of the substance, but averaging about 3.5 instead of 2. Sutherland (P. M. [5] 27, 141) objects to Lorenz's formula on the ground that it assumes a mean wave length, whereas the wave length must vary in passing from ether to matter; and he maintains that while the formula holds good for both the liquid and gaseous states, it fails when change of density is produced by change of pressure. On the other hand, he thinks that Gladstone's formula meets the last requirement, but fails to bridge over the great gap in density between liquid and vapour. Ketteler (Z.P.C. 2, 905) suggests an equation of the form $(n^2-1)(v-\beta)=M$, where β is the volume actually occupied by the ponderable molecules, v that of the space in which they are contained, and M a molecular function depending on the constitution of the medium. M may have one of two values. both of which are definite constants, the one for the liquid and the other for the gaseous state. The value of β may be obtained by varying the temperature, the pressure, or the constitution of the media examined. Sutherland says that this formula gives for compressibility results less satisfactory than those given by that of Gladstone.

Zehnder (W. 34,91) investigated the influence of pressure upon the index of refraction of water at different temperatures, and on comparing the results with the co-efficient of compressibility, he

came to the conclusion that the formula $\frac{\mu-1}{d}$ was practically exact—much more so that the forenz. Röntgen and Zehnder (W. 44) have since applied the same inquiry to CS₂, and C_dH_d, ether, and a series of alcohols, with the result that in the cases of all these substances neither formula is sufficiently exact, the experimental figures occupying very nearly the mean place between those given by the two formulæ.

Perkin (C. J. 61, 287) in studying magnetic

Perkin (C. J. 61, 287) in studying magnetic rotation at widely different temperatures, observed a slight reduction as the temperature was raised. Dale and Gladstone (as long ago as 1868) had observed that the specific refractive energy of substances was similarly affected;

Nasini and Bernheimer (G. 15, 59) in 1885, and Ketteler (W. 33, 506, 662) in 1888, also found small differences in the same direction. Perkin accordingly made a series of experiments to $\mu = 1$

see if the changes in $\frac{\mu-1}{d}$ due to temperature

vary with different classes of substances, as they do in the case of the magnetic rotation. The results seem to corroborate this, though not very definitely. On trying Lorenz's formula, he found the refraction rise with increase of temperature, and he came to the conclusion that this formula is not suitable when dealing with high temperatures.

Guye (Ar. Sc. Genève, 23, 197, 204) pointed out a relation between the constants of the equation of Van der Waals and that of the formula of Lorenz for the refraction of light. He maintained that the relation of the absolute critical temperature to the critical pressure, which he termed the 'critical co-efficient,' is proportional to the molecular refraction, and that it furnishes a valuable means of determining the molecular constitution of bodies at the critical point. By a comparison of the data given by a number of observers for forty different substances he obtained the result that the critical co-efficient (K) is in proportion to \mathbf{R}_{∞} as 1 to 1.8. The experimental difficulties connected with the determination of K, and the fact that Guye had not always the data for reducing R observed to Roo, rendered the final results less uniform than they might be otherwise; but the extreme range of his factor varied from 1.6 to 2.0, giving 1.8 as a mean. Guye found, however, certain classes of bodies which gave a different factor to that above mentioned. Such gases as O, N, and CO, gave a factor ranging from 1.1 to 1.4. Water gave 1.1, and methylic alcohol 1.1 to 1.2. He also recorded some which are exceptionally high, but in most of these cases the results of the experiments seem untrustworthy in consequence of chemical action having taken place. Having determined his factor, he was able to divide by it the recognised refractions of the elements according to Brühl, and so to get what he termed the 'atomic critical co-efficient,' from which he could build up the critical co-efficient of any substance of which the theoretical constitution was determined.

Dispersion formulæ. As the observations of Gladstone extended to the solar line A, his calculations were but slightly affected by dispersion; whereas nearly all the Continental observers have used the hydrogen light, and have no actual measurements below the line a. They have, therefore, usually sought to get rid of the effect of dispersion by adopting one or other of the suggested formulæ for calculating the theoretical ray of infinite wave length, and have worked out their results both for the observed line a, and for the assumed limit. The simplest of these methods, and on that account the one most generally adopted, is that of Cauchy. It is usually expressed by the formula

$$\mu = \mathbf{A} + \frac{\mathbf{B}}{\lambda^2} + \frac{\mathbf{C}}{\lambda^4} + \dots$$

in which λ represents the wave length of any given line of the spectrum. As this formula contains three unknown quantities, it requires for their determination three simultaneous equa-

tions; but the third term is often dispensed with to reduce the amount of calculation required. By the process of elimination, the values of A, B, and Care obtained, A being Cauchy's co-efficient of refraction, and B and C the co-efficients of dispersion. This theory accords fairly with the experimental data for substances of moderate dispersive power, though by no means well with the data for substances of high dispersive power, such as oil of cassia. Wüllner (A. 133) considered that this formula was sufficiently exact for bodies of low dispersive power when two constants were used; but that for highly dispersive substances, such as carbon disulphide, it was necessary to go to the third term, as the difference between the observed value of $\mu\beta$ and that calculated from μ_{α} and μ_{γ} amounted to as much as 0.0022; while on using three terms the calculation was correct to the fourth decimal place. He also tried the formula of Christofle

$$n = \frac{n_0 \sqrt{2}}{\sqrt{1 + \frac{\lambda_0}{\lambda}} + \sqrt{1 - \frac{\lambda_0}{\lambda}}}$$

in which n_0 and λ_0 signify two selected constants, and n is the corresponding index of refraction of the wave length A; this also gave results which were not sufficiently accurate. Taking the observed indices for the lines a and γ of carbon disulphide at 0° as 1.634066 and 1.692148, this formula gave 1.669397 for β , while the observed value was 1.669076. At 20° the calculated value for \$ was as much as .000494 too high. Langley (P. M. [5] 17, 194) has discussed the relative merits of Biot's, Cauchy's, and Redtenbacher's formulæ, and has come to the conclusion that while each of them is sufficiently exact within the limits of the visible spectrum, they soon begin to give too small results for μ when that limit is passed. Biot's formula

$$\frac{1}{n^2} = a + b \left(\frac{n^2}{\lambda^2} \right) + c \left(\frac{n^4}{\lambda^4} \right) + k \left(\frac{\lambda^2}{n^2} \right)$$
 gives more trustworthy results than either of the

others.

Redtenbacher's formula $\frac{1}{n^2} = a + b\lambda^2 + \frac{c}{\lambda^2}$ has the serious objection that it reached a mini-

mum at a point corresponding to n = 1.5647 in the prism which he employed, so that for every value of n greater than 1.5647 it gave two different values for λ . Langley has experimentally determined wave lengths in the invisible prismatic spectrum far beyond the extreme limit possible according to Cauchy's formula. Brühl, after working out his results for R_{∞} as well as for R_{α} , has latterly (4. 235, 1; 236, 233) come to an adverse conclusion as to the utility of Cauchy's co-efficient of refraction for getting rid of the effect of dispersion, and he even asserts that when three terms are used the results are less to be relied upon for this purpose than when two terms only are employed. The following table shows the differences in Cauchy's A when calculated with two and three terms respectively, from the observed indices μ_a and μ_{γ} in the former case, and from μ_a , μ_{β} , and μ_{γ} in the latter; and also $\mu\beta$, as observed, compared with the same index of refraction as calculated by Cauchy's formula from μ and μ_{γ_i} showing in this case a difference of from one to five units in the third place of decimals :-

	Cauch	ıy's A	μ _β		
Substance	2 terms	3 terms	Ob- served	Cal- culated	
Benzoyl chloride Orthotoluidine Cinnamic alcohol Aniline Furfurol Cinnamic aldehyde	1.51906 1.53715 1.54307 1.54741 1.48269 1.55066	1.52700 1.54460 1.55253 1.55689 1.49816 1.58256	1.56964 1.58945 1.59998 1.60484 1.54566 1.65090	1 57097 1 59069 1 60151 1 60592 1 54824 1 65622	

In the case of substances of small dispersion, Brühl regards the use of Cauchy's formula as superfluous; with substances of medium dispersion, the results are generally preferable to those for the line a, though not to any great extent; but with substances of great dispersive power the differences are too great to be neglected in considering questions of chemical structure.

Nasini (Lincei, Atti, 18 [3]) comes to the same conclusion, that Cauchy's co-efficient of refraction ought to be rejected in studies of this nature, and that his co-efficient B is still more misleading if it be regarded as a measure of dispersion. N. (Lincei, Atti, 19 [8]) had already tested the merits of several formulæ, and given the preference to that of Lommel as affording better results than those arrived at by Cauchy's formula with two terms. The following table will show the differences between μ_s as observed, and as calculated by these two formulæ:-

Substance	μ ∌ 0ρ•	Calou	lated	Difference		
Dubstance	served	Cauchy	Lommel	Cauchy	Lommel	
	1.51339 1.60434		1·51369 1·60522	0.00065 0.00158	0.00088 0.00088	
naphthol . Dimethyl-	1.64597	1.6477	1.64678	0.00178	0-00081	
naphthalene	1.63200	1.6330	1.6324	0.00100	0.00040	

Weegmann (Bonn., 1888) gives reasons why Cauchy's formula for the limit of the spectrum is not to be relied upon; and maintains that his B, or co-efficient of dispersion, must not be taken as a measure of dispersive power, but that it should in any case be divided by the relative density.

Helmholtz has proposed the formula
$$n^2 - 1 = Q \frac{\lambda^4}{\lambda^2 - \lambda^2 m} - P \lambda^2$$

in which n and λ represent the index of refraction and the corresponding wave length, and Q, P and λ^2_m are three constants dependent upon the nature of the medium. Ketteler has

subsequently proposed another, viz.

$$n^2-1=\frac{A}{\lambda^2-B}+\frac{C}{\lambda^2-D}+\cdots$$

Brühl has tested the comparative merits of both these formulæ, as well as that of Cauchy. Taking Gladstone and Dale's observations of a solution of phosphorus in carbon disulphide, he calculated the values of the lines B, D, F, and G from the observed values of A, E, and H, and again those of A, B, E, and G from those of D, F, and H. The following table shows the differences in the two cases between the observed and calculated values (A. 286, 283):

Line	۲.	observed	Cauchy	Helmho.tz	Ketteler	Cauchy	Helmholtz	Ketteler
A B D E F G H	7·606 6·872 5·893 5·271 4·862 4·304 3·956	1.9527 1.9744 1.9941	 -3 -2 +8 +1	 -5 -4 +9 +2	 +2 -4 -7	'+19 + 9 -5 	+48 +21 -6 	+108 + 47 -9

Dispersive power. The difference between the indices of refraction of the rays of the spectrum of any refracting medium is termed the 'dispersion'; and by the dispersive power of a substance is expressed the ratio of the co-officient of dispersion to the index of refraction of the mean ray minus unity. Thus taking the full length of the visible spectrum as extending from A to H, and F as the mean ray, the dispersive power would be represented by the formula, $\frac{\mu_{\rm H} - \mu_{\rm A}}{\mu_{\rm F} - 1}$. Ketteler (Theoretische Optik, Braunschweig,

Ketteler (*Theoretische Optik*, Braunschweig, 1885) has proposed two formulæ for dispersion based upon theoretical grounds:

based upon theoretical grounds:
$$\frac{\mu_n^2 - 1}{\mu_m^2 - 1} \text{ and } \frac{\mu_n^2 - \mu_m^2}{\mu_n^2 - 1}$$

when n is a ray more refractive than m. Costa (G. 19) regards these as having the advantage over the formula $\frac{\mu_n - \mu_m}{d}$ on account of their being independent of the density of the medium.

Early experiments. Newton, Herschel, Young, Wollaston, Brewster, and others determined the indices of refraction of a large number of solids and liquids, in most cases for the yellow or brightest part of the spectrum. The index for the solids ranged from 1·111 for tabasheer to 2·974 for chromate of lead, and for liquids and soft solids from 1·0570 for ether expanded by heat to thrice its volume, to 1·678 for disulphide of carbon, and about 1·8 for chloride of antimony (Encyc. Brit., 8th edit. Optics, 558).

Arago and Fresnel applied the method of interference to the comparison of the refractive indices of moist and dry air, with the result that moist air was found to be rather less refractive than dry air. Dulong determined the indices of refraction of a considerable number of gases and vapours.

Dulong, together with Arago, Biot, and others, adopted the formula of Newton $\frac{\mu^2-1}{d}$, and came to the following conclusions; that the refractive power of a mixture of gases is equal to the mean of the refractive powers of the constituent gases calculated for the pressure to which each gas is actually subjected in the mixture, and that the refractive power of a compound gas is not equal to the mean of those of the component gases, but is sometimes greater and sometimes less.

Höck also found this formula to apply to some mixtures of liquids, and Schrauf (P. 133, 479) has more recently supported the same mode of calculating the refractive power; but Ketteler (P. 124, 390) found that liquid sul-Vol. IV.

phurous acid was not accordant with the known refraction of the gas as arrived at by the formula $\frac{\mu^2-1}{d}$, while the two came into agreement when he adopted the modern formula for the specific refractive energy $\frac{\mu-1}{d}$.

Jamin (A. Ch. [4] 3, 49) made careful experiments on the influence of the presence of aqueous vapour in the atmosphere upon the refractive index, and he found that the difference between the refractive indices of dry air and air saturated with aqueous vapour was only '000000726, a quantity so small that it may be safely neglected in the calculation of atmospheric refraction for astronomical purposes.

Ketteler (B. B. 1865) adopted Jamin's interference refractor for the measurement of the index of air, CO₂, CN, H, and SO₂, for the line D. He calculated the lithium and thallium lines from the wave length of the sodium line, according to the proportional number of rulings that coincide, it being found that this is independent of density and is a true function of the wave lengths.

RELATIONS BETWEEN PHYSICAL AND CHEMICAL PROPERTIES AND REFRACTION OF LIQUIDS.

Dale and Gladstone (T. 1858, 887) made an elaborate series of experiments on what they then termed the sensitiveness of liquids, or the effect of temperature on the refractive index. The principal conclusions they arrived at were that both the index of refraction and the length of the spectrum, $\mu_{\rm H} - \mu_{\rm A}$, diminish as the temperature rises, and that the sensitiveness of a substance is independent of its specific refractive or dispersive power. These conclusions were the result of observations on CS₂, Et₂O, H₂O, P, alcohols, &c. The following table and the table on p. 226 give the data for CS₂ and H₂O: —

Carbon disulphide (b.p. =43°)

Temp.	$\mu_{\mathbf{A}}$	$\mu_{ m D}$	μ_{H}	Sensi- tiveness	μ _Η μ _Α	Disper- sive power
0° 5° 10° 15° 20° 25° 30° 35° 40° 42.5°	1.6217 1.6180 1.6144 1.6114 1.6076 1.6036 1.5995 1.5958 1.5919 1.5900	1·6442 1·6397 1·6346 1·6303 1·6261 1·6220 1·6182 1·6140 1·6103 1·6082	1·7175 1·7119 1·7081 1·7035 1·6993 1·6942 1·6896 1·6810 1·6778	0·0045 0·0051 0·0043 0·0042 0·0041 0·0038 0·0042 0·0043 0·0042	0·0958 0·0939 0·0937 0·0921 0·0917 0·0906 0·0901 0·0894 0·0891 0·0878	0·1489 0·1467 0·1467 0·1463 0·1460 0·1457 0·1456 0·1459

The sensitiveness is reckoned from the indices of the line D, and the dispersive power repre-

sents $\frac{\mu_{\rm H} - \mu_{\rm A}}{\mu_{\rm D} - 1}$. The sensitiveness indicates an anomaly in water, which has also been observed by others.

Gladstone and Dale (T. 1863. 317) extended the same inquiry to the following five points:—
(1) the relation between sensitiveness and the change of volume by heat; (2) the refraction and dispersion of mixed liquids; (3) the refraction, dispersion, and sensitiveness of different members of homologous series; (4) the

Water											
Temp.	μ_Α	μ _D	μ _Η	Sensi- tiveness	$\mu_{\mathbf{H}}^{-}\mu_{\mathbf{A}}^{-}$	Disper- sive power					
00	1.3291	1.8330	1.3438	0.0001	0.0147	0.0429					
100	1.3290	1.3329	1.3436	0.0002 0.0003	0.0146 0.0146	0.0439					
15° 20°	1·3284 1·3279	1.8324	1.3431	0.0004 0.0003	0.0147 0.0148	0.0445					
25° 30° 35°	1.3276	1.8317	1.3420	0.0008 0.0006	0.0145 0.0145	0.0438					
400	1.3264	1.8308	1.3410	0.0008 0.0008	0.0146 0.0148	0.0449					
50°	1.3250 1.3241 1.3235	1.3288	1.8396	0.0008 0.0009	0.0146	0.0448					
60°	1.3223	1·3271 1·3259 1·3249	1·3380 1·3367	0.0012 0.0010	0.0145	0.0441					
70° 80°	1·3203 1·3178	1.8287	1·3344 1·3321	0.0012 0.0012A	0·0141 0·0143	0.0135					

refraction, dispersion, and sensitiveness of isomeric liquids; (5) the effect of chemical substitution on these optical properties. As to the first point of the inquiry, they came to the conclusion that the specific refractive energy

is a constant not affected by temperature. They made the reservation, however, 'that there is some influence, arising wholly or partially from dispersion, which gives rise to the slight progression of most of the calculated products, and perhaps to the non-inversion of the sensitiveness of water at 4°, remarked on already by Jamin and ourselves.' As to the second point of inquiry, it was found 'that the specific refractive power of a mixture of liquids is the mean of the specific refractive powers of its constituents,' subject to some slight exceptions, as in the case of sulphuric acid and water. The inquiry into the refraction, dispersion, and sensitiveness of different members of homologous series showed a progressive increase in the specific refractive energy as the series advanced; but that the influence of each addition of CH2, which is observable throughout the series of the methyl group, does not necessarily hold good with reference to substances of quite another type. The observations on isomeric bodies showed that some of them are widely different in their optical properties; but that those which have a close chemical relationship show an identity in their optical properties also. In the last branch of the inquiry it was sought 'to determine the amount of change in the optical properties which results from a replacement of one element by another, the type remaining the same,' in order to 'attain to a knowledge of the influence of the individual elements on the rays of light transmitted by them.' The general conclusion of Gladstone and Dale was that every liquid has a specific refractive energy composed of the specific refractive energies of its compound elements, modified by the manner of combination, and which is unaffected by change of temperature, and accompanies it when mixed with other liquids.' This opened up a question which has since occupied the attention of many observers.

Landolt (P. 117, 122, 123) took up the inquiry, adopting the formula $\frac{\mu-1}{2}$ used by

Gladstone and Dale, and applied it in the first instance to an investigation of the values of C, H, and O. He simplified the investigation by multiplying the specific refractive energy by the molecular weight of the compound, which gave P $\frac{\mu-1}{2}$ as the molecular refractive power, or refractionequivalent, terms which had already been applied by Berthelot and Schrauf to the formula P $\frac{\mu^2-1}{d}$. By comparing series of homologous compounds, acids of the series $C_nH_{2n}O_2$, alcohols of the series $C_nH_{2n+2}O$, and esters of the series

C_nH_{2n}O₂, L. obtained the increment due to each addition of CH2 as shown in the following selection from his data:-

_	Substance	S.G.	μ	$\frac{\mu_a-1}{d}$	Ra	Diff CH
Acids C _n H ₂ nO _s	Formio Acetic Propionic Butyric Valerianic Caproic Cannthylic		1·3955 1·4022 1·4116	0 3518 0 8860 0 4116 0 4319 0 4449	21·11 28·57 36·22 44 05 51·61	7·20 7·40 7·61 7·8: 7·50 7·79
Alcohols CnH _{2R+2} O	Methyl Ethyl Propyl Butyl Amyl		1·3794 1·3940	0.4501 0.4717 0.4879	20.70 28.30 36.11	7·53 7·60 7·81 7·78
Esters C _n H _e nO _e	Methyl acetate Ethyl acetate Methyl butyrate	0.9015		0 3967 0 4110		6.81
	Ethyl butyrate Methyl valerate	0.8906 0.8809	1.3940	0·4424 0·4458	51.32	7 35
	Ethyl valerate Ethyl formate Ethyl		1.3580	0*4554 0 3944	29.18	7·49 6·99
	Methyl butyrate Methyl butyrate Methyl valerate Ethyl	0.8976 0.8809	1.3869	0.4110 0.4311 0.4458	43.97	7.74
	butyrate Ethyl valerate.			0·4424 0·4554		7.88

The mean of Landolt's comparisons gave the value R=7.60 for CH2. Dealing in the same manner with compounds differing by one atom of carbon, by two atoms of hydrogen, and by one atom of oxygen, respectively, he assigned to the several elements the following refractionequivalents; C = 5.00; H = 1.30; Subsequent investigations would lead one to reject several of the substances which he brought into comparison.

Landolt also adopted a modification of Biot and Arago's formula for mixtures of gases, by substituting n for n^2 throughout, so that $\frac{N-1}{D}P = \frac{n-1}{d}p + \frac{n'-1}{d'}p' + \cdots$

$$\frac{N-1}{D}P = \frac{n-1}{d}p + \frac{n'-1}{d'}p' + \dots$$

N, D and P representing the index of refraction, density, and 'molecular weight' of the mixture, and the small letters the same values of the several substances forming the compound. In this manner he calculated the value of n_a or μ_a for mixtures of liquids as shown in the follow-

ing table. It will be seen that the calculated and observed values correspond very closely.

_			p	Obse d	erved , μ.	Calcu- lated μ_{\bullet}
Methyl alcoho Amyl "	l Mixture	:	96 88 184	0.7964 0.8135 0.8038	1.3279 1.4057 1.3640	1.3644
Ethyl alcohol Amyl "	 Mixture	:	92 88 180	0.8011 0.8135 0.8065	1.3605 1.4057 1.3822	1.3821
Ethyl alcohol Amyl "	Mixture	:	46 176 222	0.8011 0.8135 0.8104	1.3605 1.4057 1.3961	1-3960
Acetic soid Butyric ,	 Mixture	:	60 88 148	1.0518 0.9610 0.9930	1·3706 1·3953 1·3850	1.3847
Ethyl alcohol Formic acid	 Mixture	:	46 46 92	0·8011 1·2211 0·9602	1.3605 1.3693 1.3610	1:3612
Oil of bitter al Formic acid	monds. Mixture	:	106 46 152	1.0474 1.2211 1.0876	1.5391 1.3693 1.4900	1.4900

Landolt subsequently (A. 4, Suppl.) reversed the process, and applied it to the quantitative analysis of mixed liquids by means of their refractive indices and specific gravities. The following will serve as instances of the percentage results obtained:—

			μa-1	Percentage		
-	μα	d	$\frac{\mu_d-1}{d}$	Calcu- lated	Actual	
Amyl alcohol . Methyl ,, . Mixture .	1.4057	0.8185	0·4987	47·4	47·8	
	1.3279	0.7964	0·4117	52·6	52·2	
	1.3640	0.8038	0·4529	100·0	100·0	
Acetic soid . Butyric , . Mixture .	1.3706	1.0518	0·3523	40·3	40·5	
	1.3955	0.9610	0·4116	59·7	59·5	
	1.3850	0.9930	0·3877	100·0	100·0	
Ethyl alcohol . Formic acid . Mixture .	1.3606	0.8011	0.4501	49·8	50·0	
	1.3693	1.2211	0.3024	50·2	50·0	
	1.3610	0.9602	0.3760	100·0	100·0	
Oil of bitter almonds . Formic acid . Mixture .	1.5391	1.0474	0·5147	69·8	69·7	
	1.3698	1.2211	0·3024	30·2	30·3	
	1.4900	1.0876	0·4505	100·0	100·0	

Adopting these values, Haagen (\dot{P} . 131, 117) deduced values for Cl, Br, and I from some of their organic liquid compounds, and then deduced values for other elements from observations of R for their chlorine compounds. The following are the results he obtained for R_{∞} and R_{\bullet} :—

Ele	men	t	R_{∞}	Ra	
Chlorine Bromine Iodine.	:	:	•	9·53 14·75 23·55	9·79 15·34 24·87
Sulphur Phosphorus	•	•	•	14·74 14·60	16.03 14.93
Arsenic Antimony Silicon Tin Sodium	•	•	•	18·84 7·81 18·64 4·71	20·22 25·66 7·90 19·89 4·89

ments. Gladstone (T. 159, 13; Pr. 18, 49) applied Landolt's P $\frac{\mu-1}{}$ to inorganic compounds in order to arrive at the refractionequivalents of the several elements, and also to determine whether or not any of the elements had more than one definite refraction-equivalent. As most of his observations were made upon crystalline salts, many of which were doubly refracting, Gladstone adopted the plan of dissolving them in water, or alcohol, and deducting from the R of the solution the R due to the solvent. This mode of operation was supported by the consideration that in the cases of rocksalt and sugar, where the refraction-equivalent of the substance has been obtained both in the solid and dissolved condition, it is found to be the same. On comparing a series of salts of K and Na, it was evident that, while the refractionequivalents of the compounds differed very widely, according to the nature of their negative constituents, the refraction-equivalents of the compounds of Na differed pari passu with those of the K compounds, and hence it was

concluded that the negative constituent has the

same effect on light, whichever metal it is united

with. Values were arrived at for forty-six of the elements, a number which has been subsequently somewhat increased; and more receninvestigations have led to slight modifications in the figures. The latest list is as follows:—

Refraction-equivalents of the ele-

Element		At. w.	R
Aluminium		27.5	7.7
Antimony	. !	120	24.5
Arsenic	.	75	15.4
Barium	.	137.2	15.8
Beryllium		9.1	5.1
Bismuth		208	38.2
Boron, in borates .		11	about 4
Bromine		80	15.3
Cadmium		111.6	13.1
Cæsium		132	19.1
Calcium		40	10.0
Carbon	.	12	5.0
" doubly linked		,,	6.1
Cerium	.	138.2	19.6?
Chlorine		85.2	9.9
Chromium	. 1	$52 \cdot 4$	15.4
" in chroma	tes.	,,	about 22
Cobalt	. 1	<i>5</i> 8·7	10.4
Copper	.	63· 4	11.5
Didymium	•	145	23.3
Fluorine	-	19	1.6?
Gold	•	$196 \cdot 2$	23.1
Hydrogen		1	1.3
Iodine		127	24.5
	om-		1
pounds		56	11.6
	om-		1
pounds		,,	19.4
Lanthanum		138	22.9
Lead	• (207	24.3
Lithium	• ;	7	3.5
Magnesium	•	24	6.7
Manganese	. 1	55	11.7
" in perma	nga-		
nates	•	30	about 25
		•	Q 2

Element		At. w.	RA
Mercury		200	19.4?
Nickel		58.7	10.0
Nitrogen		14	4.1
" in bases, oxide	s.		1
&c	•	,,	5.1
Oxygen, singly linked		16	2.8
., doubly linked		,,	3.4
Palladium	. 1	106	21.6?
Phosphorus		31	18.3
Platinum		195	24.7
Potassium		39.1	7.85
Rhodium		103.4	23.4?
Rubidium		85.4	12.1
Selenion		78	30.1
Silicon		28	7.4
,, in silicic acid		"	about 6
Silver		108	13.2
Sodium		23	4.4
Strontium		87.5	13.0
Sulphur		32	16.0
" singly linked		,,	14.1
Thallium		203.6	20.4
Tin		118	27.0?
			18.6?
Titanium		48	24.6
Uranium		237.6	19.4
Vanadium		51.3	24.8?
Zinc	•	65.3	9.8
Zirconium	•	90	21.3
WII COMINIE		00	. 410

Anomalies were pointed out in the refractionequivalents of the hydracids, and of the benzenoid carbon compounds. With regard to the specific refractive energy it was remarked :- First, hydrogen has more than double the energy of any other element, even in the lowest number that can be assigned to it. Second, phosphorus, vanadium, titanium, and sulphur have singularly high energies, and they are substances that present certain chemical analogies. Third, there are several pairs of analogous elements having nearly the same energy; thus bromine and iodine, arsenic and antimony, potassium and sodium, manganese and iron, nickel and cobalt. Fourth, an element in altering its valency alters its energy. Fifth, if the metals are arranged in the order of their energies, they are, with few exceptions, in the inverse order of their atomic weights.

Benzenoid hydrocarbons.—The anomalous refraction and dispersion of the hydrocarbons already referred to was treated in a paper by Gladstone (C. J. 23, 147), the tables in which indicated that the values for all these bodies and their derivatives were largely in excess of the normal, calculating $\mathbf{R}_{\mathbf{A}}$ of carbon at 5.0. The refraction-equivalents of the typical hydrocarbons were expressed as in the following table; but while the physical fact indicated has been fully verified by subsequent research, the particular numbers in the third column have been modified, and a plus has been assigned to the olefines.

A similar table is given illustrating the increased influence on light of carbon which is combined with two atoms of hydrogen or one of oxygen, in a series of oxidised bodies obtained from essential oils, differing from one another only in the quantity of hydrogen.

Hydrocarbo	ns	Typical Formula	R
Paraffins . Olefines . Terpenes . Benzenes . Naphthaleue Anthracene		C_nH_{2n+2} C_nH_{2n} C_nH_{2n-4} C_nH_{2n-6} C_nH_{2n-12} C_nH_{2n-8}	Normal ,, + 3 ,, + 6 ,, + 14 ,, + 17

Bedson and Williams (B. A. 1881. 155) tested the question whether the specific refraction of a solid body could be determined from the refractive power of its solution as stated by Dale and Gladstone, as this method had been called in question by Janovsky (Sitz. W. 82, 148). They found the specific refractive energy for the ray of infinite wave length, calculated from solutions of NaCl, Na₂B₄O₂, H₃BO₃, and NaPO₃ to be in substantial accord with the results derived from observations on the solids. They also found that the specific refraction of liquid phenol is practically identical with the value calculated from solutions in alcohol and in acetic acid.

In taking the index of refraction of solid NaPO₃, B. and W. immersed it in a liquid of greater refractive power, and then added a feebly refractive liquid until the refractive power of the mixture appeared to be the same as that of the solid suspended in it. The index of refraction of the mixture was then determined. The result being found satisfactory, the prisms of rocksalt and of fused borax were also tried by the same process, as well as in the ordinary way, and with similar results.

Double values of carbon and oxygen. Brühl, in 1879 and 1880 (A. 200, 139; 203. 1, 255), conducted a long investigation into the chemical constitution of carbon compounds in reference to their relative densities and their power to refract light. Taking up the observations of Gladstone, and also referring to some of the anomalous results in the tables issued by Landolt, whose mode of investigation suggested that the grouping of the atoms exercised a certain influence upon the refraction of light, Brühl set himself to inquire: What is the typical peculiarity which must be common to all the bodies that refract light in an abnormal manner? He usually determined the indices for the three H lines, Hand μ_{β} , and μ_{γ} , and for the Na line μ_{D} , and calculated the refraction-equivalents both for the line a, and for Cauchy's A, viz. R_∞ . The following table is so arranged that between each horizontal line the first-named substance has the highest molecular weight, while those below differ only in having a smaller quantity of hydrogen, and, therefore, contain doubly-linked carbon atoms.

It will be observed that the density, the index of refraction, and the co-efficients of refraction and dispersion of Cauchy always increase with the diminution of H, and that the loss in specific refractive energy which is suffered by the disappearance of two atoms of H is at least fully made up by the consequent double linking of the carbon atoms. For the calculated values of the refraction-equivalents, Brühl has adopted the following values for the elements, viz.—for R_a C=5.0, H=1.3, O=3.0, Cl=9.8, Br=15.3,

		s.G.		Cauchy's	Cauchy's	u _a -1	1	R.	
Substance	Formula	3.σ. μ		A A	B	$\frac{\mu_a-1}{d\psi}$	Ob- served	Calcu- lated	
Propyl alcohol . Allyl alcohol .	C,H,O C ₃ H ₆ O	0·8044 0·8540	1·38345 1·41051	1·37542 1·39881	0·34630 0·50453	0·4767 0·4807	28·60 27·88	28·4 25·8	
Propyl aldehyde. Acrolein	C ₃ H ₄ O	0·8066 0·8410	1·36157 1·39620	1·35344 1·38010	0·35065 0·69427	0·4483 0·4711	26·00 26·38	25·8 23·2	
Propyl ether Allyl ether	$C_3H_7(OC_2H_5)$. $C_8H_5(OC_2H_5)$.	0·7386 0·7651	1·36758 1·38565	1·35975 1·37547	0·33758 0·43882	0·4977 0·5041	43·80 43·35	43·6 41·0	
Propyl acetate . Allyl acetate .	C ₃ H ₇ (C ₂ H ₃ O ₂) C ₃ H ₅ (C ₂ H ₃ O ₂)	0·8856 0·9276	1·38235 1·40205	1·37427 1·39151	0·34832 0·45458	0·4317 0·4334	44·04 43·34	44·0 41·4	
Propyl chloride . Allyl chloride .	C ₃ H,Cl C ₃ H ₅ Cl	0·8898 0·9379	1·38659 1·41245	1·37813 1·40007	0·36474 0·53369	0·4345 0·4398	34·11 33·64	33·9 31·3	
Isobutyric acid . Methacrylic acid .	$C_4H_8O_2$ $C_4H_8O_2$	0·9490 1·0153	1·39093 1·42815	1·38259 1·41400	0·35971 0·61013	0·4119 0·4217	36·25 36·27	36·4 33·8	
Amylene (34° to 35°) Valerylene	C,H ₁₀ C ₃ H ₈	0·6476 0·6786	1·37330 1·39763	1·36352 1·88565	0·42173 0·51660	0·5764 0·5680	40·35 39·85	38·0 35·4	
Hexane Diallyl	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	0.6603 0.6880 0.8799	1·37337 1·39180 1·49668	1·36538 1·38589 1·47562	0·34462 0·52733 0·90816	0·5655 0·5787 0·5645	48·63 47·45 44·03	48·2 43·0 87·8	
Triethylamine . Aniline	C ₆ H ₁₅ N	0·7277 1·0216	1·39804 1·57948	1·38803 1·54741	0·43179 1·38319	0·5470 0·5672	55·25 52·75	55·3 44·9	

N = 5.8; the first three being the values given by Landolt, the two halogens those by Haagen, and that for N being determined by Brühl himself. From the observations recorded in the foregoing table and others, Brühl has arrived at the following conclusions:-That the closing of the chain of atoms in the form of a ring, with which so marked an increase in the density and index of refraction is associated, exercises a very special influence on the power of the substances to disperse light; that the mol. refraction of the substances in which the existence of one or more pairs of doubly-linked carbon atoms is recognised is greater than the sum of the refractions of the constituent atoms by 2.0 in the case of Cauchy's limit, and 2.3 in the case of the line Ha for each such pair. For combinations of the acetylene type, including the propargyl derivatives, he gives a rather lower value, viz.: 1.8 and 1.9 respectively. These data give the following atomic refractions for zarbon, viz .-

4.86 5.00 . 5.86 6.155.76 5.95

Nasini and Bernheimer (Lincei, Atti, 18) do not, however, admit Brühl's views as to the constitution of the propargyl compounds, as well as some of the other conclusions at which he has arrived. They give tables of observations on naphthol compounds, styrol, cinnamic alcohol, anethol, anisol, thymol, and diamylene, and have worked out the results by the two increases the greater is the difference between

 R_{∞} and R_{∞} , as well as showing the small differences both in specific refraction and specific disperson due to differences in temperature. The principal conclusions at which they arrive are as follows:-The statement that every double bond, and only every double bond, causes the mol. refraction to increase by a constant quantity is not confirmed by complete series of substances, because in compounds containing only four or five double bonds the increase in refraction indicates a much greater number. That the increase in mol. refraction grows step by step as the compounds become poorer in H; but no proportion exists between the increased values and the number of pairs of atoms of H that is lost; nevertheless, a simple relation exists with the other structural changes in the formula, expressed by double bonds, closed chains, &c. That the augmentation increases more or less rapidly from the paraffins to the olefines, and thence to the benzenes, and again to the naphthalenes, but by what law is not evident, and is not capable of expression by our structural formulæ. They consider it established, however, that the principle is approximately valid, that in every series of compounds each atom in the molecule furnishes its fixed contribution of refraction, and that in the paraffin series the mean values of the atomic refractions coincide with those derived directly from the elements themselves. They remark that even in the paraffin series the constancy of the atomic refractions is not absolute; because the more the mol. w. ordinary formulæ for Ra and Ra, and also for the calculated and experimental values. For other series the same values no longer apply, being too low, and hence other constants are required.

Kanonnikoff (Bl. [2] 36, 557) has published observations confirming Brühl's calculation of the increase due to the double bond between carbon atoms, and of the loss in specific refractive energy due to the elimination of two atoms of hydrogen being counterbalanced by the increase due to the consequent double linking of the carbon atoms.

Landolt (P. 117, 122) had determined the refraction of oxygen from a comparison of the refraction-equivalents of various substances the empirical formulæ of which differed by one atom of oxygen; and from acids of the formula $C_nH_{2n}O_2$ by deducting the value of n times CH_2 . The results were somewhat discordant, the first series giving a mean value for oxygen of 2.76, and the latter of 3.0. The former of these manifestly represented the atomic refraction of oxygen in hydroxyl; and if oxygen varied in its refraction according to its valency it is evident that a general average resulting from these two modes of determination could not be taken. Brühl shows that, adopting Landolt's second plan for arriving at the value of oxygen in the aldehydes and ketones of the formula $C_nH_{2n}O$, the \tilde{R}_a for oxygen will average 3.4; and that chloral, butyl-chloral, and the oxychlorides give substantially the same result. On the other hand, deducting the refraction-equivalents of the aldehydes from those of their corresponding acids, of the alcohols from the glycols, and of the acids from the oxyacids, he obtains 2.75 as the mean value for oxygen. An approximate result, 2.80, is obtained if, from the ethers, $C_nH_{2n}O_2$, the value of n times CH_2 + the higher value (3.4) for one atom of oxygen be deducted. For oxygen united to carbon by a double bond, Brühl therefore assigns the refraction-equivalent 3.4 for the line a; and for oxygen in hydroxyl, or generally for oxygen united to two other atoms, he gives the value 2.8; he represents the former as O'' and the latter as O'.

Hence C=0 = 5.0 + 3.4 = 8.4. C=0 = 5.0 + 2.8 = 7.8.

Whether, however, the increased result of the double linking of the carbon and the oxygen is due to one of the elements only, or whether both of them share in it, he leaves an open question, though he regards the latter as the more probable. If two atoms of H are removed from a substance without involving the double linking of carbon atoms, as in the case of an alcohol being converted into an aldehyde or ketone, there follows a very marked reduction of the specific refractive energy. This reduction, however, becomes less and less marked as the quantity of C increases in the higher members of the series. When the removal of two atoms of H involves the double linking of the C, the loss of refractive energy is more than compensated by the increase due to the doubly-linked C.

A comparison of a considerable number of isomeric bodies of the paraffinoid series of acids shows that the indices of refraction of the primary and normal compounds are a little greater than those of the secondary and iso-compounds, and that the densities are proportional. In the case of unsaturated compounds of similar

empirical composition, however, the double linking of the carbon atoms increases the refractive power more than the density.

In a subsequent paper (A. 235, 1), Brühl treated of the molecular refraction of organic liquids of high dispersive power. He arrived at the conclusion that the dispersion of different substances stands in no clearly recognisable relation either to their refraction or chemical constitution; and that it cannot be numerically expressed. This, however, is contested by Gladstone (Pr. 42, 401), who, while admitting that there are difficulties in the investigation of dispersion which are not felt in dealing with refraction, still holds (1) 'that dispersion, like refraction, is primarily a question of the atomic constitution of the body: the general rule being that the dispersion-equivalent of a compound is the sum of the dispersion-equivalents of its constituents; (2) that the dispersion of a compound. like its refraction, is modified by profound differences of constitution, such as changes of atomic valency; (3) that the dispersion frequently reveals differences of constitution at present unrecognised by chemists, and not expressed by our formulæ.' Bruhl held that the differences which still appear between the calculated and experimental values when Lorenz's formula is used, and which he showed to be less in proportion than when the old empirical formula is used, are principally due to the influence of dispersion, and that they would in great measure vanish if this disturbing cause could be overcome; that with substances with a dispersion equal to about that of cinnamic alcohol the molecular refraction for the line a would serve as a help to the determination of the chemical constitution, but beyond that limit the numerical results are too uncertain to be of use except as collateral evidence; that there is a direct and simple proportion between the number of unsaturated groups of atoms in any compound and the increase of the molecular re-

fraction $\left(\frac{\mu^2-1}{\mu^2+2}\right)\frac{p}{d}$ over the value derivable

from the empirical chemical formula; that the effect on refraction of the acetylene combination, which he represents by \equiv , is analogous to that of the ethylene, or doubly-linked carbon, =, and that it is somewhat greater, but not twice as great, as the latter.

Gladstone questions Brühl's assumption that both cinnamic alcohol and cinnamic aldehyde have four pairs of doubly-linked carbon atoms, on the ground that they have widely-different dispersions; also that allyl paracresolate and anethol are similarly constituted, for the same reason. He also thinks that the evidence as to the constitution of cymol and hexahydro-naphthalene from Brühl's observations is not conclusive as to the one having three pairs, and the other two pairs, of doubly-linked carbon atoms, seeing that they have nearly the same dispersion. Nasini (*Lincei*, Rc. 3, 164) in like manner holds that these particular compounds do not justify the conclusions that Brühl has come to respecting them, and agrees with G. that the dispersion may be a constant, and a much more sensitive one than the specific refraction; and that as such it is affected more and varies more

by certain little differences of composition and constitution of which the specific refraction does not feel the influence.

Brühl (Z. P. C. 1, 6) maintains that Erlenmeyer's constitutional formula, with five ethylene combinations, correctly represents the structure of naphthalene, and those derivatives in which hydrogen is replaced by monovalent elements. But the results of the observations on which he bases this conclusion all show a larger increment than can be accounted for by this formula; the excess he attributes mainly to the influence of dispersion. In this and all the other inquiries he has adopted Lorenz's formula, and the solar line C, without attempting to get rid of the influence of dispersion by the use of Cauchy's co-efficient of refraction. In the same paper he formulates the following six propositions:-(1) Isomerides of similar atomic arrangement (Stellungsisomere) have similar specific and molecular refractive powers; on the other hand, those isomerides which show differences of saturation (Sättigungsisomere) have various specific and molecular refractive powers; (2) polymeric combinations never show similar specific refractive energy, or multiple molecular refraction corresponding to the molecular weights; (3) transformations of multiple linkings of atoms into simple ones have as their result a diminution of refraction, equally, whatever the kind of chemical change may be (polymeric or isomeric change); (4) the optical effect of the accumulation of multiple linkings is the same, whether it produces open chains (amylene-diamylene) or ring formations with one or more groups of rings (as in paraldehyde, cymhydrene, menthol, &c. in the one case, and in turpentine and cajeputol in the other); (5) the molecular refraction of a fully saturated body is very nearly that of the sum of the refractions of the atoms of the empirical chemical formula, each of these being reckoned as united by single bonds—all those are to be regarded as saturated bodies in which there is no multiple linking of the atoms, such as the true paraffins, or derivatives of the general formula $(C_nH_{2n+2})-xH_2$; (6) all unsaturated bodies show an increased refraction which is proportional to the number of ethylene, acetylene, or carbonyl combinations present; the proportion, as a rule, is the more exact the less the substances in question are distinguished by a specially great dispersion. While holding to the opinion that the monovalent elements have virtually each but one refraction equivalent, and that the change of valency in the others is the chief cause of their variable refraction, Brühl considers that exact values cannot be obtained, because the mode of the grouping of the atoms may not be altogether without in-

In the meantime, Gladstone (C. J. 45, 241) had brought together a large number of observations on organic compounds, both liquids and solutions of solids, and placed against them the refraction-equivalents calculated on the basis of the following values for the elements:—

Carbon, saturated.		$\mathbf{R}_{A} 5.0$
$\lim_{n \to \infty} C_n H_n$.		. 5.95
. doubly linked	•	. 6.1
Hydrogen		. 1.3
Oxygen, singly linked		. 2.8
" doubly "		. 3.4

Nitrogen						4.1
"		bases,	NO2,	&c.		5.1
Chlorine						9.9
Bromine					•	15.3
Iodine						24.5
Sulphur,						14.1
	do	nhlv li	nked			16.0

Gladstone's observations confirm Landolt's original determination for H, and also for C in saturated compounds. Gladstone points out that the hydrocarbons of the formula C_nH_n scarcely come up to Brühl's previous estimate of the higher value of C, so that in these cases he has adopted the figure 5.95 instead of 6.1, as the difference is more than he considers can be accounted for either by dispersion or by the distance from the boiling-points. In the case of those substances where the C atoms outnumber those of H, all of which show an inordinately high refraction and are at the same time enormously dispersive, Gladstone points out that Kanonnikoff's view that the increase is due to dispersion alone is not established by that observer's own figures for the theoretical limit of the spectrum according to Cauchy's formula. He admits, however, that if the limit were carried back considerably further, though still fairly within what Langley's observations would allow, the values would so far approximate as to render it not impossible that the refraction-equivalent of C might prove to be the same in these compounds as in the benzenoid series, or wherever it is doubly linked. As to the halogens, the figures in the table are generally consistent with the values which were originally assigned by him to Cl, Br, and I, viz. 9.9, 15.3, and 24.5 respectively. These numbers are very nearly the same as those arrived at independently by Haagen, viz. 9.8, 15.3, and 24.8. The values for O are taken from Brühl's original determinations, though the evidence of the alcohols rather favours his old estimate of 2.9 for the line A.

With regard to the phenomena of dispersion, Gladstone points out 'that the double linking of carbon atoms, while it does not much affect the specific refraction, widens the spectrum greatly, to such an extent indeed that where there are three pairs of such carbon atoms the specific dispersion is about double that of a saturated compound; where, however, the carbon atoms are actually in excess the dispersion becomes far more rapid. This is evident throughout, but the following typical hydrocarbons may illustrate it:—

Hexane . C.H. sp. ref. 0.5626, sp. disp. 0.0242 Amylene . C_5H_{10} 0.5708 0.0300,, ,, Benzene . C.H. 0.55950.0486,, ,, NapthaleneC10H8 0.5870 0.0782,, ,, Difluorene C26H18 0.6057 0.1108

or the following alcohols :-

Isobutyl alcohol C.H., O sp. ref. 0.4878, sp. disp. 0.0209 Allyl alcohol C.H., O "0.4734" 0.0275 Phenol C.L., O "0.5090" 0.0075 Naphthol C., H., O "0.5090" 0.00832

'The effect of the halogens in lowering both the specific refraction and dispersion is very marked, and that of NO₂ is still more remarkable.'

Gladstone (C. J. 59, 290, 589) has since published other long series of observations; including a record of experiments on the effect of changes of temperature on the refraction of

some highly-refractive substances, which seem to confirm the opinion previously expressed by Dale and himself. Among the substances contained in these tables the benzyl and benzal camphors of Haller call for special notice, as they have a very high refraction, and a dispersion which is altogether excessive. The latter compound has in like manner a very high rotatory power. The hydracids in aqueous solution are shown to have not only a higher value than the normal, a fact which has long been observed, but also that the value rises with dilution; and that in the dispersion the excess is still more striking. The same fact was observed by Perkin in regard to the molecular magnetic rotation of these acids dissolved in water, both as to the initial excess, and to its increase with the dilution up to a certain point where it remained nearly constant.

Other investigators have availed themselves of the optical properties in order to determine the constitution of compound bodies, or to confirm the views arrived at on other grounds. Knops (A. 248, 175) has determined the indices of refraction of the methyl, ethyl, and propyl salts of fumaric, maleic, mesaconic, citraconic and itaconic acids. A comparison of these acids with their ethereal salts leads to the conclusion that fumaric and maleic acids contain only one doubly-linked carbon atom; and that fumaric acid stands in the same relation to maleic acid as mesaconic acid does to citraconic acid.

Kanonnikoff (J. pr. [2] 31, 321, 497) has made a long series of observations on solutions of solids, of which he tabulates seventy-nine different substances, using water, alcohol, benzene, and chloroform as the solvents. As a preliminary inquiry, he reviewed the work of former investigators, and in some cases added experiments of his own to confirm previous conclusions: that the refractive energy of a substance is not changed in passing from the state of a liquid to that of a gas or vapour; that the same applies to bodies in passing from the solid to the liquid condition; that the refraction-equivalent of a mixture of liquids is the sum of the refraction-equivalents of its constituents; and that if the refraction-equivalent of a mixture and that of one of the components, together with the proportion of the same, is known, the refraction-equivalent of the other component can be calculated; that the same law applies to solutions of solid bodies, and that like results will be obtained whatever the solvent may be; that a derivative will be comparable with the substance from which it is derived, though the one may be a liquid and the other a solid body, as in the case of mesaconic acid and ethyl mesaconate. From the results of the observations above referred to, K. has determined the refraction-equivalents of fourteen of the elements, as follows :-

Element	Ro	R_{∞}	Element	\mathbf{R}_{σ}	R_{∞}
Lithium . Sodium . Potassium Copper . Rubidium . Silver . Cæsium .	8·16 4·22 7·75 — 12·04 18·05 19·55	3·00 4·00 7·40 11·30 11·60 12·53 18·70	Magnesium . Calcium . Zinc . Strontium . Cadmium . Barium . Mercury .	7.08 9.82 9.80 11.61 13.03 15.84 19.20	6·47 9·05 9·40 11·25 12·66 15·28 17·90

K.'s values for the solar line C will be found to be generally slightly lower than those determined by Gladstone for the line A. The refraction-equivalents for the theoretical limit of the spectrum were also calculated by Cauchy's formula from observations of the lines α and β of hydrogen. K. also gave determinations of refraction-equivalents of thirty organic liquids, showing the excess due to the double linking of the carbon atoms wherever this occurred.

Nasini (Lincei, Rc. 1, 1) points out that the suggestion thrown out by Gladstone that there may be a third and higher value of carbon when its four bonds are satisfied by other carbon atoms, is supported qualitatively by the researches of Bernheimer and himself, but not quantitatively, and that, therefore, this hypothesis is not a sufficient explanation of the facts. He argues as follows: naphthalene derivatives have two such atoms, while cinnamic alcohol has but one, whereas the values differ but very little. Anethol with one such carbon atom has a much greater excess than the naphthalene derivatives, while styrol shows so slight an excess, that on this hypothesis it could hardly be allowed to have any carbon atom in this condition. He compares two pairs of isomeric bodies, both as to refraction and dispersion,

Substance		Ra	$\frac{\mu_{\beta}-\mu_{\alpha}}{d}$
Allyl phenylate.	•	70·45	0·0173
Cinnamic alcohol		73·83	0·0220
Allyl paracresolate	•	78·79	0·018
Anethol		82·95	0·0257

from which he infers that when the allyl group is directly united to the benzene nucleus the molecular refraction greatly increases, whereas when the union takes place through the intervention of oxygen no augmentation takes place; so far, therefore, while supporting the hypothesis of Gladstone, he does not find any proportionality between the number of such carbon atoms and the increase of refraction.

Nasini (Lincei, Rc. 3, 164) criticises some of the conclusions of Brühl in the paper on the molecular refraction of organic liquids of high dispersive power (A. 235, 1) already referred to; and in particular he points out that the elimination of dispersion, if that were possible, would not get rid of the discrepancies between the observed and calculated values for the refraction of substances of high refractive power, and he cites observations on a series of substances in which he finds the constant fact that on adding a saturated side chain to an aromatic nucleus the specific dispersion diminishes while the refractive power increases.

Costa (G. 19) has made observations with the special object of testing the effect on dispersion of adding saturated side chains to a benzene or naphthalene nucleus. For this purpose he used compounds of amyl with benzene, thymol, eugenol, resorcin, and naphthol, and found in each case a still further increase in the refraction-equivalent than could be accounted for by the nucleus itself, while at the same time the specific dispersion showed a decrease.

Dispersion equivalents. In the same way that Gladstone arrived at the refraction-equivalents of the elements, he has sought also to determine the dispersion-equivalents of several of them; and (Pr. 42, 401) he has given the results of his observations upon nine of them, together with the values for the combinations CH_2 and NO_2 . The values are as follows:—

Substance	$\mathbf{R}_{\mathbf{A}}$	Disp. Eqt. H-A	
Phosphorus	18.3	3.0	
Sulphur, doubly linked	16.0	2.6	
" singly "	14.0	1.2	
Hydrogen	1.3	0.04	
Carbon	5.0	0.26	
,,	6.1 ?	0.51	
,,	6.1	0.66	
Oxygen, doubly linked	3.4	0.18	
" singly "	2.8	0.10	
Chlorine	9.9	0.50	
Bromine	15.3	1.22	
Iodine	24.5	3.65	
Nitrogen	4.1	0.10	
CH.	7.6	0.34	
NO.	11.8	0.82	

It will be observed that G. gives three dispersion equivalents for C, which he considers pretty well established, and he raises the question whether there may not be an intermediate refraction-equivalent corresponding to the dispersion-equivalent 0.51, which is found in such bodies as the allyl compounds and olefines. In his previous paper (Pr. 31, 327), G. suggested that there is a still higher refraction-equivalent of C in those cases in which it has four bonds satisfied by other C atoms, as in naphthalene and pyrene. Whatever may be the truth of this view, it appears that the dispersion-equivalents of these bodies are enormously high — much beyond what is recognised in the above table.

Brühl (Z. P. C. 7, 2, 140) has since modified his views upon the subject of dispersion; and has worked out carefully the atomic dispersions of some of the most important elements that enter into organic compounds. In the course of this investigation he has been led to revise his previous figures for the refraction as well as the dispersion. They now stand as follows:—

	Symbol	Ra	Ry-Ra
Carbon, singly linked .	C'	2.365	0.039
Hydrogen	H	1.103	0.036
Oxygen (hydroxyl) .	0'	1.506	0.019
" (ether) .	0< 0"	1.655	0.012
" (carbonyl) .	O"`	2.328	0.086
Nitrogen, linked singly			
with C	N'	2.76	0.19
Chlorine	Cl	6.014	0.176
Bromine	Br	8.863	0.348
Iodine	I	13.808	0.774
Ethylene bond	-	1.836	0.23
Acetylene bond	=	2.22	0.19

It will be observed that B. retains Lorenz's formula; and that he takes the difference between the refraction of the hydrogen lines γ and α as his measure of dispersion. He now acknowledges that the molecular dispersion can be re-

garded and employed in like manner with the molecular refraction as a specific expression of the material composition of chemical bodies; and that the dispersion is decidedly more sensitive to structural influences than the refraction. His inquiries have convinced him that refraction and dispersion are in no general sense correlative properties; but that certain structural peculiarities that have much influence upon refraction have little upon dispersion, while others have the opposite effect, and, as a rule, exercise a much stronger influence upon the dispersion. It is evident that this must be so, from the very high relative dispersion of O" as compared with that of O' or O<, and the large proportionate dispersion due to the ethylene bond. This figure, ·23, B. does not, however, recognise as a constant value. These revised values have since been used by him in his inquiries into the relations between the optical properties and chemical constitution of epichlorhydrin, the aldehydes and benzoyl (B. B. 24, 656), in which he draws attention to the concurrence of the evidence derived from chemical, calorimetrical, and optical investigations.

Essential oils. Gladstone (C. J. 17, 1; 25, 1) gave a long list of observations in which he recorded the circular polarisation as well as the index of refraction. The power of the oils in rotating the plane of polarisation appears to be very diverse, both in degree and direction; but he suggests that a comparison of the specific gravity and of the length of the spectrum may in some cases be of service in determining the genuineness or otherwise of the oils. The hydrocarbons obtained from these oils (of which a long table is given) divide themselves into two great groups, which are more fully investigated in the second paper.

Subsequently (C. J. 49, 609) the effect of the higher refraction and dispersion of C when united by double bonds was applied to the determination of the structure of the essential oils. Taking 2.2 as the increased refraction for the solar line A, and 0.8 as the increased dispersion. H-A, due to one pair of doubly-linked carbon atoms, the terpenes indicate, both by their refraction and dispersion, that they contain only one pair of carbon atoms doubly linked, thus corroborating the views of those who, on chemical grounds, have held that a terpene requires the addition of only two atoms of hydrogen, or their equivalent, to saturate it. The refraction and dispersion of solid inactive camphene also show that this substance does not contain more than one pair of doubly-linked carbon atoms, which is consistent with the fact that it combines with only one molecule of hydrochloric acid, and is directly converted by oxidation into camphor C10H18O, which is certainly a saturated compound both by chemical and optical evidence. In the same way, the citrenes give a refraction and dispersion indicative of the presence of two pairs of doubly-linked carbon atoms. Allied to these are several hydrocarbons having different amounts of hydrogen, and the number of doubly-linked carbon atoms varying accordingly. Thus, while citrene $C_{10}H_{16}$ has two pairs, cymene $C_{16}H_{14}$ has three, while menthene C₁₀H₁₈ has but one pair of doubly-linked carbon atoms, and cymhydrene C10H20 has none, being a saturated body. Pentene and isoprene C_5H_8 are identical in their optical properties, and indicate that no less than four of the five carbon atoms are doubly linked, which accords with the fact that these compounds combine with four atoms of bromine or two molecules of hydro-phloric acid. The following table will illustrate the foregoing conclusions:—

μ_{IA} 2·408449 μ_{Na} 2·417024 and 2·417227 μ_{T1} 2·425487

These figures give R=4.8, a result slightly lower than his earlier observations on the diamond, in which the index for the red part of the spectrum is recorded as 2.434, and the S.G. 3.55.

Selenion. Sirks (P. 143, 429) made

			Experi	mental	Pairs of C	Theor	eoretical	
Hydroca	rbon	8	Specific refrac-	Specific dis- persive energy	atoms doubly linked	Specific refrac- tive energy	Specific dis- persive energy	
Cymhydrene			0.543	0.0246	none	0.543	0.0243	
Menthene .			0.548	0.0313	one	0.547	0.0298	
The terpenes			0.537	0.0295	do.	0.537	0.0296	
Terebenthene			0.537	0.0294	do.	,,	,,	
Camphene .			0.528	0.0269	do.	,,	,,	
The cedrenes			0.538	0.0296	do.	,,	,,	
The citrenes			0.551	0.0334	two	0.553	0.0354	
Isoterebenthene	٠.		0.552	0.0337	do.	,,	,,	
Caoutchene			0.554	0.0366	do.	,,	,,	
Cymene .			0 ·560	0.0406	three	0.558	0.0413	
Isoprene .			0.592	0.0470	four	0.585	0.0472	

The advance in these experimental figures with the double linking is evident, though it is obscured in the first two cases by the larger amount of hydrogen contained in cymhydrene and menthene. The influence of the double linking upon the specific dispersive energy is apparent at a glance, and renders this property more valuable than the refractive energy in determining the rational constitution; though, of course, it is in the agreement of the two that the most conclusive evidence must be sought.

Kanonnikoff (Bl. [2] 36, 557) had already pointed out the differences between the calculated and observed refraction equivalents for the tollowing substances:—

Substance	R_{∞}	Calcu- lated	Differ- ence
Peppermint camphor . Cajeputene hydrate .	77·6 74·8	77·1 74·5	+ 0.3
Terebenthene hydrate, lævogyrate . French terebenthene,	77.1	74.5	+ 2.6
lævogyrate Hydrocarbon from	71.7	69.2	+ 2.5
oleum camillæ Hydrocarbon from oleum thymiani .	71·9 71·4	69·2 69·2	+2.7
Isoterebenthene	73·6 74·8	69·2 69·4	+4.4
Cymene from camphor.	72.0	66.6	+5.4

K. has drawn the conclusion that the first two contain no double bond, that the two terebenthenes and the two hydrocarbons contain one double bond, that isoterebenthene contains two, and that carvol and cymene contain three, double bonds.

Carbon. Schrauf (W. 22, 424) made observations on the refraction of a Brazilian diamond of the first water, the S.G. of which was 3.516, with the following results:—

observations on the refraction and dispersion of solid Se. By pressing a small quantity of melted Se between two glass plates, he prepared a thin film of the substance, which exhibited Newton's interference bands. From these he was able to calculate the index of refraction for the solar lines A, a, B and C, with an estimated error not exceeding 0.003; and for D, with some difficulty and uncertainty, to the second place of decimals only. The values obtained were:— $\mu_{\rm A}$ 2.654, $\mu_{\rm a}$ 2.692, $\mu_{\rm B}$ 2.730, $\mu_{\rm C}$ 2.787, $\mu_{\rm D}$ 2.98. From these S. calculated the dispersion as compared with water, calcspar, and CS₂, the figures in each case being reduced to the standard of $\mu_{\rm O}$ — $\mu_{\rm A}$ = 100:—

	$\mu_a - \mu_A$	$\mu_B - \mu_a$	μ_{C} — μ_{B}	$\mu_{D}-\mu_{C}$
Water Calcspar Disulphide of	36	31	83	88
	36	30	84	90
carbon Selenion	34	31	35	99
	29	30	42	1 4 6

Comparing Se with CS₂, he found that while in the case of CS₂ $\mu_{\rm D}$ — $\mu_{\rm c}$ differed very slightly from $\mu_{\rm c}$ — $\mu_{\rm A}$, in the case of Se $\frac{\mu_{\rm D}$ — $\mu_{\rm c}}{\mu_{\rm c}}$ = 1·46.

In testing Cauchy's and Christofie's formulæ for the index of the ray of infinite wave-length by the above observations, the former gave 2.733 and the latter 2.433. It will be observed that Cauchy's co-efficient of refraction is somewhat higher than the observed value for the solar line B, which is 2.730, the co-efficient of dispersion in this particular case being a minus quantity. Phosphorus. Gladstone and Dale $(P.\ M.$

Phosphorus. Gladstone and Dale (P.M. 1859) measured the index of refraction of both solid and liquid P, an element remarkable not only for its very high refractive power, but for an unprecedented amount of dispersion. For solid P at 25° they obtained for u_A 2·1059, for u_D 2·1442, and for the extreme limit visible in the violet

portion of the spectrum 2.3097. Assuming the latter to be equivalent to $\mu_{\rm H}$, the length of the spectrum $\mu_{\rm H} - \mu_{\rm A} = 0.2038$. For liquid P at 35° they obtained the following:—

μ_Δ	μ_{D}	$\mu_{\mathbf{F}}$	μ_{G}	End of violet
2.0389	2.0746	2.1201	2.1710	2.2267

The change from the solid to the liquid state is attended with a considerable diminution of density, the ratio between the density and the refraction being about the same in the two conditions. The R_A of P is taken at 18·3 from the above observations on liquid P, but Gladstone points out (C. J. 23, 101) that when combined with O it seems to have a much smaller refraction, HPO₃ giving a less value (18·0) than the P it contains, while H₃PO₄ also has a refraction-equivalent of only 23·6.

Sulphur. It was pointed out by Gladstone in 1869 (T. 159, 9) that while S, whether solid or liquid, has a refraction-equivalent of 16·0 or 16·3 for the solar line A, the equivalent for S deduced from CS_2 was 15·85; that the difference between KCN and KCNS gave a value of 16·24, and that S_2Cl_2 gave a similar figure. On the other hand, however, it was evident that in the two gases $H_2S = 14\cdot28$, and $SO_2 = 14\cdot91$, or in liquefied $SO_2 = 14\cdot59$, the refraction-equivalent of S cannot amount to 16·0, nor yet in H_2SO_4 , which has a refraction-equivalent of only 21·9.

Wiedemann and Nasini have subsequently given considerable attention to the values of S in combination. Wiedemann (W. 17,577) investigated the sulphur substitution products of the carbonic esters, and obtained the following results:—

influences, in a very distinct manner, the value of the index of refraction. From the refraction-equivalents given above W. obtains the value 14.04 for S' in the compounds II. and III.; and for S'' 15.20, 16.31, and 17.45 respectively. Using Lorenz's formula he gets the value of 7.94 for S', and 9.09, 9.44, and 9.33 for S'. Whichever formula be adopted, the figures show that the atomic refraction of S, like that of O, is greater when S is doubly linked to an atom of carbon. It should be observed that the above results are based upon the recognised values of C, H, O', and O'', which give

$${\rm CO}_{{\rm OC_2H_5}}^{{\rm OC_2H_5}} = {\rm R}_{\infty}$$
 45.91, and ${\rm R}_{\infty}$ 27.76;

and also that W. takes as the value of S' the mean derived from the observations II. and III. If Wiedemann had adopted a direct comparison of the actual observations, the values of S" would approximate more closely according to the old formula, and less closely according to Lorenz's formula.

Nasini (G. 13, 296; Lincei, Rc. 1, 1) has determined the value for S in various compounds. From sundry organic sulphides and mercaptans he obtained the atomic refraction of S corresponding to alcoholic oxygen. From CS₂, and from the compound CS(OC₂H_b)₂, he deduced the refraction of S corresponding to aldehydic oxygen:—

Sulphur	Ra	R_{∞}	Ra	R∞
S' (singly linked) . S" (doubly linked) .	14·10 16·05			7:65 8:84

		d	μ _{Li}	μ _{Na}	μ	R_{∞}	R _∞
I.	COOC ₂ H ₃	0.975	1.3837	1.3858	1.3876	45.66	27.86
II.	COSC,H,	1.0285	1.4479	1.4515	1.4544	56-95	84.11
III.	COSC,H,	1.085	1.5168	1.5237	1.5287	69 ·1 5	40.57
IV.	CSOC,H,	1.0325 ?	1.4563	1.4601	1.4632	57.82	84.56
v .	CSOC.H. CSOC.H. CS.C.H.	1.085	1.5304	1.5370	1.5431	70.26	41.23
VI.	CSSC ₂ H ₃	1.152	1.6105	1.6210	-	82.73	47.56

From these figures it will be seen (1) that the index of refraction rises whenever an atom of O is replaced by an atom of S, and that so much the more, the more S there is already in the compound; (2) that the refractive indices of the combinations which contain the radicle CS are greater than those of their analogues containing CO; (3) that the refractive index likewise increases if, besides the radicle, first one and then a second atom of S replaces O, and in the latter case more than in the former; (4) the isomeric bodies II. and IV., as well as III. and V., have quite different indices of refraction, and in both cases the compound that has the S in the radicle has the higher index. This indicates that the position of the S in the compound

But when examining other compounds, such as H_2SO_4 , SO_4 , and SO_2 , N. found that whatever hypothesis he adopted as to their constitution—or, in other terms, whatever value he took for the atomic refraction of O—he obtained values for S, not very discordant among themselves, but totally different from those given above. Treating the sulphur as divalent, R_∞ varied from 8·10 to 9·01; as tetravalent, from 6·94 to 8·43; as hexavalent, from 6·63 to 7·85. Calculating by Lorenz's formula, R_∞ ranged from 5·24 to 6·27 in the first case, from 4·51 to 4·91 in the cond, and from 3·13 to 3·79 in the third. Some observations recorded by Kanonnikoff (J.~R.~15,~434) substantially support these figures.

Nasini and Scala (*Lincei*, Rc. 1886. 617, 623) record observations on the sulphocyanides, which show that the mustard-oil compounds have refractive and dispersive powers in excess of those of their isomerides, while phenylic isosulphocyanide has a specific refraction and dispersion far above the calculated values-exceeding, in fact, oil of cassia or CS2. In this combination the union of the group S=C=N, endowed with a high refractive power, with the phenylic group, also highly refractive, appears to produce a heightening of the refrangibility, a phenomenon which N. regards as analogous to the increase in refraction which he observed when in the benzene group an unsaturated side chain is joined to an unsaturated carbon atom. In compounds containing four atoms of S there is an excess of refraction above what is estimated from the value of S in CS2, the monothicarbonates of Et and Pr giving a value of 17.4 and 17.5 for the refraction-equivalent of sulphur.

Nasini a. Costa (Ist. Ch. Rome, 1891) made

lished (Ar. N. 3) a long series of very careful determinations of the indices of refraction of mixtures of H2SO, and H2O, in various proportions for all the principal lines of the solar spectrum from A to H inclusive. These show a progressive increase in refraction and dispersion with every addition of H2SO4, until a maximum is reached at a point corresponding, apparently, with the monohydrate of the acid, after passing which the refraction and dispersion suddenly decline, while the density of the mixture continues to increase. Out of the seventeen series of determinations at different concentrations, the seven most conc. are given below, the central one of the series representing the nearest approach to the maximum. The indices are all reduced to a temperature of 18.3°. The specific gravity given in the line below is for 0°, water at the same temperature being taken as 1.0. In the cases of the two G's and the two H's, the higher one in each corresponds with the lines so designated by Fraunhofer.

	•	•					
μ	63·69 p.c. SO _a	71.97 p.c. SO.	81.41 p.c. SO _s	85·93 p.c. SO _s	88-97 p.c. SO ₂	91.43 p.c. SO _s	94·72 p.c. SO,
A	1.40819	1.41930	1.43049	1.43279	1.43151	1.42918	1.42683
a	1.40928	1.42042	1.43168	1.43385	1.43270	1.43035	1.42781
В	1.41016	1.42133	1.43263	1.43476	1.43357	1.43114	1.42868
C	1.41112	1.42227	1.43360	1.43579	1.43444	1.43198	1.42944
D	1.41342	1.42466	1.43596	1.43807	1.43669	1.43426	1.43163
\mathbf{E}	1.41618	1.42740	1.43877	1.44081	1.43944	1.43701	1.43431
ь	1.41669	1.42793	1.43928	1.44130	1.43996	1.43749	1.43484
F	1.41838	1.42967	1.44103	1.44311	1.44168	1.43922	1.43649
G	1.42058	1.43186	1.44327	1.44534	1.44392	1.44144	1.43869
G	1.42233	1.43364	1.44507	1.44706	1.44569	1.44316	1.44037
H	1.42421	1.43561	1.44703	1.44901	1.44759	1.44512	1.44229
\mathbf{H}	1.42567	1.43696	1.44841	1.45040	1.44883	1.44640	1.44347
s.G.	1.55438	1.64925	1.76066	1.80676	1.83123	1.84485	1.84994

further studies of S and its compounds. Thev conclude that the oxygen compounds, which are generally distinguished by a low refractive and dispersive power as compared with those not oxygenated, may be divided into two principal groups: those which may be regarded as derived from SO, have the higher, and those derived from SO, have the lower, refractive and dispersive powers. Kanonnikoff (S. Ph. Ch. Russ. 16, 129) assumed that the refraction of S varies with the change of valency, and assigned the following values for Ra:-

S divalent 14.10

S tetravalent 8.72 (in sulphones)

S hexavalent 4.85 (in H₂SO₄)

but this is not accepted by N. and C. They find that the accumulation of S in the molecule is almost always accompanied by an increase in the refractive and dispersive power. In this respect, though not in others, S appears to have much analogy with carbon. Throughout this

investigation N. and C. tested the dispersion formula of Ketteler, $\frac{\mu_a^2-1}{\mu_b^2-1}$, in which a and b represent rays of different wave-length, and found that it gave very uniform results; but from a physico-chemical point of view they do not regard this fact as an argument in favour of the formula.

Sulphuric acid. Van der Willigen pub-

Van der Willigen pointed out that the formula $(100-p)\frac{n^2-1}{d}+p\frac{n'^2-1}{d'}=100\frac{n''^2-1}{d''}$ will not apply to mixtures of H2SO, and water, where

100-p and p are the respective weights of the two mixed liquids, n, n', and n'' the indices of refraction, and d, d', d'' the densities of the two liquids and the mixture respectively. Dale and Gladstone had formerly stated that the same was

the case in respect of their formula.

Nickel and iron carbonyls. Ludwig Mond has prepared Ni(CO), and Fe(CO), both liquids of a highly-refractive character. former has been examined optically by Nasini (Lincei, Rc. 7, 411), and the latter by Gladstone. Ra for the nickel compound was found to be as high as 57.7, which, deducting 8.4 for the value of each CO, gives Ra = 24.1 for the metal itself, instead of about 10.0, as determined from the chloride, sulphate, and nitrate. This would seem to confirm the hypothesis that the refractive power varies with change of valency, increasing with the increased power of combination. In this compound the Ni is regarded by Mond a. Nasini as octovalent, and as justifying Mendeléeff in placing it in Group VIII. of his system of classification. Gladstone, on the other hand, is disposed to think that the metal in both compounds is, as usual, divalent, and that it is the CO which has a greatly increased value. He obtained for Fe(CO)_s, $R_{\rm A} = 67.33$, $R_{\rm D} = 69.40$. These figures show not only a very high refraction, but also very great dispersive power.

Silveriodide, bromide, and chloride. Wernicke (A. 142, 560) measured the re-fraction and dispersion of light by these salts in two different ways: (1) by examining the light reflected from glass plates covered with a layer of these salts, by means of a spectroscope, and noting the position of the dark interferencebands; (2) by direct measurements of small prisms made of the fused salts. The portion of the spectrum observed ranged from the solar line C to G. The dispersion $\mu_{G}-\mu_{D}$ was about twenty-seven times that of water in the case of the iodide; that of the other two salts is much less, though considerably greater than that of flint glass. AgI seems capable of existing in two conditions: at temperatures above 138° it gives a short spectrum limited to the red, yellow, and green, while below that temperature it gives a spectrum twice as long, showing all the colours. W. believes that AgI is partly dissociated at the higher temperatures.

Bromine. Weegmann (Diss. Bonn, 1888) determined the refraction-equivalent of Br by Lorenz's, as well as by the older, formula, for both the hydrogen lines, a, and Cauchy's limit, from a series of ethylene and acetylene compounds, adopting Brühl's values for C, H, and the ethylene linking. The following are his figures for Br compared with those arrived at by Brühl:—

Bromine	R_{∞}	R_{α}	\mathbf{R}_{∞}	Ra
Weegmann .	14·78	15·38	8·56	8·83
Brühl	14·81	15·39	8·70	8·95

Fluorine. The writer of this article has collated the observations made by various observers on bodies containing F in order to arrive at the refraction-equivalent of this element (P. M. 1885. 481). Those of Brewster on cryolite, of Brewster, Wollaston, Fizeau, Stefan, and Kohlrausch on fluorspar, and of Gladstone on aqueous solutions of KF, give a refractionequivalent for F varying from 0.29 to 0.32. Those of Topsöe and Christiansen on crystalline silicofluorides of Cu, Ni, Zn, Mg, and Mn, of the formula RSiF, 6H₂O, give a refraction-equivalent for F varying from 0.62 to 0.84; while (NH₄)₂SiF₆ gives 0.59 for F. The latter figures must be accepted with caution, as a very slight difference in the values ascribed to the other elements will account for the whole of the discrepancy. It is evident, however, that in any case the refraction-equivalent is of exceedingly small amount, and that the specific refraction, even if the highest value be taken, can scarcely be the half of that of any other element known.

J. H. and G. Gladstone (P. M. 1891. 1) have investigated the optical properties of fluorbenzene and allied compounds. The molecular refraction for C₈H₅F, for six lines of the solar spectrum, is given in the following table; from this is deducted the value for benzene less one atom of H, leaving a small balance as the atomic refraction of fluorine.

These figures are remarkable as giving a negative value for the dispersion R_B-R_A, namely the

	R	Ro	$R_{\mathbf{D}}$	R	$R_{\boldsymbol{G}}$	$R_{\mathbf{H}}$
CaHsF CaHs	42·92 42·29	43·33 42·70	43·70 43·17	44·68 44·20	45·61 45·17	46.40 46.05
r	0.63	0.63	0.53	0.48	0.44	0.35

quantity, -0.28. According to Lorenz's formula $\mathbf{R'_A} = 0.92$ and $\mathbf{R'_H} = 0.84$, showing in like manner a negative dispersion. The observations of Baille, Stefan, and Sarasin on fluorspar, and those of the authors on solutions of KF and NH,F, all distinctly confirm the foregoing evidence as to the anomalous dispersion; while a comparison of the refraction of the three lines C, D, and F of the solid silicofluorides measured by Topsüe and Christiansen, and the author's observations on aqueous solutions of $\mathbf{H_zSiF_{e}}$, seem to point in the same direction.

Liquefied gases. Bleekrode (Pr. 37, 339) adopted a special apparatus for measuring the indices of refraction of liquefied gases, consisting of a modification of a Faraday tube, forming a vessel having two parallel side plates, free from leakage even at high pressures. By observing, through a microscope provided with a graduated scale and a micrometer screw, test objects, such as lepidoptera scales, attached to the inner surface of the glass plates of the cell, the variation in the focus between the direct observation and that through the cell and liquid was found, and, with a correction for the effect of the glass, he obtained the data from which to calculate the index of refraction according to

the equation $\frac{D}{D-d} = \mu$, in which D represents the thickness of the layer of liquid and d the apparent displacement of the test object. The observations were taken both with a sodium flame and with daylight, and the calculated probable error of the result amounted to \pm 0.003. The first table on p. 238 gives the results of B.'s observations compared with those of Dulong, Ketteler, Croullebois, and Lorenz on the same substances in the gaseous state; the specific refraction is calculated by both the formulæ in common use.

B. drew attention to the remarkably low S.G. of ethylene, as the lightest fluid substance known; his determinations gave 0.335 at 8°, 0.361 at 6°, and 0.386 at 3°. B. also gave determinations of the indices of refraction of the following substances taken by the same process; and commented on the high indices of the organo-metallic compounds as inconsistent with the opinion expressed by early observers that a high index of refraction is indicative of a body being combustible.

Substance			Temp.	μ_{D}
Hydrobromic acid Do. Hydriodic acid Bromine Methylamine Dimethylamine Trimethylamine Zinc ethyl Zinc methyl			10° 15° 16·5° 13° 17·5° 17° 16° 12·5° 14°	1·325 1·330 1·466 1·571 1·342 1·350 1·353 1·485
Aluminium ethyl Aluminium methyl	:	:	6.5° 12°	1·480 1·432

Substance			Liquid		$\frac{\mu-1}{d}$		$\frac{\mu^2-1}{\mu^2+2}\times\frac{1}{d}$		
				s.G.	$\mu_{_{\mathrm{D}}}$	Liquid	Gas	Liquid	Gas
Sulphurous anhydride . Cyanogen Hydrocyanic acid Nitrous oxide Carbonic anhydride . Hydrochloric acid . Chlorine Ammonia Ethylene Hydrogen phosphide . Hydrogen sulphide .		•	:	1·359 0·866 0·697 0·870 0·863 0·854 1·33 0·616 0·361 0·622	1·351 1·327 1·264 1·204 1·196 1·257 1·367 1·325 1·180 1·323 1·390	0·252 0·378 0·379 0·235 0·227 0·300 0·270 0·528 0·498 0·519 0·429	0·236 0·350 0·368 0·255 0·221 0·277 0·240 0·490 0·526 0·500 0·413	0·153 0·234 0·238 0·150 0·145 0·190 0·169 0·327 0·321 0·322 0·262	0·157 0·233 0·246 0·170 0·147 0·185 0·160 0·327 0·350 0·333 0·275

Liveing and Dewar (P. M. 1892. 205) have determined the refractions of three liquefied gases, at their respective boiling-points, by direct observation through a hollow prism. They obtained the following results for the sodium line D:—

	μ_{D}	R _D	R _D
Oxygen Nitrous oxide . Ethylene	1·2236	8·182	2·024
	1·3345	11·587	7·163
	1·3632	17·556	10·752

Alums. The indices of refraction of some of the alums have been measured by several observers, amongst them Grailich, Fock, Kohlrausch, Stefan, Topsöe, and Christiansen. most complete series of observations on inorganic sulphuric alums is that of Soret (Ar. Sc. Genève, 12, 553; 14, 96; 20, 517), including those containing Rb, Cs, In and Ga. His mode of determination was by total reflection, which he considered the best adapted for the purpose; he gave indices which he calculated to be correct to two units in the fourth place of decimals. His observations were given for eight rays of the solar spectrum, viz. the lines a, B, C, D, E, b, F, and G. He remarked on the very low value of the index of refraction of the Na-Al alum. He gave the following comparative table for μ_D , showing that the different series of alums vary in general with much regularity. The horizontal lines contain the same metal, and the vertical lines the same metal or alkaline radicle.

	NH.	K	Rb	Os	Tì
Aluminium Indium Gallium Chromium. Iron	1:4594 1:4664 1:4688 1:4842 1:4848	1:4564 1:4653 1:4814 1:4817	1.4566 1.4638 1.4658 1.4815 1.4823	1.4586 1.4652 1.4649 1.4810 1.4838	1·4975 1·5066 1·5228 1·5236

Calculating for each salt the difference between its μ_D and that of the ammoniacal alum on the same horizontal line, Soret obtained:—

	NH ₄ -K	NH4-Rb	NHCs	NHTI
Aluminium Indium Gallium Chromium Iron	+0.0080 0.0080 0.0028 0.0081	+0.0028 0.0026 0.0025 0.0027 0.0025	+0.0008 0.0012 0.0034 0.0082 0.0010	-0.0381 0.0383 0.0386 0.0388

These figures show a close accord, except in the case of Cs, in which the Ga and Cr compounds

appear to be too low. Soret did not consider that the impurity in the specimens, if any existed, could be due to the Cs.

Gladstone (P. M. 1885. 162) worked out the refraction-equivalents from the data given in the first named of Soret's papers, and also from those by Topsöe and Christiansen, in order to test his former conclusions that a salt has the same specific refraction whether it is in a solid form or in solution, and that the refraction-equivalent of a compound body is the sum of the refraction-equivalents of its constituents. As to the first point, in the case of the two alums given below he found the specific refractions, after deducting the amount due to the solvent, to be in substantial accord:—

Ammonium-aluminium alum 0.2780 Crystallised O.2784 Sodium-aluminium alum 0.2613 0.2604

To test the second point, the alums were regarded as compounds of the sulphates of two metals of different kinds with twenty-four molecules of water. Taking the R_A of water in the uncombined state at 5·926—and taking the observations of Kanonnikoff, Topsöe, and Christiansen, as well as his own, of the several sulphates—Gladstone arrived at the results given in the following table. The other observers did not take the A line, but their observations have been reduced to A for purposes of comparison. The agreement between the calculated and observed refraction-equivalents is as near as might be expected, except in the case of Cs.

		R₄ ob and re	served duced
Substance	R _A calcu- lated	Soret	Topsöe and Christiansen
Ammonium-aluminium alum Sodium " " Methylamine " " Potassium " " Rubidium " " Ammonium-chromium alum Potassium " " Rubidium " " Cæsium " " Ammonium-iron alum . Potassium " " Ammonium-iron alum . Potassium " " Rubidium " "	252·0 239·3 267·2 245·1 254·0 268·0 254·0 257·1 266·0 280·0 270·6 263·7 272·6 286·6	252·2 238·5 267·7 246·8 253·7 262·3 265·3 266·7 275·5 269·1 265·0 276·0	268·6 261·4

Gladstone also sought, by means of Soret's observations, to determine the refraction-equivalents of In and Ga, and (B. A. 1892) he gave the following constants:—

				At. W.	RA
Indium Gallium	:	•	:	113·4 69·9	13·7 11·6

These figures, however, can only be looked upon as approximate, as all experimental errors are accumulated on these residual numbers.

Perrot (Ar. Sc. Genève, 1880) has made a series of observations on the refraction and dispersion of an isomorphous series of biaxial crystals—the double sulphates of Zn with K2, Rb₂, (NH₄)₂, Cs₂, K₂, and of Mg with Rb₂, and in each case 6H₂O. He has followed the plan adopted by Soret in his observations on the alums. The results conform satisfactorily to the values computed on the same principle as that adopted by Gladstone with respect to the alums, except in the case of cæsium and rubidium, in both of which, especially the former, the observed values are lower than the theoretical. The mean ray adopted for this calculation is, however, the arithmetical mean of the three indices observed for each of the lines of the spectrum, and has, therefore, no real physical significance. A glance at the tables of observations will show at once what a wide scope for error there is in arriving at a mean value by this process.

Analogy between refraction and dispersion and magnetic rotation. Gladstone and Perkin have recently observed an analogy between the magnetic rotation and the refraction and dispersion of light. They have tested this in the case of organic compounds containing nitrogen (C. J. 55, 750), with the result that they have found a correspondence that points to some connection between the rotation of the polarised ray under magnetic influence and the retardation of the rays of light in passing through a material substance. These three properties are really comparable with one another, for in each case the observed value is divided by the density of the substance, and it is determined, not for equal weights, but for an equal number of molecules. It has been found that each property is determined in the first instance by the atomic composition of the substance, so that it may be laid down as a primary law that the molecular magnetic rotation, like the molecular refraction or dispersion, of a compound is the sum of the molecular magnetic rotation, refraction, or dispersion of its constituents. The values are, however, subject to large modifications dependent upon differences in the structures of the compounds. Thus a change of valency is attended by a marked change of value in these optical properties, and they are apparently affected by some circumstances which are not as yet recognised in structural formulæ.

The following values have been deduced by Perkin for the molecular magnetic rotations of certain elements in the paraffin group of carbon compounds; and against them are placed the recognised values for their molecular refraction and dispersion:—

	Molecular mag- netic rotation	$\mathbf{R}_{\mathbf{A}}$	R _{II-A}
CH ₂	1·023	7·6	0·34
	0·515	5·0	0·26
	0·254	1·3	0·04
	0·194	2·8	0·10
O, aldehydic. Cl . Br . I N, in amines	0·261	3:4	0·18
	1·733	9:9	0·50
	3·562	15:3	1·22
	7·757	24:5	2·62
	0·717	5:1	0·38

These figures show a certain analogy; and when modifications are introduced by changes in the mode of combination, the change that occurs in the one property is noticeable also in the other two, and that in the same direction, though not necessarily to the same extent. Thus there are in all cases two different values for oxygen in alcohol and aldehydes, and a very great increase of the values in the case of unsaturated carbon compounds. The values also of CH, in the first and second members of homologous series (such as the alcohols, paraffinoid acids, &c.) are different from its value in the higher members of the same series, and always in the same direction.

P. also applied the same inquiry to solutions of HCl, HBr, and HI, which when dissolved in water were known to give higher values for refraction and dispersion than the normal. The same was found to be the case in the magnetic rotations. When HCl was dissolved in isoamyloxide, however, all three values were found closely approximating to the normal, as will be seen in the following table:—

Hydrochloric acid	Molecular magnetic rotation	R_	R _{H−≜}
In water , isoamyloxide .	4·412 2·238	14·45 11·36	1·12 0·51
By calculation for free acid	2.187	11.20	0.54

It is, therefore, laid down as generally, if not always, true, that where there is a departure from the normal values in regard to one or other of these properties, it is to be found in the other two. The different properties are evidently similarly affected by change in chemical constitution. The investigation seems to indicate another close relationship between electro-magnetism and the velocity of light. G. G.

OPTICAL METHODS.—Section 2: SPEC-TROSCOPIC METHODS.

Historical notes.—Spectrum analysis is an optical method of making a diagnosis of the chemical composition, and in certain cases the chemical constitution, of either (a) a self-luminous body, or (b) an absorbing medium, whether self-luminous or not. Newton first discovered that light could be analysed when passed through a prism, and, in 1752, Thomas Melvill made an optical analysis of alcohol flames coloured with

salt and with nitre. Wollaston, in 1802, observed with the prism certain obscure bands in the solar light which had traversed a narrow slit in his window shutter. Fraunhofer, in 1814-15, was the first to employ a very narrow and accuratelyformed aperture with a prism, telescope, and divided circles, and he not only observed the dark lines in the solar spectrum, but measured their indices of refraction or refrangibilities. He examined, subsequently, the spectra of the stars, of electric light, and of the exterior envelope of the flame of a wax candle. The latter spectrum he discovered to consist of homogeneous yellow light, composed of two distinct rays very close together, and coincident with the two dark lines in the solar light which he had termed D. Brewster noted the presence of this yellow light in the combustion of many substances, but it was Fox Talbot who, in 1826, clearly proved that the source of this light was sodium. He likewise connected a red ray with the presence of potassium and an orange ray with strontium, and concluded that a glance at the prismatic spectrum of a flame may show it to contain substances which it would otherwise require a laborious chemical analysis to detect. Herschel had previously described the spectra of a few of the metals and of boracic acid. In 1834 Fox Talbot described the strontia and lithia spectra, and stated that optical analysis could distinguish the most minute portions of these two substances with as much, if not greater, certainty than was afforded by any other known method.

Wheatstone, nearly at the same time, published some experiments similar to those of Fox Talbot, and Swan first estimated the amount of sodium which can produce the D lines. At the Manchester meeting of the British Association in 1842, Brewster read a paper On Luminous Lines in Certain Flames corresponding to the defective Lines in the Sun's Light. He discovered the dark lines of absorption in nitrogen peroxide gas when white light is directed through it, and he observed and mapped a number of atmospheric lines in the solar spectrum. In 1849 Foucault obtained a reversal of the D lines in the electric arc. Stokes, about 1850, gave in his lectures the true explanation of the coincidence of sodium lines with the dark lines D. Angström in 1853 stated that an incandescent gas emits luminous rays of the same refrangibility as those which it absorbs, and Balfour Stewart, 1858-9, based the following statement on experimental data: The radiating power of a body for any ray is equal to its absorbing power for the same ray.

In 1859, Kirchhoff and Bunsen gave a great impetus to spectrum analysis by publishing their researches on the constitution of the sun, and also by designing an extremely simple spectroscope and describing its use. Chemists were further made aware of the new power placed in their hands by Bunsen's discovery of two new metals, rubidium and cæsium, in 1860, by the discovery of thallium by Crookes and by Lamy in 1862, of indium by Reich and Richter in 1863, and of gallium by Lecocq de Boisbaudran in 1875.

Fundamental laws. Spectrum analysis primarily depends on the following two laws:—
1. Every solid and liquid when heated begins by emitting rays of longest wave-length, which are

the result of comparatively slow vibrations, but as the temperature rises the vibrations become more rapid, and shorter waves are emitted, so that at the highest temperatures the sum of the radiations is white light, consisting of waves of all possible lengths, the effect of which is a continuous spectrum. A mass of iron gradually raised to its melting-point would be an example of this. 2. Every substance vapourised by heat emits rays of definite wave-lengths, and these rays are characteristic of each substance, because it alone can emit them. In other words, every substance vapourised by heat has its own definite periods of vibration, by which it can be absolutely identified, because no other substance can vibrate in the same manner and with the same periods. All flame and spark spectra furnish examples. From this second law it follows that the spectrum of an element is a constant of very great importance. The determinations of the spectra and of the atomic weights present similar difficulties, which arise from the necessity for obtaining the elements or their compounds in the highest possible degree of purity, and also in many cases from the complexity of the spectra, and the careful observations which are necessary in identifying and measuring the wave-lengths of the rays.

Determinations of the absolute wave-lengths of the bright lines in any spectrum are now usually made either with a plane or a concave diffraction grating ruled on speculum metal. When a sufficient number of accurately-measured lines is known for any region of the spectrum, determinations of the wave-lengths of lines in new spectra may be made by observing with a prism-spectroscope, and by the use of an interpolation curve the wave-lengths corresponding to the positions of the lines measured may be found. From time to time the curve will require verifying and correcting.

There are two classes of spectra, emission and absorption spectra.

Emission spectra consist of: (1) Continuous spectra and (2) Interrupted spectra. 1. Continuous spectra are characteristic of most incandescent solids, liquids, and gases at great pressures, or of a cloud consisting of solid glowing particles. 2. Interrupted spectra consist of: (a) Bands of light or 'fluted band' spectra, characteristic of certain vapours, especially the vapours of compound substances rendered glowing at comparatively low temperatures; (b) bright line spectra, characteristic chiefly of transparent vapours at high temperatures or at low pressures; (c) reversed line spectra, which are dark lines seen on a brilliantly-illuminated ground. When a bright line spectrum is observed in front of a more brilliant continuous spectrum, the lines appear dark on a bright ground of colour, instead of bright and coloured on a dark ground, and they are then said to be reversed.

Absorption spectra. When rays capable of giving a continuous spectrum are made to traverse (a) transparent solids, (b) transparent liquids, (c) transparent gases at ordinary temperatures, it is frequently observed that a definite band of rays is absorbed at either end of the spectrum. This is called a continuous absorption. Quite as frequently a band of rays is cut out of the middle of the spectrum; this is called a.

selective absorption, and the region of the rays absorbed is said to be occupied by an absorption hand.

The production of emission spectra. Emission spectra may be produced in the following ways.

(a) By flames of low temperature, e.g. by a spirit lamp or Bunsen burner, acting on solids, or the sprsy of solutions. The flame may be fed with chlorine, bromine, or iodine, or with hydrochloric acid when salts of the halogens are examined. (b) By flames of high temperature: blow-pipe, and oxyhydrogen flame. (c) By sparks in vacuum tubes containing residual gases or vapours. (d) By sparks of low tension acting on solids or solutions. (e) By sparks of high tension. (f) By solids in the electric arc. (g) By electric discharge on solids in vacuo; these are phosphorescence spectra.

As a rule, line spectra are the spectra of elements, band spectra are the spectra of compounds.

Salts of the alkali metals in the Bunsen flame emit line spectra characteristic of the metals, while the alkaline earth salts emit lines and bands which are characteristic of the metallic oxides. On the other hand, the burning of hydrogen and of hydrogen compounds causes the emission of more than 160 lines in the ultraviolet region, which can only be assigned to the vapour of water at a high temperature. Carbon is an element which emits bands in the coloured region, in hydrocarbon flames. A list of compounds which emit characteristic banded flamespectra is given on p. 246. Erbia and erbium phosphate emit luminous bands when the solid substances are heated in a Bunsen or blow-pipe flame. Discharges in vacuum tubes sometimes emit a continuous spectrum; oxygen does so at the lowest temperature at which it is luminous. Most gases emit a banded spectrum with low tension sparks at ordinary pressures; the spectrum alters with diminution of pressure until finally nothing but a line spectrum is visible—for example, this is the case with air in the ultraviolet region. Sparks of low tension from solids, such as metallic electrodes, and from solutions, emit lines chiefly in the visible region, while those of high tension are generally richest in radiations of high refrangibility, chiefly in the ultra-violet region.

Low-tension sparks may be produced in two ways from solutions, first by wires of platinum moistened by the liquid, secondly by pieces of charcoal giving the flaming spark used by Bunsen. Many elements have been shown to emit a few bands or lines in the infra-red. The difference in action of high- and low- tension sparks is chiefly, if not entirely, due to temperature; the highest temperatures cause the emission of rays of the highest refrangibility. Solid elements, with few exceptions, yield spectra in the electric arc. The lines exhibited by some of the metals are exceedingly numerous. Reversed line spectra are common in the sun, and afford evidence of the presence of metals whose bright line spectra are coincident therewith. The number of bright lines seen reversed in the sun's spectrum indicates the strength of the evidence of the existence in the sun of certain elements.

As to the distribution of known kinds of | Vol. IV.

matter in space, the following list shows the elements observed in the sun with the numbers of their reversed lines seen in the solar spec trum: Fe 730, Ti 230, Ca 94, Mn 85, Ni 102, Co 91, Cr 71, Ba 26, Na 11, Mg 17, Cu 7, H 5, Pd 5, V 4, Mo 4, Sr 4, Pb 3, U 3, Al 18, Ce 2, Cd 2, Be 2, Sn 1.

Methods of spectrum analysis. For efficient work with the spectroscope an instrument is to be preferred which admits plenty of light, has a rigid collimator, and immovable prism. Compound prisms of Grubb's and of Thollon's construction are among the best. The arc should be divided on the vertical edge of the prismtable or on an edge inclined at an angle of 45°. A vernier travelling over the arc and a mounted lens, for convenience in reading the scale and vernier, should be fitted near the telescope. A beautiful instrument made by the Societé Génevoise pour la Construction des Instruments de Physique has a circle completely divided, and fittings for investigating ultra-violet spectra, either by photography or with a fluorescent eyepiece of Soret's design. It is provided with two verniers, and an automatic arrangement admitting of the prism being placed at the minimum angle of deviation for each ray under examination. This is necessary in order to secure the best definition. The instrument, of whatever form, should be capable of dividing and measuring the lines D' and D2 of sodium, the first, second, and third lines of the yellow band; and in the carbon spectrum the first, second, third, and fourth lines of the green band, and the first, second, and third lines of the blue band, which are seen in the green inner cone of a powerful Bunsen burner. An instrument with a photographic scale, if it is sufficiently delicate, which is seldom the case, is of great use. One of these, constructed by Duboscq, was employed by Lecocq de Boisbaudran. A lens of short focus should be used to concentrate the light, from whatever source, upon the slit of the instrument. A test tube about 12 mm. diam., supported vertically, if filled with water and placed close to the slit of the instrument, answers this purpose. The telescope should be capable of easy movement without vibration, and be supported for two-thirds of its length from the object glass; neither clamping nor the use of the focussing screw should be capable of displacing its position. The rays should pass straight down the collimator and completely fill the lens. Before commencing any serious work, the capabilities of the instrument and its use for obtaining measurements should be carefully studied.

Supports of platinum wire fused into glass tubes are used for examining concentrated solutions and fused beads of salts in ordinary flames. Bunsen burners with steatite tops are particularly useful. Blow-pipe flames with a vertical direction, urged by bellows worked by the foot, are useful for examining the alkaline earths. Hydrogen gas from a constant generator made of lead and of large size is better than coal-gas for all flame spectra. Whether coal-gas or hydrogen be used, it is advantageous in examining salts of the alkaline earths, particularly chlorides, to pass the gas through a litre of strongest hydrochloric acid into a Bunsen burner, which is made of hard glass tube tipped

with platinum. The flame is thus supplied with hydrochloric acid, and the chlorides are volatilised apparently without decomposition, so that they emit rays of great brilliancy which yield simple For high temperatures a very line spectra. small gas blowpipe may be supplied with oxy-In this case the supports of platinum would be fused, and it has therefore been found convenient to use a wire of iridium, twisted into a loop, the compound being placed in the loop. In this manner manganese dioxide yields the bands of the dioxide, the spectrum of which is observed in the Bessemer flame. Calcium fluoride and copper oxide readily yield characteristic spectra. The alkaline earths yield very brilliant spectra, which are due to the metallic oxide in each case. When volatile substances, such as salts of lithium and the other alkalis, are under observation, it frequently happens that a difficulty in measuring their lines occurs owing to the evanescent appearance of their spectra; more lasting spectra may be produced by converting the volatile salts into silicates or borates by heating in a platinum spoon, or even in a porcelain crucible, with crystals of boracic acid. Beads of the borates will continue to illumine a flame for a lengthened period; with lithium and thallium, measurements have been made for as long as twenty minutes without renewing the material. Should the alkalis be accompanied by lime, strontia, or baryta, the spectra of these earths are suppressed, but towards the end of the observation sometimes the green colour of boric oxide appears. Sparks, both of low and high tension, may be used with Delachanel and Mermet's fulgurator. This instrument has the advantage of using very little solution, and losing none. Some solutions, such as salts of yttria. do not yield their characteristic spectrum to lowtension sparks, and it is necessary to strengthen the current by interposing a Leyden jar in the circuit, or, what is better, a pane of glass coated on either side with 930 square centims. of tinfoil. The spark should be 3 to 6 mm. in length. The lines of platinum, hydrogen, sodium, and a few of chlorine are visible, but these are known and are measurable. When the solution becomes contaminated with platinum, it may be removed and purified to be used over again if necessary. For such work it is best to use a coil which can yield sparks 170-180 mm. long in air.

For observations on ultra-violet spectra the arrangements described by Soret and by Cornu consist of an optical train of quartz and a prism of 60°, which is composed of two halves each of 30°; one of the two lenses and one half of the prism are cut from a right-handed crystal, the other corresponding parts from a left-handed crystal, so that the double refraction of one is corrected by that of the other. One half of the prism may be fixed on the collimator close to the lens, the other in front of the object-glass. For ordinary observations a fluorescent eye-piece is necessary, but generally photography is more convenient. Cornu has used lenses achromatised with colourless fluorspar, but this is not necessary, because the peculiarity of the camera employed by Hartley enables whole spectra to be accurately focussed on one plate if only one prism of 60 degrees be used. The camera back is sloped at an angle of about 22° to the normal,

which brings one end of the plate nearest to the rays of shortest wave-length, and the locus of the foci of all the rays, from the least to the most refrangible, lies upon the plate. There is thus a saving of time in the observation of complicated spectra, with the additional advantage that angular measurements are abolished and linear measurements substituted. Moreover, an increased dispersion and separation of the rays is obtained. This arrangement is most suitable for use with two half-prisms of quartz of 30°. Six hundred lines in the spectrum of iron have thus been photographed with such excellent definition that a magnifying power of 25 diameters was used in their measurement. A quartz lens 75 mm. in diameter and of 75 mm. focus is placed in front of the slit. The camera back is so constructed that by a rack and pinion motion a succession of twelve or fifteen spectra may be taken on one plate. The lenses, which are not achromatised, are of 425 mm. focus for yellow light, and 50 mm. in diameter, the prisms being of corresponding dimensions.

Sparks of high tension obtained in the manner already described are necessary. They may be passed between metallic electrodes 3 to

6 mm. apart.

For photographing spectra yielded by solutions electrodes of graphite are used, or wicks of fine wire drawn from pure gold. The solution is placed in a small J tube; in the shorter limb the electrode is placed, and a platinum wire passes down the longer limb to make connection with the coil. By means of a pipette the tube is easily charged. The graphite electrodes are cut from good sound pieces of Ceylon or Siberian graphite, and are chisel-shaped, with grooves scored along them with the point of a pin. The opposite electrode may be of graphite or of any suitable metal. In order that the strongest discharges may pass from below upwards the lower should be the negative electrode. The admirable work of Lecocq de Boisbaudran on flame and spark spectra (Spectres Lumineux, 1874) not only gives the practical details, but maps of prismatic spectra, and the same applied to a scale of wave-lengths, as well as a complete description of the spectra and a description of each line. The publication proves to be all that a guide to the use of the spectroscope should be, but it deals only with the visible region. In photographing spark spectra the trouble of making drawings is unnecessary, since the photographs themselves are better than any reproduction; all that is required is a description and a map of the principal lines on the scale of wave-lengths. In photographing the spectra of solutions by means of graphite electrodes it is, of course, necessary to ascertain that the material does not yield any of the numerous lines in the spectrum of iron, generally speaking four lines of magnesium with wave-lengths 2801.1 to 2789.6 are visible. Although photography has been applied almost exclusively to the ultra-violet rays and to the infra-red, there is no reason why it should not be used for visible rays, since gelatin plates stained with cyanine, eosine, erythrosine, and some other dyes, render the plates sensitive to the orange, red, and yellow rays. Cyanine is a dye which is sensitive to most of the spectrum as far as the rays of very long wave-lengths.

On the sensitiveness of emission spectra.

The following table gives the sensitiveness of the emission spectra of various elements in fractions of a milligram. It will be observed that radiations in different regions, and excited by different means, greatly vary in emissive power even with the same element.

Visible flame spectra.

	Mgr.		
Ba	2,000	Kirchhoff	a. Bunsen
Са	1 50 000	**	••
Cs	25,000	**	**
-Cu	1 285	Simmler	
[n	<u>1</u> 2,000	Cappel	
Li	600,000	Kirchhoff	a. Bunsen
Mn	1 83	Simmler	
K	3,000	Kirchhoff	a. Bunsen
Rb	7,000	**	**
Na	14,000,000	**	**
\$r	3 0,000	**	27
Tl	5 0,000	Lamy	

Visible spark spectra (Cappel).

			•
	Mgr.		Mgr.
Ba	1 930,000	Li	1 40,000,000
Bi	70,000	Mg	1 500,000
•Cd	18,000	Mn	1 200,000
Ca	10,000,000	$\mathbf{H}\mathbf{g}$	10,000
-Cr	4,000,000	Ni	600
·Cs	4,000	K	1 400
-Cu	20,000	$\mathbf{R}\mathbf{b}$	1,000
€ Co	1 15,000	Sr	1 100,000,000
Au	1 4,000	Tì	8 0,000,000
In	90,000	Sn	17,000
Fe	$\frac{1}{26,000}$	Zn	1 600,0 0 0
Pb	1 20,000		

Photographed ultra-violet spark spectra (Hartley).

Mgr. 1 100,000,000

On ultra-violet spectra. The conditions necessary in photographing ultra-violet spectra, are: 1st, a sensitive salt exerting an equality of action throughout the whole range of the spectrum; 2nd, a perfectly diactinic vehicle to hold the salt; 3rd, a process of development which will not exert a preferential action on any portion of the spectrum. These conditions are fulfilled

by the use of silver bromide emulsion, the gelatin being of the purest description, and the image being developed by pyrogallol, hydroquinol, or hydroxylamine hydrochloride and caustic soda. The new developer eikonogen is suitable. Ferrous oxalate, as usually employed, is not to be recommended, because strong lines become developed much more strongly than weak lines. The exposure varies slightly with the sensitive nature of the plates, the width of the slit, the conducting power of the electrodes, the emissive power of the rays to be observed, and lastly with the excellence of the developer. Having once ascertained by comparative exposures how to obtain the best effect, there is very seldom any necessity for altering the exposure.

The following facts have been established regarding ultra-violet spark spectra: 1. When carbon or metallic electrodes, which emit short lines, are moistened or partially immersed in water, the short lines are lengthened. 2. The impurities in a metal, or the minute constituents of an alloy, generally appear as spectra composed of short lines. When both electrodes are of the same material, the impurities in minute proportion appear at the negative pole only, but when the proportion of the impurity or alloy is increased, the long lines are seen as short lines at both poles; a further increase in proportion brings them out as long lines. 3. Of the nonmetallic elements, boron, carbon, and silicon exhibit line spectra. The line spectra of the elements are seen in spectra of borates and silicates, and in boracic and hydrofluosilicic acids. 4. The spectra of carbon and silicon may nearly always be detected in photographs taken from electrodes of metallic iron. 5. Insoluble compounds which are not volatile, such as ferrous sulphide, ferric oxide, and ferrous phosphate, do not yield spark spectra. Insoluble, though somewhat volatile, compounds, such as thallium chloride, yield a feeble spectrum of metallic lines. 6. With the exception of boron and silicon, the non-metallic constituents of a salt do not affect the spark spectra of solutions. 7. The spectrum of a metallic chloride, nitrate, or sulphate yields spectral lines identical in number and position with the principal lines of the metal itself. Some of the short lines become long lines, but whether the spectra are produced by metallic electrodes or solutions their character is identical. 8. The effect of diluting solutions of metallic salts is first to weaken and attenuate the metallic lines, then, with a more extensive dilution, to shorten them, the length of the longest and strongest lines generally decreasing until they finally disappear. There are two lines in the spectrum of copper which disappear by attenuation only, and a similar pair of lines in the spectrum of silver behaves in like manner. Both pairs of lines have been observed as short lines in alloys containing very small quantities of these metals. 9. Accidental differences in the passage of the spark, or in the time of exposure of the photographic plate, when the normal period varies from half-a-minute to five minutes, do not cause sensible variations in spectra obtained from the same substances, under any ordinary circumstances. 10. Spectra of metallic electrodes and of solutions are liable to contain the lines of calcium, with wave-lengths

2967.3 and 3933, also occasionally those with wave-lengths 3736.5 and 3705.3. These calcium lines are due to dust when the electrodes are dry, and to impurity in the acids when solutions are prepared therewith and examined. 11. Five modified spectra have been observed in photographs taken from graphite electrodes, which apparently belong to no other element but carbon. The first and second were produced under identical circumstances, and were taken from dry points in air. The first contains 20 lines, the second only 12. The third was taken from wet electrodes in air, and consists of 20 lines, some of which are lengthened. The fourth was taken from dry electrodes in oxygen, and consists of 17 lines, some being much lengthened. The fifth was taken from dry electrodes in carbon dioxide, and consists of 13 lines, all much lengthened and strengthened (T. pt. i. 1884).

The examination of metals and alloys. When a metal is to be examined it may be filed or hammered into electrodes of a convenient shape, and held between screw clips or between the points of cheap drawing pens. According to circumstances, both electrodes need not be of the same metal, but one may be of gold, platinum, or a tin-cadmium or lead-cadmium alloy containing in either case 20 p.c. of cadmium. Such an alloy gives convenient reference lines. If it be required to bring out the lines of the constituent of an alloy present in the metal in minute proportion, or present as an impurity in the metal, it is best to partially immerse the lower electrode in water contained in a small glass J tube, as if a solution were being photographed. Under these circumstances the short lines become much lengthened and consequently more prominent. The best form for electrodes is flat with curved edges, or even somewhat rounded discs, such as blow-pipe globules flattened.

The identification of lines in photographed spectra. When the spectrum of an alloy, metallic precipitate, or solution has been photographed, it is seldom easy to determine what substances are present or absent, partly on account of the complexity of the spectra and partly from the absence of colour. It is necessary in such cases to have recourse to one or both of the following methods of identification. (a) Method of identifying the elements by lines photographed from a known specimen. A series of photographs taken from metallic electrodes and from solutions should always be kept for reference. A substance may then be readily identified even when the wave-lengths of its lines are unknown, by superposing one photograph on the other, film to film, since there are but very few cases of actual coincidences of lines of similar character belonging to different elements. If this examination does not account for all the lines, it is necessary to proceed according to the next method. (b) Method of identifying lines when comparison-spectra are not available. By chemical processes of separation it is easy to ascertain what metals may be looked for. The wave-lengths of the unrecognised lines may then be ascertained by measurements and reference to an interpolation curve. The measurements need not be made with a

micrometer unless great accuracy is required. It suffices generally to measure the spectra by means of an ivory scale with bevelled edges, the divisions on the scale being hundredths of an inch. With a strong magnifier we may read by judgment to tenths of a division or thousandths of an inch. The best form of magnifier is made like the compound eyepiece, of a microscope with cross hairs or fibres of cocoon silk in the field. The lens is placed very close to the surface of the scale and photograph, so that false readings caused by parallax are avoided. When all the lines in a photograph which have been recognised, or vice versa, have been pricked off, the scale is very carefully adjusted so that it gives correct readings with lines of known wavelengths at different parts of the spectrum, such for instance, with Nos. 9, 10, 11, 12, 17, and 25 of cadmium. In order to adjust the scale accurately, it is advisable to rule a line along the whole of the spectrum, which may be done when photographing by placing a wire at right angles to the slit, and one-third or one-fourth the length of the spark, above the lower electrode. edge of the scale is applied to this line, and held in position by spring clips or screws. The scale numbers of the unrecognised lines are then read off, and by reference to the interpolation curve their oscillation-frequencies or their wavelengths are obtained. From maps, or descriptions of spectra, accompanying a list of wavelengths, the corresponding lines are selected and identified.

Descriptions of the spectra of the elements. Emission spectra. The spectra of non-metallic elements, particularly those which are gaseous, have been found to vary in character with the conditions under which they have been obtained. Thus N, O, C, Br, I, S, Se, Te, and P yield band spectra or line spectra, according to variations in temperature or pressure. The chlorides of B and Si likewise give line spectra due to B and Si; under certain conditions they emit spectra of their own. The fluted spectrum of carbon has given rise to great discussion. Swan, and later Angström, attributed the bands to a hydrocarbon, so also did Morren, and Liveing a. Dewar, but they finally arrived at the conclusion first enunciated by Attfield, that the bands were caused by the element carbon. Van der Willingen, Plücker and Hittorf, Marshall Watts, Wüllner, Lockyer, and Fievez, have all adduced evidence of this. It appears, however, that a banded spectrum of cyanogen can be obtained which is distinct from that of carbon. Certain bands in the ultra-violet spark spectrum, which have been considered by Liveing and Dewar to be caused by cyanogen, have been observed by Hartley, and he is led to the conclusion that they are in reality carbon bands intensified by the presence of an atmosphere of nitrogen, but not cyanogen bands, for the reason that they cannot be obtained from various cyasagen compounds, such as soluble cyanides. They are always present when graphite electrodes are used in air, they are intensified when the electrodes are moistened, and they become particularly strong when concentrated solutions of chlorides, such as zinc chloride or calcium chloride, surround the electrodes; moreover, the strength of the bands increases with the strength of the solutions.

A review of the facts shows that large mole-

cules give banded spectra, as is evident from the spectra of compounds, and various other considerations lead to the conclusion that the elements which give such are capable of existence in different molecular conditions; the most complex molecules emitting bands of rays, the simpler molecules emitting line spectra.

Hydrogen. When sparks are passed through hydrogen, four very well known lines appear. Huggins has observed the same in some stars, together with a remarkable series of lines in the ultra-violet, and it has been surmised that the whole series, visible and invisible, belong to the one element. Cornu has recently proved this to be the case. Fifteen lines in the visible and ultra-violet constitute the only true line spec-

trum of hydrogen.

The alkali metals, Li, Na, K, Rb, Cs. These metals are distinguished by the fact that most of their salts are decomposed in the flame, and the metal set free, or the metallic oxide formed from it, is sufficient to colour the flame intensely. The spectra are very simple, and all the principal lines are in the visible region. ultra-violet spark spectra are insignificant.

The alkaline earth metals, Ca, Sr, Ba. The spectra of these elements are obtainable by flames. When the chlorides are fused and heated in a flame supplied with hydrochloric acid, lines due to the metallic chlorides are visible. Without hydrochloric acid, the spectrum is at first a simple line spectrum, but subsequently bands, shown by Mitscherlich to be due to the oxides, take their place. Feeble sparks give line spectra with solutions of salts. In the ultraviolet, a series of well-marked groups of doublets and triplets is seen.

The magnesium group, Be, Mg, Zn, Cd. The element beryllium stands at the head of two families, that of the alkaline earths and the magnesium family. It occupies a position similar to that of lithium with regard to the alkali metals and the copper and silver family. Its spectrum is similar in many respects to that of lithium, being very simple and composed of isolated lines. There are two in the blue and five in the ultra-violet not visible in the flame, but obtainable by the action of condensed sparks. Magnesium and the compounds of the other metals of this group do not easily yield flame spectra, but give characteristic groups of lines when solutions are submitted to feeble or condensed sparks. In the ultra-violet these consist of well-marked groups of isolated lines, pairs, triplets, and groups of four and of five lines.

The aluminium elements, B, Al, Ga, In, Tl. Boron, which stands at the head of this group, gives a flame-spectrum only of its oxide, but the boron compounds, such as borates, emit three lines in the ultra-violet due to this element. Aluminium gives no flame spectrum, but gallium, indium, and thallium emit rays in the violet and green. Sparks taken from solutions of indium and thallium yield characteristic spectra in the ultra-violet consisting of many isolated lines and pairs of lines.

The tin elements, C, Sn, Ge, Pb. Carbon presents a very simple spectrum when condensed sparks are used; most of the lines are in the ultra-violet. It is a remarkable fact that under the same conditions the number of lines some-

times varies, and no satisfactory reason for this can be assigned. Four spectra of graphite have been mapped from photographs described as follows: 1. Dry electrodes in air. 2. Wet electrodes in air. 3. Dry electrodes in oxygen. 4. Dry electrodes in carbon dioxide. The strength and length of the lines vary under certain circumstances; in 1 the lines are all short, in 2 some lines are lengthened, in 3 some lines are much lengthened, and in 4 all the lines are lengthened and strengthened.

The spark spectra of tin, germanium, and lead contain numerous lines which are not characterised by any evident special grouping. The ultra-violet spectrum of germanium has yet

to be photographed.

The group of rare earths, Ce, La, Di, Sm, Yt, Sc, Er, Tr. These elements give no flame spectra, and but weak spectra with feeble sparks. Ce, Di, Yt, Sm, Sc, Tr yield very complicated spectra to powerful sparks, the spectrum of cerium in the ultra-violet being remarkable for the immense crowd of lines. Yttrium and scandium chlorides give spectra composed of bands which are very characteristic in the visible region. Didymium, samarium, erbium, and thallium will be dealt with under the head of Absorption spectra. Lanthanum gives a large number of lines in the visible region, but very few in the ultra-violet.

The nitrogen elements, N, P, As, Sb, Bi, V. Nitrogen gives a large number of lines under the action of condensed sparks. The ultra-violet portion of the spectrum has been photographed. Nothing of the same kind is known of phosphorus. Phosphorus in a current of hydrogen gives a beautiful banded flame spectrum, especially when the flame is cooled. This reaction is exceedingly sensitive. The spectra of arsenic, antimony, and bismuth in the ultra-violet are very well marked, and they each contain a large number of lines, some of which are nebulous, others are mere dots, and there is a characteristic background of continuous rays. Vanadium yields spark spectra, with a large number of lines.

The chromium elements, Te, Cr, Mn. The line spectrum of tellurium obtained from the element is very complex, and by far the greater number of lines are in the ultra-violet; the character of the spectrum resembles those of arsenic, antimony, and bismuth. The chromium and manganese spectra are very complex and similar to those of the iron group.

The iron, nickel, and cobalt group. These metals present very complicated spectra when condensed sparks are employed in the visible, and more particularly the ultra-violet, regions. The grouping of the lines in the dif-

ferent spectra is similar.

The palladium, gold, and platinum These elements present complicated spectra, though less so than the foregoing. The strong lines of gold are few in number. These strong lines of gold are few in number. These metals do not excel in emissive power. Feeble sparks with them give feeble spectra. Iridium gives a spectrum only with condensed sparks, which in the ultra-violet consists of a great crowd of lines.

The spectra of many elements, such as tungsten and uranium, have yet to be examined thoroughly. It may here be remarked that elements usually associated by their natural occurrence, or by classification according to their chemical properties, exhibit spectra with characters in common.

A very important paper by Mitscherlich (A. 121, 459) first drew attention to the fact that a number of compounds emit distinct spectra, for the most part band spectra. A list of such compounds here follows, most of the spectra of which were originally described by Mitscherlich. Diacon also (Thèses de Physique et de Chemie, Montpelier, 1864), using a flame the interior of which was fed with chlorine, obtained distinct spectra of chlorides, such as those of the alkaline earths, gold, lead, iron, cobalt, and nickel; the alkalis gave no spectra excepting where they became converted into oxides or metal, as in the mantle of the flame.

Compounds emitting distinct spectra (Mitscherlich). 'Venetian-blind' or degraded band spectra. Flame. BaO; BiCl₃, BiBr₃, BiI₃; B₂O₃; CuCl₂; CuBr₂, Cu₂I₂, CuO; AuCl₂; Fe₂O₃; MgO; MnO₂; PbCl₂, PbO; MgCl₂.

Plain band spectra. BaF₂, BaCl₂, BaBr₂; BiI₃; CaF₂, CaO; CuF₂; SrO.

Line spectra, or lines with bands. BaCl2, BaBr₂, BaÎ₂; CaCl₂, CaBr₂, CaI₂; SrF₂, SrCl₂,

SrBr₂, SrI₂.

Other observers have discovered and described specific spectra emitted by the following compounds:—Al2O3; feeble sparks (Wüllner and others). NH3; flame (Dibbits). CO; spark (Plücker and others). CN; flame (Fox Talbot, Draper, Dibbits, and others). Er₂O₃ and ErPO₄; flame (Bunsen a. Bahr). MgO and hydride of Mg; flame and spark (Liveing a. Dewar). PH₃; SiCl4, flame cooled (Christofle a. Beilstein). SiBr, SiI, ; spark (Plücker, also Salet). H₂O; flame (Liveing a. Dewar; Huggins).

Harmonic relations between the lines of a spectrum. The self-luminous nature of a molecule is connected with the vibrations of those parts of the molecule which are called atoms. The number of vibrations is indicated by the inverse wave-lengths of the lines in the spectra. The spectrum of iron contains more than 1,200 distinct lines, and it follows, therefore, that the molecule or its atoms must simultaneously excite as many different vibrations. We are acquainted with similar relations in acoustics. For instance, it has been observed by Hipkins that the simplest vibration of a pianoforte string will cause it to emit its own proper note, and as many as twenty-four overtones or harmonics. Johnstone Stoney first discovered such harmonics in spectra. The visible spectrum of hydrogen contains the three lines Ha wavelength 6563.9, $H\beta = 4862.1$, $H\delta = 4102.4$, these are the 20th, 27th and 32nd overtones of a vibration, of which the wave-length = 0.013127714 mm. But the fourth line $H_{\gamma} = 4341$, and the lines in the ultra-violet do not fall in with this system. Soret has calculated the harmonics of the three groups of triplets in the magnesium spectrum to be the 20th, 27th, and 31st harmonics of the fundamental vibration. Similar groups occur in the cadmium spectrum, which are the 20th, 23rd, 27th, and 31st of a fundamental vibration. Schuster has observed several harmonics in the spectrum of strontium; also in the spectra of

bismuth, calcium, thallium, and gold. According to Mascart, sodium has five series of doublets which are the 132nd, 138th, 143rd, 158th, and 163rd harmonics. One of the most remarkable spectra is the absorption caused by chromyl chloride. Johnstone Stoney and Emerson Reynolds have shown that under great dispersion and lime-light illumination it gives a series of harmonics which are similar to those of a violin string, and of which the order lies between 628 and 733. Liveing and Dewar have observed that many ultra-violet spectra contain lines which are harmonically related, not so simply related perhaps as the harmonics of a uniform stretched string, but like the overtones of a string which is not of uniform thickness, or is loaded at different points, similarly related in origin though not exact harmonics. Lithium, potassium, calcium, magnesium, and zinc are such elements. The alkali metals have each one series in the visible spectrum, and another in the ultra-violet. It may happen in other cases that two or more such series overlap, and it may be very difficult to distinguish and separate them. There is a remarkable fact to be noted with regard to ultraviolet spectra, namely, that similar groups of lines, evidently harmonically related, are alternately sharp and nebulous in character.

Huggins discovered that the four hydrogen lines in the solar spectrum are part of a much longer series of lines which occur in the spectra of white stars. Cornu discovered the whole series in the spectrum of pure hydrogen, and Balmer showed that the whole series of lines can be expressed by the following formula:

$$N_{m-2} = k\left(1 - \frac{4}{m^2}\right)$$
, in which k is a constant

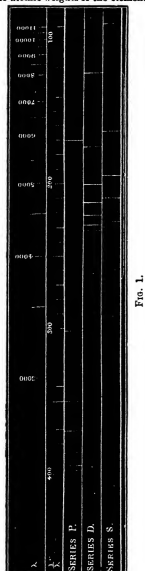
for the whole series, and by substituting the whole numbers 3, 4, 5, 6, &c., for m, we obtain values for N which are the oscillation-frequencies of the successive lines.

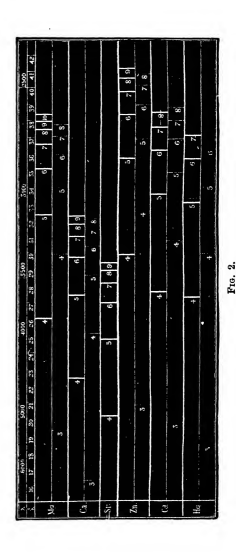
Relations between the spectra of different elements. Lecocq de Boisbaudran has observed such a relationship between the lines in the different spectra of the alkali metals and their atomic weights, that he has been able to calculate the atomic weight of gallium from its spectrum. It may be stated that the atoms of greater mass vibrate more slowly, and therefore emit rays of proportionally greater wave-lengths. Ciamician made an extended series of observations on the spectra of elements usually classed together on account of their chemical properties, and he described several of these spectra as being homologous, that is to say, similarly constituted. A. Grünwald observed mathematical relations between the spectrum of water vapour and the line spectra of H and O, which led him to predict the existence of many lines in the spectrum of water vapour which had not been photographed. By prolonged exposures, Liveing and Dewar obtained photographs of a number of lines approximating very closely to those calculated by Grünwald.

Hartley, in 1883, observed the law of homology in the visible and ultra-violet spectra of the magnesium, zinc, and cadmium group. Elements with properties in common exhibit spectra with similar groupings of lines, but the dispersion of the lines and the refrangibility of the strong lines

In each group vary with the atomic weights of the elements. In each spectrum the differences between the oscillation-frequencies of the individual lines of each triplet is a constant quantity. The law also applies to the series of doublets. The differences between the oscillation-frequencies of individual rays in the series of triplets increases with the atomic weights of the elements to which

who it appears recognised it independently in 1885. J. S. Ames, in 1890, proved the perfect homology, line for line, of the arc spectra of zinc and cadmium. Kayser and Runge have shown that the series of triplets referred to in the magnesium, zinc, and cadmium spectra are in reality three series of harmonic vibrations, a principal series and two subordinate series over-





the triplets belong. A survey of these facts led to the conclusion that as the molecules of these elements contain but one atom, the atoms have a complex constitution, and that those elements the spectra of which are homologous are composed of the same kind of matter in different states of condensation, the molecules having similar modes, but different rates, of vibration. The law of homology has been studied by J. R. Rydberg,

lying each other. The magnesium series they place along with those belonging to the calcium, strontium, and barium family, but according to the periodic law, the magnesium spectrum might be expected also to stand at the head of the spectra of the family which includes zinc and cadmium. The series of lines in the alkali and alkaline earth metals do not fall in with Balmer's law for the hydrogen lines. Homology has been observed

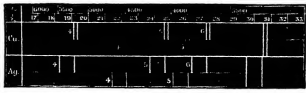
in the spectra of the following groups: lithium, sodium, potassium, rubidium, cæsium, also in aluminium, indium, thallium. Runge announces that doublets and triplets existing in the spectrum of an element can be arranged in series which show an appearance of great regularity, and are seemingly analogous to the overtones of a vibrating body (see figs. 1, 2, and 3). In fig. 1 the spectrum of sodium is shown with a scale of unequal parts, A corresponding to wave-lengths,

and a scale of equal parts 1 corresponding to

oscillation-frequencies. The lines forming the entire spectrum are arranged in three series. Each line in the maps indicates a closely-placed pair of lines. Series P is the principal series, series D is the subordinate series of diffuse lines, series S the subordinate series of sharp lines. The three series are regular, the intervals between the lines decrease as the wave-lengths of the lines diminish from right to left. In fig. 2 we have similar maps, but the scales run the reverse way, from left to right. The principal and subordinate series of lines are shown in the spectra of Mg, Ca, Zn, Cd, and Hg. Corresponding lines in two different series of the same spectrum are numbered alike. The homology of the same group. In fig. 3 the homology of | When it is excited by exposure to a brilliant

determined the composition of some homogeneous alloys of gold and silver, also of lead and cadmium, by comparison of their spectra with check pieces of known composition. Sir J. G. Alleyne estimated phosphorus in iron and steel, and Ballman attempted the estimation of lithium by the spectroscope. Liveing and Dewar made observations on a spectroscopic method of estimating sodium by measuring the width of the sodium lines. Hartley determined the different spectra emitted by solutions of different strengths, and found these constant when the ultra-violet lines were photographed. Accordingly, by comparison, the metals may be determined in solutions made from determinate weights of material, diluted to a definite volume. Quantitative determinations of the basic constituents of limestones, of copper in pyrites cinder, and of lead, silver, and copper in zinc blende, have been made with quantities of matter so small that no other method could have dealt with them. It will be seen later on, that absorption spectra are serviceable in quanti tative analysis.

Spectra of the infra-red region and spectra of solids in vacuo. There are three methods of spectroscopy which have not yet been much employed. 1. The method of E. Beequerel for observing the infra-red rays. of the spectra is evidenced, and also variations | This is accomplished by the use of an eyepiece following increase of atomic mass in elements covered with Balmain's phosphorescent paint.



the spectra of copper and silver is well shown. The difference of two consecutive oscillationfrequencies decreases as these values increase, and there seems to be a finite limit to the oscillation-frequencies of a series. If n represent whole numbers, the oscillation-frequencies of a series may, with great accuracy, be represented by the formula $A-Bn^{-2}-Cn^{-4}$, where A, B, C, are positive constants. B has nearly the same value for all the series of the different spectra. A is the limit towards which the oscillation-frequency tends when n increases. For elements that are chemically related the series are distinctly homologous, both in appearance of the lines and the values of A, B, C, and with increasing atomic weight they shift towards the less refrangible end of the spectrum. The spectra show relationships between the elements similar to those between their chemical proper-

Quantitative analysis by the spectroscope. In 1862 W. A. Miller first attempted to apply the spectroscope to the quantitative estimation of the constituents of an alloy. Jansen, in 1870, proposed two methods of quantitative analysis. Champion, Pellet, and Grenier successfully applied a spectro-photometrical method to the estimation of alkalis. Lockyer and Roberts-Austen

light, it glows, but as soon as any radiation of very low refrangibility falls upon it the phosphorescence is destroyed. 2. Abney observes the same rays by means of specially-prepared photographic plates, sensitive to exceedingly slow vibrations. Even the radiations from a kettle of boiling water can impress themselves permanently on his preparation of silver bromide. V. Schumann and also Waterhouse have used photographic plates prepared with cyanine, erythrosine, and other dyes for the same purpose with some success. 3. Crookes has made much use of the phosphorescent spectrum obtained by the action of an electric discharge in high vacua, and has made a most interesting series of observations on the rare earths by this means. Among other results he has proved the wide distribution of yttria, its occurrence in strontianite, coral, oyster shells, and limestones. He considers that there are several kinds of yttria with different phos-phorescent spectra, but as they all give the same spectrum when excited by the spark in the usual manner, they must be considered as being derived from the same elementary yttrium.

Absorption spectra. In dealing with absorption spectra, we may observe the effect on a continuous spectrum of (a) rases, (b) liquids, (c) solids. When rays enter a transparent mediumeither gaseous, liquid, or solid—they do not leave it in a condition identical with that in which they entered it. They may be either partially absorbed or altered in refrangibility. No known substance is capable of transmitting radiations of all known wave-lengths; those media which do not absorb appreciably any of the coloured rays, or any of the rays in the ultra-violet which are capable of being photographed, absorb a portion of the infra-red spectrum. It may be remarked that substances like air and water which in thin layers do not appear to absorb the rays in the visible spectrum, in thicker layers are seen to absorb the rays of a limited region. The A and B groups of lines in the solar spectrum have been shown to be due to oxygen in the atmosphere, while nine lines lying between the two D lines in the solar spectrum were observed, and proved by Josiah P. Cooke, in 1866, to be caused by water vapour in the atmosphere amounting to not more than 1.5 g. in 100 litres of air. With 0·18 g. only one line was visible, and with 0·55 g. two lines. The successful application by Piazzi Smyth of the spectroscope to weather forecasts is based upon such facts.

Many cases of absorption are known in the infra-red and in the ultra-violet, when no visible rays are absorbed. Colours may be either visible or invisible, since we must regard every substance which absorbs rays as coloured. The eye cannot always distinguish colour, because the most sensitive eye cannot observe the rays which are absorbed by its own aqueous humour, and are thus prevented from affecting the optic nerve. Hence the infra-red rays have no effect upon the retina, and most of the ultra-violet cannot affect it. The colour of a substance is determined by the combined effect of the rays which are not Cobalt glass, which absorbs the absorbed. yellow and green rays, transmits the red, blue, and violet, but as it transmits comparatively little of the red its colour appears blue. The purity of the light by which the substance is examined is of importance. Chromium solutions which are green by daylight are red by gaslight, because the yellow and red rays are predominant in gaslight, and as the red rays and the green are transmitted by pure white light, the dominant colour is red by light of less purity. There is but little importance to be attached to the general absorption of rays, but well-defined absorption bands afford evidence both of the composition and the constitution of substances.

The theory of absorption spectra It is a well-known fact in acoustics that a vibrating musical instrument, a tuning-fork, or a stretched string, will communicate its vibrations to any similar instrument near to it which is capable of similar vibrations, and a sufficient number of strings or tuning-forks can entirely absorb the vibrations of a similar string or fork. In like manner when a molecule of a substance is capable of vibrating synchronously with a radiation, the rays received on the substance are absorbed by reason of the molecules being put in motion by the waves which strike them. The absorption is complete if the direction of vibration of the rays and of the molecule is the same, but the phase is opposite. A general absorption appears (in the ultra-violet at any rate) to be due to the vibration of the molecule, while selective absorption appears to be caused by the vibrations of atoms within the molecule. When a substance in solution absorbs all the rays of a certain region of the spectrum which fall upon it, it is because the molecules are vibrating synchronously with these rays, and the number of molecules present is sufficient to damp all the vibrations. When the rays traverse a smaller quantity of substance-as, for instance, when a solution is diluted—the number of molecules present is not sufficient to damp all the vibrations, and some rays pass through. If, however, certain atoms within the molecule are vibrating synchronously with certain rays, we shall have a selective absorption of these rays after the general absorption has been so weakened by dilution as to allow them to pass. It has been shown by Hartley that the vibrations of the atoms depend upon the rate of vibration of the molecule, or in other words, the rate of vibration of the molecule determines the rate of vibration of its parts. Four absorption bands in the three hydrocarbons benzene, naphthalene, and anthracene are evidence of this. The mean rate of vibration of the rays absorbed by naphthalene and anthracene is less than that of the rays absorbed by benzene, and the vibrations of the carbon atoms within the molecule are not independent of, but are a consequence of, the molecular vibrations; they must, in fact, be regarded as overtones or harmonics of the fundamental or molecular vibrations. When the rate of the fundamental vibration is reduced by an increase in the mass of the molecule, the rate of vibration of the carbon atoms is reduced in a similar ratio. Greater amplitude of vibration means a louder note, or in the case of luminous radiations a brighter light. Consequently the converse of this may be considered to hold good, that a greater intensity of absorption is caused by a greater amplitude of vibration in the molecules of the absorbing medium, the number of atoms or molecules remaining constant.

An examination of the three hydrocarbons benzene, naphthalene, and anthracene shows that the mean rate of vibration of the rays absorbed by benzene is greatest, naphthalene comes next, and anthracene is least; but the absorptive power is in the reverse order, that of anthracene being greatest. Hence we conclude that the amplitude of vibration is greater as the rate of vibration is slower (C. J. 1881). The approximate rates of vibrations of the three hydrocarbons are given as follows:—

	ave-length absorbed.		lar vibra- r second.	Molecular weights.
Benzene	2526	1,248	billions	78
Naphthalene	2687	1,177	,,	128
Anthracene	8439	910	11	178

The various absorption spectra may be referred to in no better order than that of the refrangibility of the rays absorbed.

Infra-red absorption spectra. Abney and Festing have photographed rays extending down to λ 12000; the visible region ends about λ 7600. They studied the absorption spectra of water, hydrochloric acid, chloroform, carbon tetrachloride, cyanogen, and a number of hydrocarbons and their derivatives. The compounds

containing hydrogen show the same group of lines, which, however, do not appear in carbon compounds containing no hydrogen; they do not all appear in some hydrogen compounds. The inference is that they belong to hydrogen. When oxygen is present as a part of hydroxyl it obliterates the rays between two of those lines which are due to hydrogen. When it forms part of the carbon nucleus of a compound, as it does in aldehyde, the spectrum is inclined to be linear, or the bands are bounded by well-defined lines. There appear to be characteristic bands which indicate the carbon nucleus of a series of substances. Certain radicles have a distinctive absorption lying near \$\lambda 7000 and others about λ 9000; thus in the three compounds C_8H_6 , C_8H_5 , H_2N , C_6H_4 , C_2H_5 , H_2N we have the following coincident bands probably due to the benzene nucleus; the line λ 8670 is the principal one; λ 8670, λ 8670 to 8720, λ 8720 to 8880, a fourth band about λ 9300, a fifth about λ 10400-10660. In benzene and ethyl-aniline there occurs a band also at λ 10970 to 11050. If the line λ 8670 is associated with a band, it is almost certain to be caused by the benzene nucleus. Ethyl compounds are indicated by absorption at A 7410, A 8950 to 9030, 9040 to 9070, 9130 to 9180, 9270 to 9300.5, 9320 to 9420. The solar spectrum shows an absorption at A 8660, and, with the exception of the line at 7410, the absorptions noted above have been observed to be coincident with bands or lines in the solar spectrum.

Visible and ultra-violet absorption spectra. The absorption spectra of the coloured region were long since carefully studied by Gladstone. These spectra pass gradually into the extreme violet and ultra-violet. In dealing with the subject it will be convenient first to consider the absorption caused by gases and vapours, elementary or compound, then the absorption by inorganic salts, and lastly the spectra of organic

substances.

Elements which exhibit absorption spectra.

Gases and vapours.

Br, Cl, I, O, ozone, K,
Na.

Metals, the salts of which show banded spectra. Cr, Er, Sm, U, dysprosium, holmium, neodymium, praseodymium, thulium.

Compounds which exhibit absorption spectra.

Gases and vapours.

ICl, IBr, NO₂, H₂O vapour, oxides of Cl.

Salts the acids of which show banded spectra.

Chromates \ Ultra-Nitrates \ violet.

Nitrites (some); per-

manganates.

Organic suostances which exhibit absorption spectra. All benzenoid hydrocarbons and tertiary bases derived from the same. All haloid, hydroxyl, carboxyl, and amide, derivatives of benzenoid hydrocarbons. Cyanuric and uric acids, with colouring matters derived therefrom. Albumens and allied substances. Organic colouring matters of unknown constitution, such as chlorophyll and hæmoglobin. Many alkaloids and their derivatives, e.g.: Aconitine, pseudaconitine, japaconitine, morphine, narcotine, codeïne, papaverine, oxynarcotine, apomorphine hydrochloride, tetracetylmorphine, diacetyl-

codeïne, quinine, quinine sulphate, cinchonine sulphate, quinidine sulphate, cinchonidine sulphate, veratrine, piperine, brucine, strychnine.

The relation of absorption spectra to the chemical constitution of carbon compounds. 1. Substances with a nucleus consisting of an open chain of carbon atoms transmit continuous spectra and are highly diactinic. Examples: the alcohols, acids, ethereal salts, haloid ethers, and the carbohydrates. 2. In homologous series the greater the number of carbon atoms the shorter the transmitted spectrum. 3. In substances with the same carbon nucleus that containing the most oxygen transmits the shortest spectrum, as, for instance, in acids the spectrum is shorter than in alcohols, and in dibasic shorter than in monobasic acids. 4. Substances with a closed chain nucleus when not more than two atoms of carbon are doubly linked transmit continuous spectra, but their absorptive power is much greater than that of open chain compounds; e.g. camphor, benzene hexachloride, and the terpenes, also hexahydropyridine. 5. The more closely the carbons in a closed chain are linked, the greater is the absorptive power of the molecule; e.g. the terpenes withstand dilution more strongly than benzene hexachloride, and camphor. 6. The carbon nuclei of benzenoid hydrocarbons possess strong absorptive power, and after great dilution exhibit absorption bands; e.g. four bands are seen in the spectrum transmitted by benzene when 1 part is diluted with 2,400 parts of alcohol, three bands in naphthalene with a dilution of 1 in 100,000 parts, one band in phenanthrene with 1 in 500,000 parts, and one band in anthracene with 1 in 5,000,000 parts of alcohol. 7. The introduction of radicles in place of hydrogen and of side-chains into the benzenoid nuclei diminishes the number of absorption bands, but increases their width and intensity. 8. The simple union of carbon to nitrogen does not cause selective absorption; e.g. in hydrocyanic acid and cyanides. 9. When a benzenoid nucleus contains an atom of nitrogen in place of an atom of carbon, the molecule exhibits absorption bands, and its absorptive power is generally not impaired, but is sometimes increased, e.g. picoline, pyridine, quinoline, and their derivatives. 10. Isomerides exhibit different spectra, e.g. the terpenes and isomeric benzene derivatives. 11. When the condensa-tion of the carbon atoms in the quinoline nucleus is modified by the combination therewith of four atoms of hydrogen, the intensity of the absorption bands is reduced but is not destroyed. 12. Nitrogen combined with oxygenas in nitroxyl, nitrites, and nitrates—exhibits absorption bands. 13. When several carbon atoms are united by oxygen and nitrogen atoms, the resulting compounds exhibit intense absorption bands, e.g. cyanuric acid, uric acid, and derivatives, including murexide. 14. Molecules of compounds—that is to say, molecules composed of dissimilar atoms—vibrate as wholes or units, and the fundamental vibrations give rise to secondary vibrations which stand in no visible relation to the chemical constituents of the molecule, whether these be atoms or groups of atoms. Hence it appears that a molecule is a distinct and individual particle which is not fully represented by our usual chemical formulæ,

since these only symbolise certain chemical reactions and fail to express any relation between physical and chemical properties. As certain molecular groupings are characterised by the absorption of rays of particular wave-lengths (absorption bands), it is evidently possible to draw conclusions as to the constitution of substances from their absorption spectra. Many of the foregoing statements may be usefully and clearly summarised by means of formulæ. These render evident how the optical properties of a compound are determined by the skeleton of carbon atoms.

compounds which exhibit selective absorption of the ultra-violet rays are coloured. The subtlety of the colour, however, is such that the eye cannot perceive it without the aid of photography or a fluorescent screen.

That which in the ordinary acceptation of the term is a coloured substance is merely one in which the absorption of rays extends into a region limited by the red and violet ends of the spectrum, or between wave-lengths 7000 and 4000. According to this view, the colour of a substance may be invisible to the unaided eye. As bands of selective absorption appear to be the

Typical carbon nuclei, transmitting continuous spectra.

Typical carbon nuclei, exhibiting absorption bands.

Nuclei containing oxygen and nitrogen, exhibiting absorption bands. R representing any radicle.

$$NO_2$$
 RNO₂ RNO₄ OH $NH.CO.C.NH$ $NH.CO.C.NH$ $NH.CO.C.NH$ $NH.CO.C.NH$ $NH.CO.C.NH$ $NH.CO.C.NH$ $NH.CO.C.NH$ $NH.CO.C.NH$ $NH.CO.C.NH$

On the cause of colour in organic compounds. According to O. Witt, the tinctorial character of aromatic compounds is conditional upon the simultaneous presence of a colour-producing group and a salt-forming group in the molecule. A group of atoms of the former kind he terms a chromogen, and of the latter a chromophor. He says: 'For instance, NO₂ is the chromophor of nitraniline and nitrophenol, but nitrobenzene is the chromogen of these bodies. The different nitrodiphenylamines are all acids and dye-stuffs, and their tinctorial power increases with the number of nitroxyls.' In this connexion it is of interest to study the relation of the hydrocarbons to the more complex compounds which are colouring matters derived from them.

If a source of light emits all luminous and invisible vibrations capable of being transmitted through 3 feet of air, a perfectly colourless substance will transmit these rays without impairing their intensity. A coloured substance is one which absorbs rays at either end of the spectrum, or selects rays of a definite wave-length from the middle of the spectrum. Every fluorescent substance is therefore coloured, and benzene, benzenoid hydrocarbons, phenols, and other derived

effect of vibrations taking place within the molecules of a substance, and these are dependent upon the rate of vibration of the molecules themselves, they are called in general 'molecular vibrations.' If, then, it be desired to convert a substance such as benzene, the colour of which is invisible, into a compound with a visible colour, it is necessary to slacken its rate of vibration so that the molecule will absorb rays oscillation-frequencies (inverse lengths) occurring within the limits of visibility. That which is called a chromogen is an invisibly-coloured substance, and that termed a chromophor is an atom or group of atoms capable of so affecting the molecule as to reduce its rate of vibration, so that it absorbs rays within the limits of visibility. Under certain conditions of combination, nitrogen and oxygen are chromophors, hence also nitroxyl and hydroxyl; that is to say, they are themselves coloured either visibly or invisibly. When two benzene molecules are doubly linked by two nitrogen atoms, as in azobenzene, their mode of vibration is profoundly modified, and a brilliant colour as low down in the scale as the yellow rays is the result. A similar modification takes place when two atoms of oxygen replace two of hydrogen, as in | HASSELBERG, B. quinone, which is of a golden-yellow colour. The effect of linking two or more benzene nuclei by carbon atoms has been already dealt with; the colours are not rendered visible, but the molecular vibrations are greatly reduced in rapidity, and the amplitude of the vibrations is much increased. That is to say, the absorption bands are transferred to less refrangible rays, and withstand a much larger amount of dilution. These modifications are both observed in the curve of benzene as modified in triphenylmethane. The vibrations of this radicle are reduced in rapidity in rosaniline hydrochloride, Hofmann's violet, iodine green, and other such derivatives in which the carbon nucleus remains intact.

There is a family likeness in the curves of azo-benzene, azo-naphthalene, chrysoïdine, and other derivatives of these substances, which is quite in accordance with these observations. In short, in a general way, the change of colour effected by any given series of reactions may be predicted from an inspection of the curve of molecular vibrations of any benzenoid hydrocarbon. W. N. H.

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OPTICAL METHODS .- Section 3: ROTA-TION OF THE PLANE OF POLARISATION OF LIGHT .- When a ray of plane polarised light is passed through a plate of quartz cut at right angles to its optical axis, it is found that the plane of polarisation of the emergent ray is not coincident with that of the incident ray; the angle through which the plane has been rotated is called the angle of rotation. If the rotation takes place in the same direction as that in which the hands of a watch move when the face of the watch is looked at, the quartz is said to show dextrorotatory power; if the rotation is in the opposite direction, the quartz is said to show lævorotatory power. The symbol + is used to express dextrorotation, and the symbol - to express lævorotation. Substances which rotate the plane of polarisation of a ray of light passed through them are said to be optically active.

The polarimeter is an instrument wherein a ray of light can be polarised, the position of the plane determined, the ray passed through a determinate quantity of an optically active substance, and the position of the plane of the

emergent ray determined.

The angle of rotation depends upon (1) the nature of the active substance, (2) the thickness of the layer of the substance through which the light passes, (3) the wave-length of the light used, and in most cases (4) the temperature.

The composition of the substance examined must then be known; the length of the column of liquid used if the substance be liquid, or the thickness of the layer if the substance be a solid, must be measured; the temperature must be noted—observations ought to be made at a common temperature; and monochromatic light should be employed.

Suppose we are dealing with a liquid carbon compound. Let l = length of column of liquid in decimetres, d = sp.gr. of liquid (referred to water), and a = angle of rotation of the plane of polarisation of light of stated wave-length; then

$$[\alpha] = \frac{a}{l \cdot d}$$

The value of $[\alpha]$ is usually called the specific rotatory power of the substance. It is customary to indicate the light employed by placing a letter to express the wave-length below the bracket; thus $[\alpha]_D$ means spec. rot. power for light of wave-length D. The spec. rot. power of a liquid as thus defined is the angle through which the plane of polarisation of a ray of light of stated wave-length is rotated by passing through a column one decimetre long of the liquid containing one g. of the substance in 1 c.c.

If a solid compound is to be examined, it must be dissolved in an optically inactive solvent. In this case l = length of column of solution in decimetres, d = sp. gr. of solution, p = g. of optically active substance in 100 g. of solution; then, assuming that the solvent is without influence on the rotatory power of the

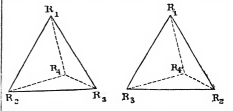
dissolved substance, $[a] = \frac{100a}{l \cdot p \cdot d}$. The question as to the effect of an inactive solvent on the rotatory power of an active substance in solution will be examined later (p. 257).

Connections between optical activity and scope, English trans.) has divided substances which rotate the plane of polarisation of a ray of light into three classes: (1) those which are active only when in the form of crystals; (2) those which are active only when liquid or in solution; (8) those which are active both as crystals and also in solution or in the liquid state. The only substance at present known to belong to the third class is strychnine sulphate. The liquid crystals examined by Lehmann (Z. P. C. 4, 462; 5, 427) probably belong to the first class.

The optical activity of carbon compounds. All compounds which exhibit optical activity unconnected with crystalline form—in other words, all compounds which are optically active when liquid or in solution—are compounds of carbon. Le Bel (Bl. [2] 22, 337) was the first (1874) to trace a definite connection between the optical activity and the constitution of carbon compounds. He was followed by van't Hoff (Bl. [2] 23, 295 [1875]). The subject has been much advanced by van't Hoff in his pamphlet, La Chimie dans l'Espace, and more especially by a small book published in 1887, Dix Années dans l'Histoire d'une Théorie (translated into English by Marsh, and published in 1891, entitled Chemistry in Space).

The hypothesis of Le Bel and van't Hoff

connects optical activity with the presence of one or more asymmetric atoms of carbon in the molecule of the active compound. An asymmetric atom of carbon is one which is in direct combination with four different atoms or radicles, these atoms or radicles being arranged so that any three are similarly situated with regard to the fourth. The conception of the asymmetric atom carries with it the conception of the arrangement in space of the parts of the molecule. The asymmetric atom of carbon is supposed by van't Hoff to be placed in the centre of a regular tetrahedron, and a different radicle is supposed to be held by the carbon atom at each summit of the tetrahedron. Two forms of this arrangement may exist, and these forms are geometrically different. Neither is superposable on the other. They bear to one another the re-lation of an object to its image, or of the right hand to the left, as shown in the figure.



This arrangement corresponds to the enantiomorphous (non-superposable) form of crystals. Ammonium malate, for instance, crystallises in two non-superposable forms, which differ in exactly the same way as the molecules of two geometrical isomerides, each containing an asymmetric carbon atom and both having the composition CR1R2R3R4, are supposed to differ. If one of the isomerides CR₁R₂R₃R₄ rotates the plane of polarisation to the right, the other rotates to the left. If this hypothesis is to be accepted, every compound of carbon which is optically active must contain at least one asymmetric carbon atom. So far as investigation has gone, this deduction from the hypothesis has been confirmed (v. van't Hoff, Dix Années &c., 31; Landolt, Handbook, 25 et seq.). Further, it has been found that compounds which do not themselves contain asymmetric carbon atoms are inactive, although they are derived from optically active compounds.

There is no doubt that compounds exist which contain asymmetric carbon atoms but do not affect the plane of polarisation of a ray of light. At first sight this fact might be supposed to be irreconcileable with the hypothesis; but the recognition of the existence of inactive compounds containing asymmetric carbon atoms has led to a most ingenious and interesting development of the hypothesis.

Some inactive compounds containing asymmetric carbon atoms can be separated each into two optically active isomerides, one of which is dextrorotatory and the other is lævorotatory. Some inactive compounds which contain asymmetric carbon atoms cannot be separated into active isomerides. Racemic acid, which can be resolved into dextrorotatory and lævorotatory tartaric acid, is an example of the first of these classes, called inactive resolvable compounds;

mesotartaric acid, which cannot be resolved into active isomerides, is an example of the second class, called inactive non-resolvable compounds.

Inactive resolvable compounds are said, in the language of van't Hoff's hypothesis, to be inactive by external compensation; inactive nonresolvable compounds are said to be inactive by internal compensation.

To understand these expressions consider

the formulæ (R₃R₂R₁) C.C(R₁R₂R₃) and $(R_2R_3R_1)C.C(R_1R_3R_2)$; an italicised C represents an asymmetrical atom of carbon. Each formula contains two asymmetric carbon atoms, and both carbon atoms in either formula are united to the same radicles; in the first formula each C is united to the radicles R_1 , R_2 , R_3 , and $C(R_1R_2R_3)$; in the second formula each C is united to the radicles R_1 , R_2 , R_2 , and $C(R_1R_3R_2)$. The structure represented by one of these formulæ is the reflected image of that represented by the other: hence if one of these geometrically isomeric compounds is dextrorotatory, the other will be levorotatory, and both will rotate the plane of polarisation to the same degree. But if a compound were formed by the combination of equal numbers of molecules of these two geometrical isomerides, that compound would be inactive, for the tendency to righthanded rotation of one part of the compound molecule would be exactly neutralised by the equal tendency to left-handed rotation of the other part. Moreover, if the inactive compound were split into two compounds, one consisting of molecules of the form represented by the first of the above formulæ, and the other of molecules of the form represented by the second formula, then each of these compounds would be optically active, and the compounds would be active in opposite directions. The hypothetical inactive compound $(R_3R_2R_1)C.C(R_1R_2R_3)$ would be said

to be inactive by external compensation.

Racemic acid, which is optically inactive, can be resolved into dextrorotatory and lævorotatory tartaric acids. Both tartaric acids have the formula C₂H₂(OH)₂(CO₂H)₂; the hypothesis we are considering represents one of these acids as (H.OH.HOOC) C.C(COOH.OH.H); and the other acid as (H.COOH.OH) C.C(OH.COOH.H); and the hypothesis represents racemic acid as made up of an equal number of molecules of the two active acids, and therefore as a substance which is inactive by external compensation.

But suppose the molecule of a compound contained two asymmetric carbon atoms, both united to the same radicles but having one half of the molecule the reflected image of the other, then the tendency to right-handed rotation belonging to one part of this molecule would be neutralised by the tendency to left-handed rotation belonging to the other part of the molecule: the molecule as a whole would be incapable of rotating the plane of polarised light, and the compound would be inactive by internal com-pensation. The general formula of such an inactive compound is $(R_2R_2R_1)C.C(R_1R_2R_3)$.

A compound of this type cannot be resolved into active isomerides, because a resolution into parts is only possible by splitting the molecule, removing one of the radicles, and so destroying the asymmetry of one, or both, of the atoms of

carbon. Mesotartaric acid, which is non-resolvable, probably belongs to this class of compounds; it may have the formula

(H.COOH.OH)C.C(COOH.OH.H).

The hypothesis therefore provides for the exis. tence of two classes of optically inactive compounds, both containing asymmetric carbon atoms; the members of one class are resolvable into active isomerides, the members of the other class cannot be so resolved.

Those compounds which contain in their molecules more than one asymmetric carbon atom, each of which atoms is united to the same radicles, whether these radicles are arranged in space in exactly the same way, relatively to each carbon atom, or not, are called by van't Hoff symmetrical compounds; and the term unsymmetrical compounds is used to designate those which contain more than one asymmetric carbon atom each of which is united to different radicles. The typical formulæ for symmetrical compounds containing two asymmetric carbon atoms

 $(R_1R_2R_3)C.C(R_3R_2R_1), (R_1R_3R_2)C.C(R_3R_2R_1), \&c.$ The typical formula for an unsymmetrical compound containing two asymmetric carbon atoms is $(R_1R_2R_3)C.C(R_4R_5R_8)$.

Unsymmetrical compounds may be either active or inactive; if inactive, they must be inactive by external compensation, and therefore they must be resolvable into pairs of active isomerides. Symmetrical compounds may be active or inactive; if inactive, they may be inactive by external, or by internal, compensation, and therefore they may be either resolvable or not resolvable.

An unsymmetrical compound containing n asymmetric carbon atoms may exist in 2ⁿ isomeric forms, which will always be grouped in pairs, one being dextrorotatory and the other lævorotatory. A symmetrical compound containing n asymmetric carbon atoms may exhibit $\frac{1}{2}$ ⁿ active isomeric forms, grouped in pairs of opposite rotatory power, and may also exist in

122 non-resolvable inactive modifications (van't Hoff; Dix Années &c., 54-5).

There are three general methods for separating inactive resolvable bodies into their dextrorotatory and lævorotatory isomerides.

In the first method advantage is taken of the differences between the actions of certain minute organisms on the two active isomerides. The second method proceeds by treating the inactive compound with an active body with which one of the isomeric constituents of the inactive compound combines more readily than the other. The third method consists in separating the inactive body into two active isomerides by crystallisation at a definite temperature (van't Hoff, l.c. 63-69).

Racemic acid, which is an inactive compound, can be resolved into equal quantities of rightand left-handed tartaric acid by each of the three methods. When penicillium is allowed to act on a dilute solution of ammonium racemate, lævorotatory ammonium tartrate is found in the solution after a time, the dextrorotatory tartrate having been destroyed by the action of the organism. When a quantity of active cinchonine sufficient to neutralise half of a determinate

quantity of racemic acid is added to a solution of the racemic acid, crystals of lævorotatory cinchonine tartrate are obtained, and dextrorotatory tartaric acid remains in solution. When a solution of racemic acid is neutralised by soda, and another equal quantity is neutralised by ammonia, and the solutions are mixed, and evaporated at a temperature slightly below 28°, crystals of two forms are obtained; one set of crystals is dextrorotatory, and the other is lævorotatory, sodium-ammonium tartrate. Hoff (l.c. 69) has shown that sodium-ammonium racemate is changed to a mixture of the active isomeric tartrates by heating the dry salt with water, in the ratio NaNH, H,C,O,H,O:3H,O (the salt crystallises with H₂O), to a little under 27°, and that the reverse change is effected by heating the mixed tartrates to a little above 27°. The changes may be represented thus-

centre of figure of the tetrahedron. When the molecule is asymmetric, the centre of gravity will not be situated on any one of the six planes of symmetry of the tetrahedron. So long as the substitution of one radicle by another does not move the centre of gravity of the molecule to the other side of one of the planes of symmetry of the tetrahedron, Guye supposes that the rotatory power is increased or diminished, but is not changed in sign, by such substitution; but that a change of sign of the rotatory power accompanies a substitution which results in moving the centre of gravity of the molecule from one side to the other side of one of the planes of symmetry of the tetrahedron. Guye gives about fifty cases to which he has applied his hypothesis successfully. The derivatives of tartaric acid afford the most complete example.

Replacement of the acidic hydrogen of tar-

$$2(\text{NaNH}_4.\text{H}_4\text{C}_4\text{O}_6.4\text{H}_2\text{O}) \stackrel{\longrightarrow}{\sim} 2(\text{NaNH}_4.\text{H}_4\text{C}_4\text{O}_6.\text{H}_2\text{O}) + 6\text{H}_2\text{O}.$$

The racemate crystallises with H₂O, and the tartrates with 4H₂O. Slight variations of temperature above or below 27° determine the direction in which the change shall occur.

The change of inactive sodium-ammonium racemate to a mixture of the active tartrates, and vice versd, is very similar to some changes which occur among inorganic compounds; for instance, when a mixture of MgSO₄.7H₂O and Na₂SO₄.10H₂O is heated to a little above 21°, it is changed to the double sulphate MgNa₂(SO₄)₂.4H₂O, and water, and this double sulphate is resolved into the two single sulphates at a little under 21°; these changes may be represented thus:—

taric acid by alcoholic radicles is accompanied by increase of rotatory power, and the greater the molecular weight of the replacing alkyl the greater is the increase of rotatory power; thus—

Methyl tartrate
$$[a]_D = +2.14$$

Ethyl , 7.66
Propyl , 12.44
Isobutyl , 19.87

Replacement of the hydroxylic hydrogen of the acid by benzoyl moves the centre of gravity to the other side of the plane of symmetry which is situated between the COOH and OH groups; this replacement is accompanied by a change of sign of the rotatory power. But if the H of the COOH group is now replaced by one alkyl group,

$$MgSO_4.7H_2O + Na_2SO_4.10H_2O \longrightarrow MgNa_2(SO_4)_2.4H_2O + 13H_2O.$$

Van't Hoff calls the temperature at which such a chemico-physical change as this occurs the transition-point of the system (v. Z. P. C. 1, 165, 227).

Extension of the hypothesis of van't Hoff and Le Bel. The hypothesis of van't Hoff and Le Bel connects the power of rotating the plane of polarisation of a ray of light primarily with the configuration of the parts of molecules, but it points to the formation of molecular aggregates, without change of molecular structure, as a cause of the disappearance of optical activity. Although optical activity is primarily dependent on the arrangement of the parts of certain molecules, and although we cannot assign a definite part of the total rotatory power of a molecule to each atom, or to each group of atoms which together form the molecule, nevertheless a consideration of the constitution of optically active compounds shows that the amount of the activity of any compound is dependent on the greater or less differences between the four radicles which are in direct union with the asymmetric carbon atom or atoms in the molecule of that compound. The greater the differences between the radicles the greater will be the asymmetry of the molecule, and, therefore, the greater will be the rotatory power. An attempt has been made by P. A. Guye (C. R. 110, 714) to connect the degree of asymmetry of the molecule of an optically active compound with the mass of each radicle, and the distance of its centre of gravity from the the centre of gravity is moved back towards the plane of symmetry already mentioned, and the rotatory power is diminished. Thus—

Dibenzoyl tartaric acid $[a]_D = -117.7$ Methyl salt of , 88.8 Ethyl , , , 60.0 Isobutyl , , , 42.0

If acetyl is used in place of benzoyl the rotatory power changes its sign, but the value of the leworotation of diacetyl tartaric acid is less than that of dibenzoyl tartaric acid; the replacement of acidic hydrogen in the diacetyl acid by alkyl radicles decreases the rotatory power, and, as the alkyl radicles of large molecular weight exert a greater effect than the acetyl group, and an effect in the opposite direction, the rotatory power again changes its sign. Thus—

Diacetyl tartaric acid $[a]_D = -23.1$ Methyl salt of , 11.3 Ethyl , , , 6.5 Isobutyl , , , 10.3

Crum Brown (Pr. E. 17, 181) has tried to connect the greater or less rotatory powers of different active compounds with differences between the constitutions of the substituting radicles. Crum Brown admits the influence of the masses of the radicles, but he tries to show that their constitution must also be taken into account. If two methyl groups are introduced into dextrorotatory tartaric acid, the salt thus obtained is still dextrorotatory; in this change

two CO.OH radicles have been changed into two radicles CO.O.CH₂. If now this dimethyl trate is treated with acetyl chloride, a strongly lævorotatory dimethyl diacetyl-tartrate is obtained; in this change the CO.O.CH₃ groups remain, and two OH groups have been changed to two groups O.CO.CH₂. In the first change, the introduction of the group CO.O.CH₃ for H altered the rotatory power but did not change its sign, in the second change the introduction of the group O.CO.CH₃ changed the sign of the rotatory power; therefore, Crum Brown argues, the effect of the radicle O.CO.CH₃ on the rotatory power is greater than the effect of the isomeric radicle CO.O.CH₃. As the masses of these two radicles are the same, it is evident that the constitution of the substituting radicle affects the rotatory power of the compound.

The specific rotatory powers of compounds often change more or less with small changes in certain physical conditions. Some active bodies become inactive by heating, and at another temperature the change is sometimes reversed. The values of the rotatory powers of active bodies dissolved in inactive solvents are dependent on the nature and the quantity of the solvent used (v. post, p. 257); the rotatory power of a solution sometimes changes on keeping until a constant value is attained (v. Landolt's Handbook of the Polariscope, 62). This readiness to change shown by the rotatory powers of carbon compounds finds some explanation in van't Hoff's hypothesis, and especially in the development of it made by Wislicenus. According to Wislicenus (Kön. Sächsischen Ges. der Wiss. 14, 1) the arrangement of the four radicles attached to an asymmetric carbon atom is conditioned chiefly by the affinities of these radicles for one another in the sense that those radicles which have the greatest mutual affinity will tend to get as near as possible to one another; but besides those configurations which are conditioned by the affinities of the radicles, and which will be comparatively stable, other configurations will probably exist conditioned by the temperature and by the collisions of other molecules—for instance, the molecules of the solvent—and these arrangements will be relatively unstable. These unstable forms may be optically active, as they are only geometrically different from the stable forms, but their rotatory powers will probably differ from those of the stable forms. This way of looking at the question of optical activity enables us to connect the fact that active compounds easily undergo changes in the values of their rotatory powers with the conception which the hypothesis of van't Hoff and Le Bel furnishes of the cause of optical activity.

Molecular rotatory power. The notion of molecular rotation, as the product of the specific rotation into the molecular weight of a substance, was introduced by Wilhelmy (P. 81, 527). Krecke (J. pr. [2] 5, 12) proposed to define molecular

rotatory power as
$$[m] = \frac{m}{100} \times \frac{\alpha}{1.d}$$
, where $m =$

molecular weight of compound, and α , l, and d have the same meaning as before (p. 253); m is divided by 100 to obviate the use of inconveniently large numbers.

Krecke endeavoured to generalise the rela-

tions between the values of [m] for certain compounds and bodies derived from, or closely connected with, these compounds. Some years before Krecke's attempt, Mulder (Z. 1868.58) put forward a statement to the effect that optically active compounds contain certain active radicles, which may be transferred to other active atomic aggregations, and the rotatory powers of the new bodies bear a simple relation to those of the original radicles. Krecke, on the basis of further experimental results, modified and extended Mulder's statement. Krecke announced two generalisations:

(i.) 'When an optically active body forms a compound with an inactive body, or when it is altered by the action of chemical reagents, either the molecular rotatory power of the body is unchanged, or the molecular rotatory power of the new substance is a simple multiple of that of the parent substance.'

(ii.) 'The molecular rotatory powers of isomerides are multiples of one and the same value.'

Most of Krecke's values were obtained from measurements of substances in solution, and, as we shall see in another paragraph, the results of such measurements are trustworthy only in a few cases. Landolt (B. 6, 1073) made a careful examination of the rotatory powers of tartrates in solution; he came to the conclusion that the value of [m] for tartrates containing a single atom of a monovalent metal is double the value of [m] for free tartaric acid, and that [m] for tartrates containing two atoms of a monovalent metal is three times [m] for the free acid. In this case, Krecke's first statement was confirmed.

In 1885, Oudemans (v. P. B. 9, 635) arrived at the conclusion that the compounds of active bases with inactive acids, and also the compounds of active acids with inactive bases, follow the simple law that 'the molecular rotatory power of a salt, in fairly dilute solution, is independent of the nature of the inactive part of the salt.' This result has been confirmed by the measurements of camphorates and tartrates made by Landolt, of salts of cholic acid by Hoppe-Seyler, of malates by Schneider (A. 207, 286), and of camphorates by Hartmann (B. 21, 221).

This result indicates that the rotatory powers of salts in solution are not affected, to anything like the extent that the rotatory powers of liquid compounds are affected, by the nature of the radicles which compose them. The rotatory power of a salt in solution approaches to being an additive property, whereas the rotatory power of a liquid compound is a constitutive property (cf. Ostwald, Lehrbuch, 1, 499 [2nd ed.]).

Optical activities of bodies in solution. The specific rotatory power of an active substance in an inactive solvent has been defined (p. 253) as $[a] = \frac{100a}{l \cdot p \cdot d}$, where l = length of column of solution in decimetres, d = spec. grav. of the solution, and p = g. of active substance in 100 g. of the solution. This formula is expressed in more general terms as $[a] = \frac{n \cdot a}{l \cdot g \cdot d}$, where n = g. of solution containing g g. of the substance; in this case, 1 c.c. of the solution weighs d g.

and contains $\frac{g.d.}{g.}$ of the active substance.

This formula assumes that the rotatory power of the dissolved active substance is not affected by the inactive solvent. In 1838 Biot (A. Ch. [3] 36, 257; cf. 59, 206) found that the rotatory power of tartaric acid in aqueous solution increased with increasing dilution; that solutions, of equal concentration, of the same active substance in different inactive solvents had different rotatory powers; and that the same inactive solvent raised the rotatory powers of some active substances, and decreased the rotatory powers of others. Results similar to those of Biot were obtained by Oudemans (A. 166, 65; 182, 33), Hoorweg (Maanblad voor Naturwet, 3, 12 [1873]), and Hesse (A. 176, 89, 189).

Landolf (A. 189, 241; B. 21, 191; Handbook of the Polariscope, 80-94) made many observations on the effect of inactive solvents on the rotatory powers of active substances; he came to the conclusion that the specific rotatory powers of such solutions are expressed by equations of the following forms: $[\alpha] = A + Bq$, or

[
$$\alpha$$
] = A + B q + C q^2 , or [α] = A + $\frac{Bq}{C+q}$, where A, B,

and C are constants, and q is weight of active substance in 100 pts. of the solution. Which formula must be employed is discovered from the results of several observations made with solutions of different concentrations in different solvents. In some cases the observed values of [a] for liquid compounds agree with the values calculated from series of observations of a for solutions of the same compounds. Thus Landolt got the following results:—

on the solubility in water of the salt used; the greater the atomic weight of the metal, the greater was the concentration whereat the change began, and the greater was the difference from the normal state of the salt. Schneider (A. 207, 257) found that the rotation of an aqueous solution of malic acid was right-handed in concentrated solutions, left-handed in dilute solutions, and at 34 p.c. was equal to zero. These observations, and others of a similar character, seem to me to show that observations of the rotatory powers of solutions of solid compounds cannot be applied, at least not in all cases, to draw conclusions concerning the connections between the rotatory powers and the chemical constitutions of the compounds themselves. There is undoubtedly an action of some kind between the molecules of the active dissolved substance and the molecules of the inactive solvent. Molecular aggregates of the solid substance may be broken down by the solvent, or the solvent may form a series of unstable compounds with the dissolved substance, or the solvent may bring about changes in the configuration of the atoms which form the molecule of the dissolved body, or the dissolved compound may be electrolytically dissociated into ions by the solvent. Some of these actions may occur in certain cases, and others in other cases. The action of much water on active salts dissolved therein seems to consist in the electrolytic dissociation of the salts into their ions.

Magnetic rotatory power. Following observations made in 1871 by A. de la Rive (A. Ch. [4] 15, 57), Becquerel in 1877 (A. Ch. [4] 22, 5) made a number of measurements of the rotations.

	[a] _D	[a] calculated from observations on solutions in					
Active substance	observed	(1) EtOH	(2) MeOH	(3) H ₂ O	(4) C _e H _e	(5) AcOH	Max. diff.
Dextro-ethyltartrate		8.27	8.42	8.09		_	22
Dextro- turpentine	14·15 37·01	14·87 36·97	_		36.97	36.89	+ .72
Lævo- turpentine Lævo- nicotine	161.55	160.83	_	161-29			72

It is not possible to determine with certainty whether the true values of [a] for solid compounds can be determined from observations on solutions of such compounds. In some cases the values obtained seem to approach constancy when the solutions become very dilute; but according to Pribram (B. B. 1887. 505; B. 20, 1840), who used a very delicate method of measuring, [a] does not certainly attain final constant values in aqueous solutions of tartaric acid, nicotin, and cane sugar, even with very great dilution. R. von Sonnenthal (Z. P. C. 9 656) determined [a] for dilute aqueous solutions of normal and acid tartrates of K, Na, Li, and NH,; he found that [a] decreased as dilution increased, but that when the solutions became very dilute (c. 4 to 2 p.c. salt in solution) [a] began to increase as dilution still further increased. The conclusion come to by von Sonnenthal was, that a change in the state of the dissolved tartrates occurred when more water was added to the very dilute solutions; this change was supposed to be of the nature of dissociation. The dilution whereat the change occurred was found to depend on the nature of the metal of the tartrate, and

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tory powers of liquid substances, and of salts in solution. The subject was taken up by Perkin in 1882, since which time a series of memoirs by this investigator has appeared (C. J. 45, 421; 49, 777; 51, 808; 53, 561; 55, 680; 59, 981; 61, 800). The methods employed in the measurements are based on the discovery which Faraday made in 1846 (T. 1846. 1), that most transparent bodies acquire the power of rotating the plane of polarisation of a ray of light when they are brought within the action of a magnet, or of an electric current which is made to pass round the body so that its plane is at right angles to the direction of the ray of light. Faraday showed that the amount of rotation is proportional to the strength of the current, or the intensity of the magnetic action, and to the length of the layer through which the light passes, and that it is dependent on the temperature and on the nature of the substance examined. Perkin has worked with liquid compounds and with solutions. fluid to be examined was placed in a glass tube about 103 mm. long. and 9 mm. diameter, the ends of which were closed by circles of glass cemented on; the ends of the tube were let into the pole-pieces of a large electro-magnet. Sodium light was employed by Perkin. (For a description of the apparatus, and the various precautions and corrections, v. Perkin, C. J. 45, 421-445.) Perkin calculates his results so that the observed rotations of liquid compounds are referred to lengths of the liquids related to one another in the same proportion as the molecular weights of the gaseous compounds obtained by vapourising the liquids; the results when thus calculated represent the magnetic rotatory powers of quantities of the various compounds proportional to the weights of the gaseous molecules of these compounds.

The molecular rotation of water is taken as unity. If r is the rotation of unit-length of liquid, and r' is the rotation of the same length of

water under the same conditions, then - is the specific rotation of the liquid; and

sp. rot. $\times \frac{M}{M' d}$ is the molecular rotation, where

is not constant; the change from a normal paraffin, CH₂,nCH₂.CH₃, to the next higher isoparaffin, CH(CH₃)₂.nCH₂.CH₃, raises Mol. R. by 1.023 + .105; in the change from a normal acid to the next higher iso-acid of the same series, CH₂ has a different value; and so on. When Cl is substituted for H in a hydrocarbon the mol. rotatory power increases, but each Cl atom has a different value from the others. So with the replacement of H in a hydrocarbon by OH; each OH group has its own value, the effect on Mol. R. of the second OH being less than that of the first. So again with the group NO3; the second NO, does not increase Mol. R. so much as the

The molecular rotatory powers of various homologous series of carbon compounds may be expressed by the general formula

(Mol. R.) = c + n1.023, where n = number ofatoms of C in the molecule of any stated compound in the series, and c is a constant for each series. Perkin gives the following table (C. J. 45, 574):-

Mol. R. of members of homologous series.

Alcohols $C_nH_{2n,2}O$ 699	
,, iso-and sec ,, '844 ,	, ,
	,
	•
,, iso	
Aldehydes $C_nH_{2n}O$ 261	,
" iso- and ketones " 375	,
	•
,, iso ,, 509 ,	
Formic ethereal salts (Et, and higher) . ,, 495 ,	•
Acetic ,, ,, ., .,, .370 ,	,
,, iso- ,, '485 ,	,
Methyl salts	,
Ethyl ,, and higher salts ,, 337 ,	,
,, iso- , ,, ·449 ,	,
Succinic methyl salts $C_n H_{n_1, n_2} O_1 = 0.93$,
" ethyl " " 196 ,	,
,, ,, iso ,, ·422	,
Chlorides $C_nH_{2n+1}Cl$ 1.988 ,	,
,, iso- and sec ,, 2.068 ,	,
Bromides $C_nH_{2n+1}Br 3.816$,	,
,, iso- and sec ,, 3.924 , Iodides $C_nH_{2^{n+1}}I$ 8.011	,
Todides $C_nH_{2^{n+1}}I$ 8.011	,
" iso- and sec " 8.099	,
,, iso- and sec , 8099 Ethyl salts, unsaturated $C_nH_{2_1-2}O_2$ 1451 ,	,

M = molecular weight of substance, M' = molecular weight of water, and d = relative density of substance. The calculation may be expressed by

one equation, thus: (Mol. R.) =
$$\frac{r \cdot M}{r' \cdot M' \cdot a'}$$

About 200 compounds have been examined by Perkin, the observation of r being repeated five to ten times for each compound, and the specific gravity of each being determined with great

Magnetic molecular rotatory powers of liquid carbon compounds. In strictly homologous normal series of carbon compounds, each increment of CH2 produces a constant increase in molecular rotatory power, amounting to 1.023. But when the addition of CH₂ is accompanied by a change in the distribution of the atomic interactions the change of Mol. R. At. R. of H.

· These results indicate that the molecular rotatory power of a liquid carbon compound is not the sum of certain constant values belonging to each atom or atomic group, but that it depends, to some extent, on the arrangement of the atoms which form the molecule. This result is confirmed by the outcome of attempts to assign values to the atomic rotatory powers of oxygen and carbon. The following data are given as examples of such attempts.

The atomic rotatory power of hydrogen may be deduced thus:

1. Mol. R. $C_nH_{2n+2} = Mol.$ R. $nCH_2 + H_2$; but Mol. R. $nCH_2 = n1.023$.

The value thus deduced for At. R. of H = ·254.

2. Mol. R. C_nH_{2n+2} minus Mol. R. C_nH_{2n+1} =

Thus, Mol. R.
$$C_sH_a = 3.577$$

Mol. R. $C_sH_7 = 3.323 = Mol.$ R. C_sH_{14}
At. R. of $H = .254$
Now, At. R. of $C = Mol.$ R. CH_2 minus At. R. of

 $\mathbf{H} \times \mathbf{2} = 1.023 - .508 = .515.$

By applying similar methods to the data for compounds of O, the following results are obtained:

The At. R. of Cl varies according to the series of compounds considered, and also as one or two atoms of H are replaced by one or two atoms of Cl, according as the H replaced is in one part of the molecule or in another part, and so on. Two values are obtained for At. R. of N, according as the atom of N is directly connected with 3 or 5 other atoms or groups. The general conclusion is that changes in the magnetic molecular rotatory powers of liquid carbon compounds are intimately connected with changes in molecular structure, so that any cause which alters this structure also alters the rotatory power.

In C. J. 61, 800, Perkin gives an elaborate study of the constitution of ethyl acetoacetate and allied compounds, which elucidates vericlearly the connections between the constitution and the magnetic rotatory powers of carbon

compounds.

Mol. R. of compounds in presence of water. Perkin has used determinations of Mol. R. of certain compounds before and after addition of water, to throw light on the question whether hydrates are formed by the action of water on these compounds (C. J. 49,777; 51,808; 55,680). In all measurements of Mol. R. the molecular rotatory power of water is taken as unity; if therefore a compound is formed by the addition of water to another compound, the Mol. R. of the new compound might be expected to be nearly equal to that of the original compound, plus one unit for each molecule of water added; if the observed Mol. R. is distinctly less than that calculated in this way, the difference may be explained by supposing that the formation of the new compound has been accompanied by a rearrangement of the atoms of the reacting molecules.

The following are examples of the application of this method. In each case the compound and

formic, acetic, and propionic acids in the ratio of equal molecules of water and acid, either a hydrate or a mixture is formed in each case; but that when H₂SO₄ and H₂O react in about the ratio H₂SO₄:H₂O (or a little more H₂O than this), a rearrangement of atoms occurs with formation of a new compound of S, O, and H (perhaps SO(OH)₄); and that reactions of this nature, involving rearrangements of atoms, also occur between HNO₃ and H₂O, and CCl₃.CHO and H₂O.

The following data are interpreted by Perkin to mean that a solution of NH, in water, or in alcohol, 'simply consists of the solvent and ammonia':

Mol. R.
$$NH_{J} + 2 \cdot 1H_{J}O = 3 \cdot 91$$

, $2 \cdot 1H_{J}O = 2 \cdot 1$
, NH_{3} $1 \cdot 81$
Mol. R. $NH_{J} + 2 \cdot 18C_{2}H_{a}O = 7 \cdot 886$
, $2 \cdot 18C_{2}H_{a}O = 6 \cdot 06$

Mol. R. of acids and salts in aqueous solutions. Perkin (C. J. 55, 680; 59, 981) gives Mol. R. for HCl as 2·187. This is arrived at by finding values for At. R. of H and Cl from observations of Mol. R. of various chlorides of alcoholic radicles. By similar methods of calculation values for Mol. R. of HBr and HI were obtained. The three results are:

The value for HCl was confirmed by measuring Mol. R. of HCl dissolved in isoamyl oxide; the result was Mol. R. HCl = 2.24.

Values were then obtained for the three acids in aqueous solutions. The numbers increased as dilution increased; in any case they were much larger than the values obtained by the method stated above. The following table gives the minimum and maximum values for each acid:

Mol. R. in aqueo solutions.	us Maximum.	Minimum.
HCl	4.419	4.045
	(15.63 p.c. HCl)	(41.7 p.c. HCl)
$\mathbf{H}\mathbf{Br}$	8.519	7.669
	(15·47 p.c. HBr)	(65.99 p.c. HBr)
\mathbf{HI}	18.451	17.769
	(31·77 p.c. HI)	(67·02 p.c. HI)

Compound and water	Mol. R. observed	Mol. R. calculated for	Diff,
HCO,H.H,O	2.666	H.CO ₂ H + 1 = 2·671	- ·005
CH,CO,H.H,O	3.554	CH ₂ CO ₂ H + 1 = 3·525	+ ·029
C,H,CO,H.H,O	4.512	C ₂ H ₃ CO ₂ H + 1 = 4·462	+ ·05
H,SO,H,O	3.188	H ₂ SO ₄ + 1 = 3·315	- ·127
H,SO,2H,O	4.113	H ₂ SO ₄ + 2 = 5·188	- 1·075
H,SO,3H,O	5.064	H ₂ SO ₄ + 3 = 6·188	- 1·124
HNO,267H,O	3.656	HNO ₃ + 2·67 = 3·85	- ·194
CCl,CHO.H,O	7.037	CCl ₂ CHO + 1 = 7·591	- ·554

water were mixed in the ratio shown by the formula given, and the Mol. R. of the product was determined; the Mol. R. of the product was then calculated by adding one unit for each molecule H₂O to the observed Mol. R. of the original compound without water.

Perkin thinks that when water is added to

It is evident that the magnetic mol. rotatory powers of HCl, HBr, and HI in aqueous solutions are much greater than the values calculated from those obtained for the same three acids from observations made with compounds of them with C and H; in the case of HCl, the value of Mol. R. in aqueous solution is much

greater than the value for the same compound in solution in isoamyl oxide. There must be an action of some kind between water and these three compounds.

It has been shown already that addition of water to H₂SO₂ results in a value for the Mol. R. of the acid different from that observed for the pure acid itself; but in this case addition of water reduces the Mol. R. of the acid. A similar reducing effect was noticed as accompanying the addition of water to HNO₃.

Perkin was unable to explain these results. Ostwald, however, has shown that they are in keeping with the electrolytic dissociation hypothesis (C. J. 59, 198). According to this hypothesis strong acids are dissociated in aqueous solutions into their ions. Aqueous solutions of

HCl, HBr, and HI, contain the ions H and Cl,

+ - + -
H and Br, and H and I, each with its electric
charge. The observed rotatory powers of these
solutions, minus the value for the water contained in them, therefore represent the rotatory
powers of these ions, and not of the compounds
themselves; and the rotatory powers of these
ions are considerably greater than those of the
non-dissociated compounds. An aqueous solution of H₂SO₄, or HNO₂, must also contain dissociated, electrically charged, ions—according to
the hypothesis; but Perkin's observations
(ante, p. 259) show that the rotatory powers of
the ions of these acids are slightly smaller than
those of the acids themselves.

Now, the hypothesis of electrolytic dissociation asserts that the metallic salts of acids are dissociated into electrically charged ions in dilute aqueous solutions; hence determinations of Mol. R. for aqueous solutions of metallic salts of HOl, HBr, HI, H₂SO₄, and HNO₃ ought to give results different from the values obtained for these salts themselves. But the values obtained for aqueous solutions of salts of HCl, HBr, and HI ought, on the whole, to be greater than the calculated values, whereas the values obtained for salts of H₂SO₄ and HNO₃ ought, generally, to be slightly smaller than the calculated values. This deduction is partly confirmed by Perkin's results.

The values of Mol. R. for NH_4Cl is calculated by Perkin as follows (C. J. 55, 743). The value for HCl is found from determinations of Mol. R. of alcoholic chlorides, which enable values to be found for H and Cl in combination; the value for NH, is taken as 1.818, which is the mean of the values found from observations on NH3 in water and alcohol (ante, p. 259); the sum of the values for HCl and NH₃, with the subtraction of .5, gives the calculated Mol. R. of NH,Cl. The number '5 is taken from the sum of the values for HCl and NH₃ because observation has shown that this is about the difference between the values for NIII and Nv, and in NH, the N atom is trivalent, while in NH,Cl it is supposed to be pentavalent. The values for the other salts of NH, are calculated in a similar way. Perkin's results for Mol. R. of NH₄I, NH₄Br, and NH₄I show that practically the same values are obtained for solutions containing from c. 30 to c. 60 p.c. of the salts. The results with the six salts were as follows:

Calculated		Observed, in aqueous solutions	Differences		
NH ₄ Cl	4·305	6·096	+ 1·791		
NH ₄ Br	6·096	7·997	+ 1·901		
NH ₄ I	8·149	9·896	+ 1·747		
NH ₄ NO ₃	2·298	2·320	+ ·022		
NH ₄ HSO ₄	8·433	8·455	+ ·022		
(NH ₄) ₂ SO ₄	4·551	4·980	+ ·439		

With regard to NH, NO, and the two sulphates, it is to be observed that the calculated values are found by adding the values for the acid and the base (NH₃), then deducting 5 for the change of NIII to Nv, and then deducting 2 for the change from the free acid and base to the combination of these compounds; now, 5 is merely an approximate value for the change from NIII to Nv, and 2 is also merely a rough approximation for the value of the process of combination of acid and base. As the differences between the observed and calculated values for NH₄Cl, NH, Br, and NH, I are large, small errors in the method of calculating Mol. R. for these compounds do not materially affect the differences in question; but as the differences are small in the cases of the nitrate and sulphates, small errors in the method of calculating Mol. R. for these compounds do materially affect those differences. Moreover, the observations of Mol. R. of the nitrate and the sulphates of ammonium in aqueous solutions were made with a single solution in each case (59.7 p.c. NH, NO₃, 66.6 p.c. NH₄HSO₄, and 40 p.c. (NH₄)₂SO₄); hence it is not certain that the numbers set down as the values of Mol. R. of these compounds in aqueous solutions are the true values.

Perkin has also (C. J. 59, 981) made measurements of the magnetic mol. rotatory powers of the ammonium salts of formic, acetic, and propionic acids in aqueous solutions, and compared the results with the values calculated. He gives the results as follows:

	Observed, in aqueous solution	Calcu- lated	Dif- ferences
Ammonium formate ,, acetate ,, propionate	3·363	3·489	- ·126
	4·247	4·343	- ·096
	5·259	5·280	- ·021

The interpretation which the electrolytic dissociation hypothesis gives of these results is that, as in their aqueous solutions the salts examined are dissociated into their ions, and as the observed results are smaller than the calculated values, therefore the magnetic mol. rotations of the ions of HCO₂H, CH₂CO₂H, and C₂H₃CO₂H are slightly less than the rotations of the acids themselves (v. Ostwald, Z. P. C. 9, 513; cf. Ostwald, C. J. 59, 201).

But it seems to me that the numbers given by Perkin as the calculated values of Mol. R. for the three salts cannot be accepted as correct. For these values were found by adding the values of Mol. R. for the free acids to the value for free NH₃; no deduction was made for the change of rotation accompanying the combination of the

acid and the base, and no deduction was made for the change of NIII to NV. Either these values cannot be accepted as the true values, or the values calculated for the salts of NH, with the halogen acids, and with HNO₃ and H₂SO₄, are not correct (ante, p. 260). If the values of Mol. R. for the salts of formic acid &c. are calculated by the same method as was used in the cases of the other salts of ammonium, then the results of this investigation are as follows:

	Observed in aqueous solution	Calcu- lated	Dif- ferences
Ammonium formate ,, acetate ,, propionate	3·363	2·789	+ ·574
	4·247	3·643	+ ·604
	5·259	4·580	+ ·679

If these results are accepted, then the differences are much greater than those given by Perkin, and they are in the opposite direction.

Measurements of the Mol. R. of formic, acetic, and propionic acid in aqueous solution would enable the question to be settled as to the effect of water on these acids; at present it seems that the magnetic rotations of the ions of these acids would be found to be greater than the rotations of the acids themselves.

This conclusion is confirmed by combining the results obtained by Perkin (C. J. 59, 986) for Mol. R. of sodium salts of formic and other acids with the value calculated by him for sodium (C. J. Proc. 1890. 141). In the following table I have calculated the values for the salts as Perkin calculated the values for NH₄Cl, NH₄NO₄ &c., by deducting 2 from the sum of Mol. R. of acid and At. R. of Na, and then further deducting .254 for the loss of H:

	Observed in aque- ous solu- tion	Calcu- lated	Differ- ences
Sodium formate , acetate , propionate	2·347 3·281 4·308	1.775 2.629 3.566	+·572 +·652 +·742
" butyrate	5.332	4.576	+ .756

In a more recent communication (C. J. 63, 57)Perkin gives the Mol. R.s of aqueous solutions of H₂SO₄, HNO₅, Na₂SO₄, and LiNO₅ of varying degrees of concentration. His results show that the Mol. R.s of these solutions do not vary with dilution in the same way as the electrical conductivities vary. The most dilute solution of H₂SO₄ examined by P. contained c. 9.2 p.c. H₂SO₄, and the most dilute solution of HNO₃ contained c. 22.5 p.c. HNO₃. Now it has been shown repeatedly, by Ostwald and others, that measurements of the conductivities of fairly conc. solutions of acids and salts cannot lead to just conclusions regarding the amount of electrolytic dissociation of the compounds in solution; hence it is not to be expected that the amount of dissociation of H,SO, and HNO, should be calculable from the Mol. R.s of these compounds as deduced from observations on solutions so concentrated as those used by Perkin. What the observations of P. show is that the magnetic rotations of certain compounds diverge from the calcu-

lated results in all cases wherein electrolytic dissociation occurs. The laws which express the electric dissociation of compounds are to be deduced from the study of very dilute solutions. and as no measurements of magnetic rotations have yet been made for such solutions, it is not justifiable to draw detailed quantitative conclusions concerning the electrolytic dissociations of compounds from the measurements of the magnetic rotations of those compounds which P. M. M. P. M. has made.

VIII. OSMOTIC PRESSURE, METHODS BASED ON; v. MOLECULAR WEIGHTS, vol. iii. p. 418; and Electrical methods, this vol. p. 185.

IX. PHOTOGRAPHIC METHODS: PHOTOGRAPHIC CHEMISTRY, this vol. p. 154. HEATS X. SPECIFIC OF

METHODS BASED ON; v. ATOMIC AND MOLE. CULAR WEIGHTS, vol. i. p. 342.

XI. THERMAL METHODS. Determinations of the quantities of heat which are produced, or which disappear, in chemical reactions help to elucidate the nature of these reactions. There are two sides to every chemical operation: there is a change in the form or distribution of matter, and a change in the form or distribution of energy. Light is thrown on the second of these changes by thermo-chemical investigations, and the connections between the two parts of the complete occurrence, and the mutual dependence of both, are thus made clearer. Thermo-chemical methods rest on the principle of the conservation of energy. When a chemical reaction occurs, heat may be produced, electricity generated, mechanical work done by expansion, and sound or radiant heat may leave the system; the sum of these forms of energy, added to the energy remaining in the system at the close of the reaction, must equal the energy originally present in the system at the moment when the change began. As a large part of the energy set free during any chemical reaction usually takes the form of heat, it follows that measurements of the heat produced during the reaction must give information regarding the differences between the initial and final energies of the system. The primary aim of thermo-chemical measurements is to determine such energy-differences; the secondary aim is to connect these differences with the differences of composition, constitution, and properties which the reacting systems undergo.

The loss or gain of energy which accompanies the passage of a system from one defined state to another is independent of any intermediate states through which the system may This generalisation was proved experimentally for several cases, so far as heat energy was concerned, by Hess in 1840 (P. 50, 385); the generalisation may be deduced from the On this statement rest principles of energy. the methods for determining the thermal values of chemical changes when these values cannot be observed directly.

Notation used in thermo-chemistry. Thom sen's notation is followed in this Dictionary; the formulæ of the reacting substances are taken to represent grams of these substances-thus HCl means 36.5 g. of hydrogen chloride; the formulæ are inclosed in a square bracket, and the figures representing the number of atoms of each ele-

ment are placed above the symbols; the reacting substances are separated by a comma. The unit of heat is that quantity which raises the temperature of 1 g. of water at c. 18° through 1°. Ostwald, in his Lehrbuch der Allgemeinen Chemie, employs a unit almost exactly 100 times greater than this, viz. the quantity of heat given out by 1 g. of water in cooling from 100° to 0°. The symbol Aq means that a large quantity of water is present. Thus [H,Cl] = 22,000 means that 22,000 gram-units of heat are produced when 1 g. H combines with 35.5 g. Cl; [H,Cl,Aq] = 30,660 means that 30,660 gramunits of heat are produced when 1 g. of H combines with 35.5 g. of Cl in presence of a large quantity of water in which the HCl dissolves; [HCl,Aq] = 17,320 means that 17,320 gramunits of heat are produced when 36.5 g. of HCl dissolve in a large quantity of water; and [HClAq,KOHAq] = 13,750 means that the neutralisation of 36 5 g. HCl, dissolved in much water, by 56 g. KOH, dissolved in much water, the products of neutralisation being allowed to remain in solution, is accompanied by the pro-The duction of 13,750 gram-units of heat. symbol H,O is used as in ordinary notation to represent 18 g. of water; thus [As2O3,3H2O] =6,800 means that 6,800 gram-units of heat are produced during the formation of 284 g. H. AsO. by the combination of 230 g. As, O, with 54 g. water; and $[As^2O^5.3H_2O,Aq] = -800$ means that 800 gram-units of heat disappear during the solution of 284 g. H₃AsO₄ in a large quantity of Thomsen's notation does not indicate the products of the reaction, the thermal value of which is set down; nor does it show the physical states of the reacting substances, or of the products of the reaction, except in the one case when the substances are dissolved in much water. Ostwald (Lehrbuch) uses the ordinary notation, and adds figures to express the quantities of heat which are produced or disappear in the reactions; he employs ordinary type for liquids, thick type for solids, and italics for gases. Thus, $H_2 + Cl_2 = 2HCl + 44{,}000$ means that 44,000 gram-units of heat are produced when 2 g. gaseous hydrogen combine with 2×35.5 g. gaseous chlorine to form 2×36.5 g. gaseous hydrogen chloride; $2H_2S + 2I_2$ = 4HI + 2S - 34,000 means that 34,000 gram- $2H_2S + 2I_2$ units of heat disappear when 2 × 34 g. gaseous hydrogen sulphide react with 2×254 g. solid iodine to form 4 × 128 g. gaseous hydrogen iodide and 2 × 32g. solid sulphur; and 2H2SAq + 2I2Aq =4HIAq + 2S + 34,000 means that 34,000 gramunits of heat are produced when 2×34 g. hydrogen sulphide dissolved in much water react with 2×254 g. iodine dissolved in much water, to form a dilute aqueous solution of 4×128 g. hydrogen iodide, and 2×32 g. solid sulphur. Ostwald sometimes indicates the temperature by figures in brackets placed after the formulæ of the substances: thus, $H_2O_{(0)} = H_2O_{(0)} + 1440$ means that 1440 gram-units of heat are produced when 18 g.

liquid water at 0° become 18 g. solid water at the same temperature.

Measurements of thermal values of chemical changes. The quantities of heat produced during chemical processes are measured by causing the processes to take place in vessels arranged so that the whole of the heat is used in raising the temperature of known masses of water, or of water and a solution the specific heat of which is known. When the process consists in a reaction between substances in aqueous solutions, the calorimeter employed usually consists of a vessel of platinum holding 500 to 1000 c c. placed inside another vessel of silver, or thin sheet iron, with water between the two vessels. The reacting liquids are brought to the same temperature, and are then mixed in the platinum vessel, and the rise of temperature of the contents of this vessel is noted, any change in the temperature of the outside water being also measured. When the thermal value of a process of combustion is to be determined, the combustion is caused to proceed in a vessel, usually made of platinum, surrounded by a determinate quantity of water; if the products of combustion are gases, an apparatusis attached wherein these gases are absorbed, and thus the quantity of substance burnt is determined.1

To calculate the thermal value of a reaction. it is necessary to know the initial temperatures of the reacting bodies, the final temperatures of the products of reaction, the masses of the reacting substances, the specific heat of water, the specific heat of the liquid formed by the reaction if the reaction be one wherein a solution is produced, and the water-equivalent of the calorimeter. The water-equivalent of the calorimeter must be determined; it is equal to the number of unitweights of water which would be raised to the same temperature as that to which the calorimeter and its accessories-thermometer, stirrer, &c.—are raised by the heat produced in the process. When dilute solutions of acids and alkalis, or similar compounds, react, the specific heat of the dilute salt-solutions produced may be taken as equal to the specific heat of the water they contain, without appreciable errors.

Let Q be the quantity of heat produced in a reaction between two solutions; let t be the initial temperature of one solution, t' the initial temperature of the other solution, and T the final temperature of the solution produced; further, let a be the calorimetric equivalent of one solution, i.e. mass \times spec. heat, b the calorimetric equivalent of the other solution, and c the water-equivalent of the calorimeter; then

 $\mathbf{Q} = (\mathbf{T} - t)a + (\mathbf{T} - t')(b + c).$ In the cases of dilute aqueous solutions, a and b represent the masses of water in the solutions mixed. If a change occurs in the state of aggregation of some of the constituents of the reacting system, e.g. if liquid water is formed from gaseous hydrogen and oxygen, the quantity of heat which is produced or disappears in this change must be taken into account in the calculation of the thermal value of the reaction. If contraction or expansion occurs, without change of state, the thermal value of such contraction or expansion must be determined and allowed for.2

Heats of combustion, and of formation, of compounds. The heat of combustion of an element or compound is the thermal value of the reaction which takes place when that element or com-

ratus v. References, p. 268.

For methods of calculating Q at one temperature when it is known at another temperature, v. 7h. 1, 69-78.

^{*} For references to descriptions of calorimetrical appa-

pound is completely oxidised to those masses of the product, or products, of oxidation which are represented by the formulæ of these products. Thus $[H^2,O] = 68,360$; [CO,O] = 67,960. In this Dic-TIONARY the quantities represented by formulæ are taken in grams.

The heat of formation of a compound is the thermal value of the reaction whereby the formula-weight of the compound is produced from the formula-weights of its constituents. Thus, the heat of formation of H₂SO, varies according to the constituents from which it is produced; we have the following statements: [SO',H-O] =21,320; $[SO^2,O,H^2O] = 53,480$; $[SO^2,O^2,H^2]$ **121,840**; $[S,O^3,H^2O] = 124,560$; $[S,O^4,H^2]$ = 192.920.

It is often practicable to measure heats of combustion directly; but in very many cases it

is necessary to determine heats of formation by indirect methods. These indirect methods rest on the principle, deduced from the laws of energy, that the total change of energy, and therefore the total thermal change, which accompanies the passage of any system from a definite initial composition to a definite final composition is independent of the intermediate states. Suppose that a system changes from a certain arrangement or configuration A to another configuration B, and that x units of heat are produced in this change. Suppose also that the same initial system then passes from the state A to a new state a, from that to b, from that to c, and, lastly, from that to B, and that there are x' units of heat produced in the first of those intermediate changes, x" units produced in the second, and x" units in the third; then x - (x' + x'' + x''') gives the thermal value of the change from the state c to the state B. If then the thermal value of a chemical process, whether the formation of a compound, or other process, cannot be determined by direct measurement, it may be determined if the process can be made a portion of a series of changes, the total thermal value of which is measurable, and the thermal values of all the portions of which are measurable with the exception of the value of that portion which is sought.

For instance, formic acid, CH₂O₂, cannot be directly produced from C, H, and O, and, therefore, the heat of formation of this acid cannot be determined by direct measurement; but C and H can be burnt to CO2 and H2O, and CH2O2 can also be burnt to CO₂ and H₂O; hence we can acquire the data needed for calculating the value of $[C,H^2,O^2]$. The data are these: $[C,O^2] = 96,960$; $[H^2,O] = 68,360$; sum = 165,320; $[CH^2O^2,O] = 65,900$. Now, suppose that the first stage of the combustion of C and H2 consists in the formation of CH₂O₂, and that this is then burnt to CO, and H2O, we should have the statement: $[C,O^2] \cdot [H^2,O] = [C,H^2,O^2] + [CH^2,O^2,O] = 165,320$. But $[CH^2,O^2,O] = 65,900$; and also $[C,O^2] + [H^2,O] - [CH^2,O^2,O] = [C,H^2,O^2]$. Hence Hence $[C,H^2,O^2] = 165,320 - 65,900 = 99,420.$

Another example will show the application of the principle to a more complicated case. It is required to find the heat of formation of gaseous hydrogen bromide from gaseous H and Br, i.e. the thermal value of the reaction H + Br = HBr.

To begin with: let $[H,Br,Aq] = \infty$, and let [H,Br]=x'; direct measurement gives [HBr,Aq] = 19,900. x-19,900=x'= heat of formation of HBr. Hence it is necessary to determine x.

Now, the thermal values of the neutralisation of HClAq and HBrAq respectively by KOHAq are found to be the same, i.e. [KOHAq, HClAq] =[KOHAq,HBrAq]. If these reactions are analysed, the first is seen to consist in (1) the splitting of HClAq into H and Cl in presence of water; (2) the separation of KOHAq into K, O, and H in presence of water; (3) the combination of K and Cl, in presence of water, to form KClAq; (4) the combination of H, H, and O, in presence of water, to form water; and the second is similar to this, only putting Br in place of Cl. These reactions may be stated in thermo-chemical notation thus:

(1) [KOHAq,HClAq] = -[K,O,H,Aq] - [H,Cl,Aq] + [K,Cl,Aq] + [H,H,O,Aq];(2) [KOHAq, HBrAq] = -[K, O, H, Aq] - [H, Br, Aq] + [K, Br, Aq] + [H, H, O, Aq].

> The first and fourth terms on the right side are the same, and the total thermal values are the same. Are the values of the second and third terms the same? To answer this question we pass Cl into KBrAq, forming KClAq and BrAq, and measure the thermal disturbance; we then analyse the reaction, and see what information it has afforded. These are the results:

> [KBrAq,Cl] = 11,500,i.e. -[K,Br,Aq]+[K,Cl,Aq]+[Br,Aq]=11,500. Now, [Br,Aq]=500, by direct measurement, [K,Br,Aq] + [K,Cl,Aq] = 11,000, i.e. to form KCl, in solution, from K and Cl produces 11,000 units of heat more than to form KBr, in solution, from K and Br.

> Now, turning back to equations (1) and (2) above, and remembering that the thermal values of these are the same, it is evident that, since 11,000 more heat-units are produced in forming KClAq than in forming KBrAq, each from its elements, 11,000 more heat-units must disappear in splitting up HCl, in presence of water into H and Cl, in presence of water, than in similarly separating HBr into H and Br in presence of water; and therefore, since the heat energy required to decompose a stated mass of a compound is equal to the heat energy that is produced when the same mass of that compound is formed, 11,000 more heat-units will be produced in the formation of HClAq from H, Cl, and Aq than in the formation of HBrAq from H, Br, and Aq; or, stated in thermo-chemical notation: [H,Cl,Aq]-11,000=[H,Br,Aq]. Now [H,Cl,Aq]=39,300 by direct measurement, [H,Br,Aq] = 89,300 - 11,000 = 28,300.

> At the beginning of this calculation we had [H,Br,Aq]-19,900=[H,Br]. We can now substitute the value of [H,Br,Aq], and write [H,Br] = 28,300 - 19,900 = 8,400. That is to say, 8,400 gram-units of heat are produced when 81 g. gaseous hydrogen bromide are formed from 1 g. gaseous hydrogen and 80 g. gaseous bromine.

> Exothermic and endothermic reactions. Chemical changes which are accompanied by production of heat are sometimes classed together as exothermic reactions, and are distinguished from changes accompanied by disappearance of heat, which changes are called endothermic reactions. The terms endothermic and exothermic are sometimes useful. It should not be forgotten

that most chemical reactions consist of portions which are exothermic and portions which are endothermic. Substances which are formed with the disappearance of heat are generally more readily decomposed by the application of outside forces than substances which are formed with the production of heat.

Interpretation of thermo-chemical measure-nts. Thermo-chemical measurements aim at determining the quantities of heat which are produced or disappear during definite and defined chemical changes. But every chemical change is inextricably bound up with more or less extensive physical changes; hence some portion of the thermal value of a chemical occurrence is always due to a process which is not, strictly speaking, chemical. A purely chemical change is a change in the distributions, configurations, and motions of atoms. But in only some cases are we able to form clear conceptions as to the configurations and motions of atoms; therefore, even if it were possible always to disentangle the purely chemical from the accompanying physical parts of a change, we should still very frequently be unable to connect the thermal values of purely chemical processes, in a clear and definite way, with measurable changes in the distributions, configurations, and motions of atoms.

The heats of formation of HCl, HBr, and HI are said to be 22,000, 8,440, and -6,050 gramunits respectively. But these thermal values are not strictly comparable, because the first represents the heat produced in forming gaseous HCl from gaseous H and Cl; the second represents the heat produced in forming gaseous HBr from gaseous H and liquid Br, and the third represents the heat which disappears when gaseous HI is formed from gaseous H and solid I. In other words, more heat is used in the second change than in the first, and still more in the third than in the second, in accomplishing subsidiary physical changes. Steam is decomposed by chlorine, with formation of hydrogen chloride and oxygen. All the reacting substances are gases. This change, which is stated in formulæ as $2H_2O + 2Cl_2 = 4HCl + O_2$, when analysed thermo-chemically, is found to consist of four parts, thus: [2H2O,2Cl2]

= -2[H,H,O] - 2[Cl,Cl] + [O,O] + 4[H,Cl]. We measure the thermal value of this complete change, but we cannot at present separate the portions of the change and assign to each its proper thermal equivalent; hence we cannot give a complete explanation of the thermochemical measurement we have made.

Notwithstanding these difficulties, attempts have been made to generalise from thermochemical measurements to statements of universal applicability. The most widely known of such attempts is that which finds expression in Berthelot's 'law of maximum work.' This so-called law asserts that 'Every chemical change accomplished without the addition of energy from without tends to the formation of that body, or system of bodies, the production of which is accompanied by the development of the maximum quantity of heat.' In another place Berthelot states the law more rigidly, thus: 'Every chemical change which can be accomplished without

the aid of a preliminary action, or the addition of energy from without the system, necessarily occurs if it is accompanied by disengagement of heat.'

The same generalisation was stated by Thomsen (v. Th. 1, 12-16) some years before it was enunciated by Berthelot. Thomsen's statement took this form: 'Every simple or complex reaction of a purely chemical kind is accompanied by production of heat.' None of these statements is strictly applicable to actually occurring chemical changes, because we never have to deal with reactions of a purely chemical kind, but with reactions that are partly chemical and partly physical-that is, with reactions which consist in part in changes in the arrangement and motions of atoms, and partly in changes in the arrangement and motions of molecules. consideration of the way in which the so-called law is applied, especially by Berthelot and his school, shows that it is taken to mean that measurements of the thermal values of various possible chemical changes enable us to predict which of these will occur. Thus, suppose we start with a system A+B+C, and suppose that from this may be formed AB+C, AC+B, or ABC; suppose also that x thermal units are produced in the first of these changes, x' thermal units in the second, and x'' thermal units in the third; finally, suppose that x'' > x' > x; then the law of maximum work is taken as asserting that the reaction A+B+C=ABC will occur, and will occur to the complete, or almost complete, exclusion of the two other possible reactions.

It is easy to show that many reactions are known to occur which would be impossible were this 'law 'a true generalisation. But it is better to treat the law as a deduction from the principles of energy, and to show that it is not a warrantable deduction.

The statements made by Thomsen and Berthelot are true only when an arbitrary separation is made of chemical changes into two parts, and one of these parts is alone called chemical. Every chemical change, however simple, consists of at least two parts, the first of which is the necessary antecedent of the second; the 'law of maximum work' ignores this duality, or, it might be more accurate to say, the law assumes that the second part of a chemical process may occur without the first. A process of chemical change may be compared to the flight of a stone from, and its return to, the surface of the earth. During the first part of this process there is a continual transference of kinetic energy from the moving stone to the surrounding medium and during the second part there is a continual transference from the medium to the stone, until the stone comes to rest, when its energy becomes a part of the total energy of the system earth plus stone. If the final resting-place of the stone is nearer the centre of the earth than the spot from which it was projected in its upward flight, then the stone contains less energy, relatively to surrounding systems, at the close of the transaction than at the beginning. On the other hand, if the starting-point is nearer the earth's centre than the final point of rest, then the transaction has resulted in a gain of energy to the stone. In both cases the second part of the transaction, that which occurs between the turning point and the coming to rest of the stone, is attended with loss of energy to the stone; but this second part does not represent the complete transaction. The 'law of maximum work,' if applicable at all, is applicable only to the second part. And, moreover, this law ignores the fact that the stone, or chemical system, does not leave its initial resting-place of its own accord; the law assumes that no work need be done, no energy need be expended, in the passage of the stone, or chemical system, from its original position to that whereat the energy-relations between it and surrounding systems come within the cognisance of the law.

The 'law of maximum work' asserts a certain condition of equilibrium for a chemical system, for the law states that equilibrium results when that change has occurred which is attended with the maximum production of heat. But the true condition of equilibrium of a material system undergoing a reversible change is laid down in the statement that equilibrium results when the entropy of the system has attained the maximum value under the conditions which prevail.

Let Q = quantity of heat added to a body at constant temperature T, then Q = gain of entropy to the body; let $Q_1 = \text{quantity}$ of heat lost by a body at constant temperature T_1 , then $Q_1 = \text{loss}$ of entropy to the body. All chemical and physical changes which occur spontaneously increase the entropy of the system. This statement holds good for non-reversible changes; and as no actually occurring change is completely reversible, the statement holds for all changes. Suppose that one of two bodies is hotter than

the other and loses heat to the colder body; the hotter body at temperature T_1 loses heat Q, therefore its entropy is diminished by $\frac{Q}{T_1}$; the colder body at temperature T_2 gains heat Q, therefore its entropy is increased by $\frac{Q}{T_2}$; but as $T_1 > T_2$, it follows that $\frac{Q}{T_1} < \frac{Q}{T_2}$; in other words, the entropy of the system is increased by the passage of heat from the hotter to the colder body.

Now, a system is in equilibrium when its entropy has attained the maximum value possible under the conditions. But, inasmuch as entropy is measured by a quantity of heat divided by a temperature, it is only at the absolute zero of temperature that dS = dQ (S = entropy, Q = quantity of heat); hence it is only at the absolute zero that thermal changes directly measure changes of entropy. When a chemical change is accompanied by the production of much heat, and the change occurs at a low temperature, the thermal change will roughly measure the change of entropy; therefore, if such a change be possible, it will occur. But if the quantity of heat produced in a chemical process is small, the change of entropy may be conditioned, to a large extent, by changes other than the thermal change. Indeed, in some cases, heat may disappear from the system, and yet the total change of entropy may be positive; in such cases, chemical change will occur with the disappear-

ance of heat, because the decrease in the entropy of the system caused by the loss of heat will be more than balanced by the increase in the entropy caused by the changes of state which the system undergoes.

The general conception of chemical change which is given by applying the law of entropy is that of a system attaining equilibrium as the result of processes taking place in opposite directions. According to van't Hoff (Dynamique Chimique, 153), the directions of chemical processes which result in equilibrium vary with variations of temperature in such a way that the lower the temperature the more is equilibrium established with the production of heat, but the changes can take place wholly in one direction only at the absolute zero. The 'law of maximum work' would then hold good for the limiting case that the change should occur at -273° As the temperatures at which most chemical changes occur are not very high, many chemical processes are accompanied by production of heat.

The 'law of maximum work' is not, then, a law of nature. But it is true that most chemical processes which occur without the expenditure of much energy from without, and which take place at moderate temperatures, are accompanied by the disengagement of heat. If it is known that a certain chemical change would be attended by the disappearance of much heat, we may conclude that this change will be difficult to accomplish; that it will be brought about only by the expenditure of a considerable quantity of energy, and that the most successful way of accomplishing the change will be to make it one part of a series of changes the sum of which is attended with the disengagement of heat.

Applications of thermo-chemical methods. The applications of thermo-chemical methods are many and varied. A few of the more important will be briefly dealt with here.

Neutralisation of acids and bases. In 1842 Hess stated the principle of the thermo-neutrality of salts in solutions (P.52,79). He said that when aqueous solutions of two normal salts are mixed the thermal disturbance is nil; in other words, that the thermal value of the reactions consisting in the exchange of the acids and bases is equal to zero. Hess supposed that the heats of neutralisation of acids were independent of the nature of the bases used. Andrews (P. 54, 208; 59, 428) thought that the heats of neutralisation were dependent only on the bases, and were independent of the nature of the acid employed. Favre a. Silbermann (A. Ch. [3] 34, 357; 36, 1; 37, 406) put the law of thermo-neutrality in its proper form by showing that the differences between the heats of neutralisation of any two bases by any acid have a constant value, and the differences between the heats of neutralisation of two acids by any base are constant. Let the composition of various salts be represented by the scheme:

A' + B $\Lambda^{\prime\prime} + \mathbf{B}$ A''' + BA + BA' + B'A'' + B'A"' + B' A + B'A'' + B''A''' + B''A + B''A' + B''A + B" A' + B'''A'' + B'''And let f(A+B), f(A'+B), &c., represent the quantities of heat produced by the union of the acid A with the base B, the acid A' with the base B, &c. Then the law of thermo-neutrality asserts that

f(A+B) + f(A'+B') - f(A+B') - f(A'+B) = 0or f(A+B) - f(A+B') = f(A'+B) - f(A'+B')and f(A+B) - f(A'+B) = f(A+B') - f(A'+B')

Exceptions have been found to this law, but these exceptions have all proved to be connected with some abnormality in the behaviour of the salts formed. The law may be stated by saying that, in the normal formation of a salt, in solution, by the reaction between an acid and a base, the acid contributes a definite portion of the total heat of neutralisation, independently of the nature of the base, and the base contributes a definite portion of the total heat of neutralisation, independently of the nature of the acid. If the statement is correct, then the heat of neutralisation of an acid, in solution, must be independent of the nature of the base, and the heat of neutralisation of a base, in solution, must be independent of the nature of the acid, unless there be some divergence from the normal condition of affairs. By the heat of neutralisation of an acid is meant the quantity of heat produced when an equivalent weight of the acid reacts with an equivalent weight of a base, both being in dilute aqueous solution. If A' represents a formula-weight of a monobasic acid, A" a formula-weight of a dibasic, and A" a formulaweight of a tribasic, acid; and if B represents a formula-weight of a mono-acid base; then the heats of neutralisation of these three acids are the thermal values of the reactions,

[A'Aq,BAq]; [½A''Aq,BAq]; and [¼A''Aq,B\q]. Sometimes it is more convenient to take the thermal values of the reactions [A''Aq,2BAq] and [A'''Aq,3BAq] to represent the heats of neutralisation of the dibasic and tribasic acids respectively.

The following table, taken chiefly from Thomsen's results, shows that the heats of neutralisation of the strong acids are practically independent of the base, provided the base used is strong; the numbers all refer to reactions between dilute solutions of the acids and bases:—

Acid	NH,	NMe,	ин,он		
HCl HNO ₃	12,200 12,300	8,700	9,200		

When weak acids neutralise strong bases, the thermal values differ from those obtained for strong acids; and when weak acids neutralise weak bases the reactions have different values from any of the preceding cases. The following numbers illustrate this:

	NaOH	NH,	
Acid	strong	weak base	
$\overset{\mathbf{H}}{\overset{\mathbf{G}}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}}}\overset{\mathbf{G}}{\overset{\mathbf{G}}}}{\overset{\mathbf{G}}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}}{\overset{\mathbf{G}}}{\overset{\mathbf{G}}}{\overset{\mathbf{G}}}}{\overset{\mathbf{G}}}{\overset{\mathbf{G}}}{\overset{\mathbf{G}}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}}{\overset{\mathbf{G}}}}{\overset{\mathbf{G}}}}}{\overset{\mathbf{G}}}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}}}}{\overset{\mathbf{G}}}}}}{\overset{\mathbf{G}}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}{\overset{\mathbf{G}}}{\overset$	10,100 7,700	10,900 7,800	8,400 6,200

The electrolytic dissociation hypothesis gives an explanation of the facts concerning the thermo-chemical reactions between acids and bases. According to this hypothesis, a dilute aqueous solution of a salt contains the ions of the salt each with its electric charge; the salt is dissociated into its ions; when two dilute salt solutions are mixed, the ions remain as they were, there is no change, and therefore heat is neither produced nor consumed. The law of thermo-neutrality holds. The hypothesis looks on a dilute aqueous solution of a strong acid, or a strong base, as containing the ions of the acic or the ions of the base; when the solutions of the strong acid and strong base are mixed, a salt is not formed in the solution, because in dilute solutions salts are wholly dissociated, but water is formed by the union of the ion H of the acid with the ion OH of the base. Thus, the com position of dilute solutions of HCl and NaOH before and after mixing, are represented by the hypothesis as follows:

$$\frac{\overset{+}{\text{Na}} + \overset{+}{\text{OH}} + \overset{+}{\text{H}} + \overset{-}{\text{Cl}}}{\text{hofore mixing}} \xrightarrow{\text{after mixing}} \overset{+}{\text{after mixing}}$$

Acid	NaOH	кон	LiOH	тон	BaO,H	½CaO,H.	½Sr0, H ,	Pt(NH _s),(OH)	S(C,H,),OH	N(CH,),OH
HCI	13,300	13,300	13,300	13,300	18,300	13,300	13,300	13,600	13,700	13,600
HBr	13,300	18,300	_			<u> </u>			_	-
HI	13,300	13,300		_			- 1			
HNO.	13,500	13,500		18,500	14,000	13.900	l _			
HClO.	13,600	13,600			14,000					
HBrO.	13,600	13,600			_			·		_
HIO.	13,600	13,600	-		_					
HCIŌ.	14.000	14.000			-	_				
H.S.O.	13,500	_			13,800					
H.PtCl.	13,600									-
n.c.H.so.	13,500				13,600					_

If concentrated solutions are employed, the heats of neutralisation, even of the strong acids, show considerable differences with different bases. If the salt which is formed by neutralising an acid by a base is allowed to precipitate, then the apparent heat of neutralisation as thus determined is not the true heat of neutralisation (v. Th. 1, 440). The following numbers represent the heats of neutralisation of two strong acids by weak bases; these numbers differ from one another, and also from the values given in the preceding tables in which strong acids and strong bases only were included:

Or, generally, if M represent the positive ion of a strong base, and A the negative ion of a strong acid; we have the composition of the dilute aqueous solutions, before and after mixing, represented by the two schemes:

$$M + OH + H + A$$
 and $M + A + HOH$
before mixing after mixing

Hence, as the only process wherein heat car be produced consists in all cases in the union of H with OH, the quantity of heat produced is always the same. The hypothesis asserts that HOH is produced because water is a non-dissociable, or nearly a non-dissociable, compound; pure water

is, approximately, a non-electrolyte.

But when a weak acid is mixed, in dilute aqueous solution, with a strong base, or when dilute solutions of a weak acid and a weak base are mixed, the processes are more complex; for according to the hypothesis, only a portion of the weak acid is dissociated into ions, and only a portion of the base is dissociated if the base is weak. In such cases the state of matters, before and after mixing, would be represented thus (supposing both acid and base to be weak)—

But the process does not necessarily stop here; there is a striving towards the formation of the comparatively non-electrolytic compound HOH, therefore more of the acid and more of the base become dissociated, so that we have, as a final condition, less or more of the acid HA, and less or more of the base MOH, remaining in the solution. The dissociation of the acid and the base, when these are mixed, is accompanied by a thermal change; and as the degree to which this dissociation proceeds depends on the nature of the acid and the base, the values of the heats of neutralisation of weak acids and weak bases differ one from another (cf. Electrical Methods, p. 189 in this vol.).

If this treatment of the heats of neutralisation of acids and bases is accepted, it is evident that the thermal values of the neutralisations of acids by a base do not necessarily measure the affinities of these acids for that base. If the acids were all equally dissociated in solution, they would all be equally strongor their affinities would be equal-and their heats of neutralisation would be equal. The statement Q = about 13,500 + A + B expresses the heat of neutralisation of any acid by any base in dilute aqueous solution. A represents the heat of dissociation of the acid, and B is the heat of dissociation of the base, into their ions; A and B may be positive or negative at any specified temperature. Under the conditions of the experiment only a portion of the acid, or of the base, may be dissociated; therefore the observed heat of neutralisation cannot measure the affinity of the acid for the base. But at the same time, it is evident that there is a connection between the heats of neutralisation and the affinities of acids and bases. On the one hand, measurements of the thermal values of the reactions between acids and bases enable conclusions to be drawn as to the distribution of two acids between one base, or two bases between one acid, and such measurements therefore lead to determinations of the affinities of acids and bases (v. infra); on the other hand, although the affinity of an acid for a base probably consists in a striving towards electrical equilibrium among the ions, yet, as electrical and thermal phenomena are closely connected, measurements of the heats of dissociation of acids and bases, in aqueous solutions, must help us to understand the relative affinities of acids and bases, to classify acids and bases in accordance with their affinities, and to connect the affinities of these substances with their constitution (cf. ELECTRICAL METHODS, especially pp. 208 and 209).

METHODS, especially pp. 208 and 209).

Monobasic and polybasic acids. When an equivalent of a monobasic acid is added to an equivalent of a base, both in dilute solutions, a certain quantity of heat is produced, and there is no further thermal disturbance on adding more of the acid to the neutral solution; but heat is either produced or disappears when a polybasic acid is added to a solution of the same acid which has been neutralised by an equivalent quantity of a base. This behaviour enables a thermo-chemical distinction to be drawn between monobasic and polybasic acids (cf. Acids, Basicity of, vol. i. p. 51).

Distribution of an acid between two bases. Thomsen found the following data:

[H²SO⁴Aq,2NaOHAq] = 31,380; [2HNO³Aq,2NaOHAq] = 27,230; [Na²SO⁴Aq,2HNO₃Aq] = -3,500.

Now, supposing that the reaction of equivalent quantities of Na2SO4 and HNO4, in dilute solution, produced NaNO₃ and H₂SO₄, and these compounds only, this reaction would be attended by the disappearance of 31,380 - 27,230 = 4,150thermal units; but as only 3,500 thermal units disappear in the reaction, it is evident that some change has occurred wherein heat has been produced, or that the whole of the Na2SO, has not been changed to NaNO, by the reaction of the equivalent quantity of HNO. If we suppose that the discrepancy in the quantity of heat consumed is due to the partial decomposition of the Na, SO, by the HNO,; in other words, if we suppose that when equivalent quantities of Na SO, and HNO, react in solution, some of the base remains united with the H.SO, and some goes into combination with the HNO3, and that no other reaction occurs, it is easy to calculate the distribution of the base between the two acids. For it is evident that, on this supposition, 3500 \$\frac{355}{4150} = 845 of the quantity of Na₂SO₄ present

has been decomposed. Thomsen found that H₂SO₄ reacts with Na₂SO₄ to form Na₂HSO₄, with disappearance of heat. He also found that this reaction is expressed thermo-chemically by the statement [nH²SO⁴Aq,Na²SO⁴Aq]

=31,380 $-\frac{n}{n+8}$ 3,300. These data give a means for calculating the distribution of the base between the two acids. The complete reaction between equivalent quantities of Na,SO₄ and HNO₃ will consist of three parts: (1) the decomposition of a equivalents of Na,SO₁, attended with the disappearance of a. 31,380 thermal units; (2) the formation of a equivalents of NaNO₃, attended with the production of a. 27,230 units of heat; (3) the reaction of a equivalents of H₂SO₄ with 1-a equivalents of Na₂SO₄ attended with the disappearance of

$$1-a\frac{1-a}{a+8}$$
 3,300 units of heat.

The complete thermo-chemical change may be stated thus:

 $[Na^2SO^4Aq, 2HNO^3Aq] = -3,500$ -3,300. =a(27,230-31,380)-(1-a)

<u>a</u> + ·8

Thomsen found that if a is taken as $\frac{2}{3}$, the number - 3550 is obtained, which is almost identical with the observed value. Hence Thomsen concluded that $\frac{2}{3}$ of the base, NaOH, went to the nitric acid, and $\frac{1}{3}$ to the sulphuric acid; or, that the affinity of nitric acid for soda (Thomsen used the term avidity) is twice as great as the affinity of sulphuric acid for the same base.

The relative affinities of various acids for different bases have been measured by Thomsen by this method (v. Affinity, vol. i. pp. 74-75).

Allotropy and isomerism. The generally accepted views regarding the allotropy of elements and the isomerism of compounds would lead us to expect that the formation of one allotropic form of an element from another form, or the formation of one isomeride from another, should be accompanied by changes of energy, and therefore, probably by the production or disappearance of heat. This expectation is confirmed by thermo-chemical measurements. The following tables present some typical data:—

Combustion of allotropic forms of sulphur, phosphorus, and carbon.

 $[S,O^{2}]$ to form gaseous SO. 71,220 S crystallised from CS2 for 71,720 rhombic S 72,300 native opaque S •• [P2,O5] to form solid P.O. 369,900 ordinary P for 362,800 red P [C,O] to form gaseous CO. 96,900 for amorphous C

diamond Combustion of isomeric carbon compounds.

93,200

 $[C^{0}H^{6},O^{15}]$ to form $6CO_{2} + 3H_{2}O$ 787,900 for benzene 883,200 dipropargyl $[C^{2}H^{6}O,O^{6}]$ to form $2CO_{2} + 3H_{2}O$ 330,400 ethylic alcohol for 344,200 methylic ether ,,

 $[C^3H^6O,O^8]$ to form $3CO_2 + 3H_2O$ 442,600 for allyl alcohol 424,000 acetone ,, 420,000 propaldehyde

These values, and many more might be given, show that the quantity of heat produced in the change from the combination of a certain number of atoms to certain other combinations of these atoms is dependent, in part, on the arrangement of the atoms in the initial combination. Isomeric molecules differ in the arrangements of their parts; and they also differ in the quantities of energy which are associated with the different atomic arrangements.

The attempts which have been made, chiefly by Thomsen (v. Th. vol. iv.), to connect in a definite way the thermal values of the combustion and formation of carbon compounds with the constitutions of these compounds have not led, as yet, to any very satisfactory results (v. also Armstrong's criticism of some of Thomsen's conclusions; P. M. Feb. 1887. 73).

Dissociation.—This very important branch of the subject is fully discussed in the article

Dissociation in vol. ii. pp. 385-410.

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XII. VAPOUR-PRESSURES OF SOLU-TIONS, METHODS BASED ON, v. MOLECULAR WEIGHTS, vol. iii. p. 420; ELECTRICAL METHODS, this vol. p. 185; and Solutions, this vol.
XIII. VISCOSITY OF LIQUIDS. A per

fect fluid, if such existed, would offer no resistance to a change of shape, and if its parts were set in motion relatively to each other by the action of any forces, the work done would be exactly equal to the kinetic energy produced. Moreover, this energy of motion would continue unchanged in amount if the fluid were then left to itself, and would not die away by being converted into heat as is the case with every known fluid. The property which causes this subsidence of relative motion is called viscosity. In consequence of the existence of viscosity a continual expenditure of energy is required to maintain the parts of a fluid in a state of steady motion relatively to each other, just as the existence of friction makes it necessary to apply a constant force to a body to keep it moving uniformly over another. From this resemblance in the two

cases the viscosity of a substance is often called its internal friction.

Imagine a layer of a liquid one cm. thick, contained between two flat plates, one of which is at rest and the other is moving parallel to the first. The liquid in contact with the plate which is at rest is at rest also, and the successive layers have velocities which increase uniformly as we go towards the moving plate, so that the liquid is being sheared, and this requires a continual supply of work; which is proportional to the area and velocity of the moving plate. If this -velocity is one cm. per second, the coefficient of viscosity is defined as the tangential force which must be applied per sq. cm. to the moving plate to maintain the motion, or if the relative velocity of the two surfaces is v, and their distance apart is l, the tangential force T per unit area required

to maintain the motion is given by $T = \eta \frac{v}{l}$ where η is the coefficient of viscosity.

From this definition it can be proved that if a circular plate or other solid of revolution be set vibrating about its axis of figure in a liquid, the amplitudes of successive vibrations will bear a constant ratio to each other, and the experimental verification of this is the best proof we have of the proportionality of the viscous forces to the relative velocity of the parts of the fluid.

The coefficient of viscosity is regarded by Maxwell (P. M. [4] 35, 133) as being the product of two factors, a coefficient of elasticity and a If between the parallel time of relaxation. plates spoken of above we have an elastic solid without viscosity, instead of a liquid, the result of moving one of the plates parallel to the other will be to distort the solid, and give rise to a force or stress tending to restore it to its original shape, and this force will be proportional to the amount of distortion and to the coefficient of rigidity of the body. Now, if the body is a viscous substance like pitch, the stress will soon die away, even though the surfaces be held in the new position, the molecules rearranging themselves so that this position becomes one of equilibrium. The time required for this to take place is what Maxwell called the time of relaxation. For solids it may be very large, extending to several hours or even days, but for ordinary liquids it is a small fraction of a second, while for a gas, such as air, Maxwell estimated it at the fifty thousand millionth of a second. The rate at which this internal stress is disappearing at any moment is proportional to the strain at that moment. If the upper surface is moved uniformly, relatively to the lower one, the strain will never disappear entirely, for, although the rearrangement of the molecules is continually going on andthe substance is tending towards its equilibrium state, the strain is being continually set up again by the relative motion of the two surfaces we are considering, and the body will soon get into a steady state in which the rate whereat the strain is being produced is equal to the rate whereat it is dying away. The stress produced by this constant strain is the tangential force required to maintain the motion, and Maxwell proved that this stress is proportional to the coefficient of rigidity of the substance and to the time of relaxation.

For the determination of coefficients of viscosity, Coulomb and others allowed some solid of revolution, such as a circular plate or a sphere immersed in the fluid, to vibrate about its axis of figure, and observed the ratios of the amplitudes of consecutive vibrations, from which the coefficient can be calculated. O. E. Meyer (W. 43, 1) modified the method by hanging a flat cylindrical box by a bifilar suspension, filling it with liquid, and observing the rate at which the vibrations died away. This method has been used by Mützel (W. 43, 15), and is capable of considerable accuracy. For absolute measurements it has some advantages in the fact that the lengths to be measured are not very small: but in most cases only relative values are required, and for obtaining these the transpiration method described below is preferable, as it is more readily applied, requires a smaller quantity of material, and lends itself better to accurate adjustment of temperature.

In the common form of the experiment the liquid is caused to flow through a capillary tube in consequence of a constant difference of pressure, P, between the ends, and the time, t, is observed which is required for a volume, v, to flow through the tube; then the coefficient of viscosity, η , is given by the equation $\eta = \frac{\pi}{8} \frac{Pr^{t}t}{lv}$,

where r is the radius and l is the length of the tube. Hence the coefficients for two liquids will be in the ratio of the times required for the outflow of the same volume with the same difference of pressure. If the liquid flows through the tube in consequence of its own weight, P will be proportional to its specific gravity, and the coefficients of viscosity will be proportional to the product of specific gravity and time of flow.

The formula given above assumes that the work done by the pressure is entirely converted into heat inside the tube, or, in other words, that the liquid flows out without any appreciable kinetic energy. If the tube is very long and narrow this may be taken to be the case, but generally a small correction has to be applied for the energy of the issuing fluid. The form of this correction is doubtful. Hagenbach (P. 109, 385) has given a formula which has been largely used, but Wilberforce (P. M. 1891. 407) has pointed out an error in his assumptions which makes the correction probably too small.

The coefficient of viscosity falls off very rapidly with rise of temperature, the average rate of fall for water between 0° and 10° being nearly 3 p.c. per degree, which shows the necessity for very accurate adjustment of the temperature during an experiment.

Using Maxwell's conception of the nature of viscosity, and making certain assumptions as to the way in which the rigidity and the time of relaxation vary with the temperature, Graetz (W. 24, 25) deduced the formula $\eta = A \frac{t_0 - t}{t - t_1}$, where

 t_1 is some low temperature at which η is infinite, and t_2 is the critical temperature. The formula-expresses the facts very well over a moderate range, but, as we might as reasonably assume

expresses the facts very well over a moderate range, but, as we might as reasonably assume the connection between η and t as between the time of relaxation and t, it can only be regarded as empirical.

Extensive observations of the viscosity-coefficients of organic liquids have been made by Graham (T. 1861), Rellstab (Inaug. Diss. Bonn, 1868), Guerout (C. R. 81, 1025; 83, 1291), Pribram a. Handl (Sitz. W. 1878 and 1879), and Gartenmeister (Z. P. C. 6, 524); but few laws have been observed beyond qualitative relations applicable to small classes of compounds. In general the viscosity increases with the molecular weight, but formic acid is an exception, as it is more viscous than acetic acid. Isomeric esters have nearly the same viscosities, that with the higher alcohol radicle having the greater. Normal compounds are generally more viscous than the corresponding iso- compounds, but the propyl halogen compounds, and a few others, form exceptions. At high temperatures these differences are less marked than at lower ones, rise of temperature tending to efface the distinction between normal and iso-compounds. Pribram a. Handl endeavoured to find the conditions of temperature under which the results are comparable, by determining the coefficients of viscosity of a number of liquids at various temperatures up to 50°. Taking the series PrCl, EtBr, PrBr, EtI, and PrI, they plotted the curve connecting viscosity with molecular weight, and found that while at the lower temperatures it was distinctly curved it got gradually flatter with rise of temperature, till at 50° it was almost a straight line, showing that at this temperature the increase of viscosity is proportional to the increase of molecular weight, whether this be due to the introduction of CH2 or of a halogen.

Though sulphuric acid is much more viscous than water, the first effect of adding water to it is, as Graham showed, to increase its viscosity, and this continues till 18 p.c. of water has been added, when a maximum is reached, and further addition of water causes a rapid diminution of viscosity. The mixture with maximum viscosity corresponds to the proportions given by the formula H₂SO₄.H₂O, and there is a maximum in the electrical resistance near the same point. Similar relations are shown by nitric and acetic acids. For mixtures of alcohol and water Graham showed that the particular proportion of the constituents which gives the greatest contraction has maximum viscosity, and Traube (B. 19, 871) showed further that aqueous solutions of most alcohols and acids of the fatty series have maxima which occur at different concentrations according to temperature.

Arrhenius (Z. P. C. 1, 285) has investigated the relation between the concentration of a solution and its viscosity; he finds that for indifferent substances the exponential formula $\eta = \mathbf{A}^{1-x}\mathbf{B}^x$ expresses the results, where η is the viscosity, x the proportion by volume of the dissolved substance, and 1-x that of the solvent, so that A is the viscosity of the pure solvent, and B is a constant for the dissolved substance which is independent of the concentration up to about 10 p.c. As the solvent is usually water, and its viscosity is taken as the unit in comparative measurements, the formula can be written in the form $\eta = B^x$. In every case investigated by Arrhenius B proved to be greater than unity, showing that the viscosity of water is increased by the addition of a small quantity of an indifferent substance. The same is generally true of salts, but not quite invariably so. A viscous fluid like glycerin has less effect than ether has; there seems, in fact, to be no relation between the value of B and the viscosity of the dissolved substance alone.

The latest and most comprehensive determinations of the viscosities of salt solutions are by Reyher (Z. P. C. 2, 744), Wagner (Z. P. C. 5, 31), and Lauenstein (Z. P. C. 9, 417). The formula of Arrhenius in the form $\eta = A^x$, where x is now the number of gram-molecules in a litre of solution, or A is the viscosity of a normal solution, is found to hold moderately well, but for many salts of acids of the aromatic series it shows divergences if the strengths of the solutions are greater than half normal.

Wagner worked with sulphates, nitrates, and chlorides of metals, and found that the viscosities of solutions of equivalent quantities of the various salts are additive quantities, the part which the base or the acid contributes being approximately constant. A further relation is shown by tabulating the viscosities of normal solutions of the chlorides arranged according to the periodic law, when the coefficients are found to diminish in any group as the molecular weight increases. From this arrangement it would seem that copper and manganese should be put in the eighth group, for their coefficients are almost exactly equal to those of nickel and cobalt.

Reyher determined the viscosities of a number of acids and their sodium salts, and found they could be divided into two classes. The strong mineral acids have viscosities about 3 p.c. less than those of the sodium salts, while the weaker mineral acids and organic acids have coefficients from 20 to 30 p.c. less.

Lauenstein has found the viscosity-coefficients of solutions of the sodium salts of many organic acids. In the fatty series he found that substitution of CH₂ for H increases the viscosity, while the replacement of H by COOH or OH diminishes it. The change from succinic to maleic acid diminishes the viscosity, but if we remove two more atoms of hydrogen, giving acetylene dicarboxylic acid, we get a large increase.

In the aromatic series the results are less regular. Introduction of a carboxyl group into the ring still diminishes the viscosity, and this diminution is greater in the para-position than in the meta-, and greater in the meta- than in theortho-; thus the viscosity diminishes regularly in the series shown by the formulæ at the top of p. 271.

According to the usual theory, the electricity which constitutes the current in an electrolyte is carried by charged ions, and it might be expected that the conductivity would be greater the more easily these ions were able to move along under the influence of the electromotive force—that is to say, the less the frictional resistance they had to overcome. On this account Wiedemann thought there would prove to be a connection between the viscosities and conductivities of solutions. Some experiments which he made seemed to point to the conductivity being inversely proportional to the viscosity, but though this was found at a later time to hold only in a few special cases, it has been shown that there is

$$C_eH_a.COONa, C_eH_a< \begin{matrix} COONa\\ COONa \end{matrix} 1:2, C_eH_a< \begin{matrix} COONa\\ COONa \end{matrix} 1:3, C_eH_a< \begin{matrix} COONa\\ COONa \end{matrix} 1:4.$$

With other replacing groups we have the following series, each being arranged in order of diminishing viscosity: -

$$\begin{array}{l} \textbf{1:2} \; \textbf{C}_{u}\textbf{H}_{4} < \begin{matrix} \textbf{COONa} \\ \textbf{NO}_{2} \end{matrix}, \; \textbf{1:3} \; \textbf{C}_{b}\textbf{H}_{4} < \begin{matrix} \textbf{COONa} \\ \textbf{NO}_{2} \end{matrix}, \; \textbf{C}_{u}\textbf{H}_{5}\textbf{COONa}, \; \textbf{1:4} \; \textbf{C}_{b}\textbf{H}_{4} < \begin{matrix} \textbf{COONa} \\ \textbf{NO}_{5} \end{matrix} \\ \textbf{1:4} \; \textbf{C}_{u}\textbf{H}_{4} < \begin{matrix} \textbf{COONa} \\ \textbf{OH} \end{matrix}, \; \textbf{C}_{u}\textbf{H}_{5}\textbf{COONa}, \; \textbf{1:3} \; \textbf{C}_{u}\textbf{H}_{4} < \begin{matrix} \textbf{COONa} \\ \textbf{OH} \end{matrix}, \; \textbf{1:2} \; \textbf{C}_{b}\textbf{H}_{4} < \begin{matrix} \textbf{COONa} \\ \textbf{OH} \end{matrix} \\ \textbf{1:2} \; \textbf{C}_{b}\textbf{H}_{4} < \begin{matrix} \textbf{COONa} \\ \textbf{OH} \end{matrix}, \; \textbf{1:4} \; \textbf{C}_{u}\textbf{H}_{4} < \begin{matrix} \textbf{COONa} \\ \textbf{OH} \end{matrix}, \; \textbf{C}_{c}\textbf{H}_{5}\textbf{COONa}, \; \textbf{1:3} \; \textbf{C}_{b}\textbf{H}_{4} < \begin{matrix} \textbf{COONa} \\ \textbf{OH} \end{matrix} \\ \textbf{OH} \end{matrix} \end{array}$$

undoubtedly a close connection between the two

properties.

This connection is shown very clearly by the work of Grotrian (P. 157, 130, 237; 160, 238), who determined the proportional rate of change with temperature of the viscosity and conductivity of salt solutions for various concentrationsthat is, he calculated from his observations the

values of $\frac{1}{f} \frac{df}{dt}$ and $\frac{1}{k} \frac{dk}{dt}$, where f is the coeffi-

cient of viscosity and k is the electrical conductivity, and found that though not equal they always vary in the same way with the concentration. If curves be drawn with the values of these coefficients at a fixed temperature for ordinates, and the concentrations for abscissæ, W. N. Shaw (Camb. Phil. Proc. 7, 21) has shown that the two curves run almost exactly parallel, any peculiarities in the shape of one being repeated in the other.

That the conductivity is not dependent on the viscosity alone is shown by the fact that if we add to an electrolyte such quantities of different non-conductors, as alcohol, sugar, gly-cerin, &c., as increase the viscosity by the same amount, the conductivity will generally be affected differently by the different substances. Arrhenius (Z. P. C. 9, 495) has investigated this effect at some length. He finds that the conductivity can

be expressed by the equation: $l = l_0 \left(1 - \frac{\alpha}{2}x\right)^2$,

where l_0 is the conductivity of the electrolyte alone, and I that which it has when x p.c. of the water is replaced by a non-conductor. Thus a may be taken as defining the change in the conductivity. The change in the viscosity of the electrolyte on the addition of 1 p.c. of non-conductor is the same as the change would be with water alone, and is hence equal to A-1, where A is the constant of the formula $\mu = A^x$.

On tabulating the values of 1000a and 1000 (A-1) for a number of different salt solutions and non-conductors, it appears that the equation, $1000 \alpha = c + 1000 c'(A-1)$ holds as a first approximation, and the electrolytes fall into four classes, for the members of any one of which the constants c and c' are the same. These classes are:

Solutions of strong acids and bases;

(2) Solutions of salts of the type KCl;

K2804; (3)BaCl2.

Arrhenius considers the relation not to be exact, but thinks it should include a term depending on the degree of dissociation, which can be neglected only in the cases of the more completely dissociated salts. J. W. C.

XIV. VOLUME-CHANGES, BASED ON, v. VOLUMES, SPECIFIC; this vol.

PHYSODIN C₁₀H₁₀O₇. [125°]. Extracted by ether from the dried lichen Parmelia physodes (or Ceratophylla] and crystallised from alcohol (Gerding, N. Br. Arch. 87, 1). Mass of rainute prisms, insol. water. Forms a yellow solution in KOHAq.

PHYSOSTIGMINE v. ESERINE. PHYTO-ALBUMOSES v. PROTEÏDS.

PHYTOLACCIC ACID. Occurs as K salt in the Poke Berry, the fruit of Phytolacca decandra (Terreil, C. R. 91, 856; cf. Claesson, Ph. [3] 10, 566). Extracted by dilute alcohol. Yellowishbrown resin.

PHYTOSTERIN v. vol. ii. p. 149. PHYTO-VITELLIN v. PROTEIDS.

PIAZTHIOLE C₆H₄ $<_{\dot{N}}^{\dot{N}}>$ S. [44°]. (206°).

Formed by heating o-phenylene-diamine with aqueous SO₂ at 190° (Hinsberg, B. 22, 2899). Colourless crystals, sl. sol. hot water. Feeble base. Reduced by tin and HCl to o-phenylenediamine.

Reference.—METHYL-PIAZTHIOLE.

PICENE C_{2.}H₁₄. [345° cor.]. (520°). V.D. 9.8 (obs.). Occurs in the highest boiling portions of brown-coal tar (Burg, B. 13, 1834), and in the residues of Californian petroleum (Graebe a. Walter, B. 14, 175). Appears also to be formed by the action of ethylene bromide and AlCl, on naphthalene (Lespeau, Bl. [3] 6, 238). White plates with blue fluorescence, insol. alcohol and ether, sol. boiling solvent naphtha (150°-170°). On oxidation with CrO_s it yields picoquinone C22H12O2, which crystallises from HOAc as a dark orange-red powder, sublimes in red needles, and forms a green solution in pure

H₂SO₄.

Di-bromo-picene C₁₂H₁₂Br₂. [206°]. Needles.

(from xylene), insol. alcohol.

Picene hydrides C.H., (over 360°) and $C_{22}H_{36}$ [175°] (over 360°) are got by heating pieene with HIAq and red P at 250° (Liebermann a. Spiegel, B. 22, 781). $C_{22}H_{34}$ is liquid.

PICHURIN OIL contains the glyceryl ether of lauric acid (Sthamer, A. 53, 390).

PICOLINE v. METHYL-PYRIDINE.

PICOLINIC ACID v. PYRIDINE CARBOXYLIC

Dipicolinic acid v. Pyridine DICARBOXYLIC ACID.

DIPICOLYL v. DI-METHYL-DIPYRIDYL.

PICRACONITINE v. ACONITE ALKALOIDS.

PICRAMIC ACID v. DI-NITRO-AMIDO-PHENOL.

PICRAMIDE v. TRI-NITRO-ANILINE.
PICRASMIN C_{3a}H_{4a}O₁₀. [204°]. A bitter substance which, together with C_{3a}H₄₀O₁₀ [209°-

212°], may be extracted by dilute alcohol from Picræna excelsa (Massute, Ar. Ph. [3] 28, 147; C. J. 58, 791). HClAq converts it into picrasmic acid C₂₁H₄₀O₄(CO₂H)₂, while fuming HIAq displaces three methyls.

PICRIC ACID v. TRI-NITRO-PHENOL.

PICRO-ACONITINE v. Aconite alkaloids.

PICROCROCIN C₃₉H₆₆O₁₇. [75°]. Obtained by extracting saffron with ether (Kayser, B. 17, 2233). Colourless prisms. V. sol. water and alcohol, sl. sol. ether. Bitter taste. By heating with baryta-water or with dilute acids it is split up into crocose and the ethereal-oil of saffron C10H16

PICROERYTHRIN v. ERYTHRITE.

PICROLICHENIN $C_{12}H_{20}O_6$. [100°] (A.). S.G. 1·176. Extracted by alcohol from the lichen Variolaria amara (Alms, A. 1, 61; Vogel, J. 1857, 515). Trimetric octahedra, sl. sol. hot water, v. sol. alcohol and ether. Tastes very bitter. Its alkaline solutions turn red in air.

PICROROCCELLIN $C_{27}H_{39}N_3O_5$. Occurs in a variety of the lichen Roccella fuciformis (Stenhouse a. Groves, A. 185, 14). Prisms, insol. water, m. sol. boiling alcohol, sl. sol. ether. Yields benzoic acid when oxidised by chromic acid mixture. Converted by boiling with HOAc and HCl into xanthoroccellin C21H18N2O2, which crystallises from alcohol in yellow needles [183°]. Boiling dilute ($1\frac{1}{2}$ p.c.) NaOHAq converts picroroccellin into $C_{21}H_{12}N_2O_3$ crystallising from alcohol in prisms [154°].

PICROSCLEROTIN. Occurs in ergot of rye (Dragendorff, C. C. 1878, 125, 141; Blumberg, Ph. [3] 9, 23, 66, 147). Dissolved in acids and is reppd. by ammonia. It is an active poison

V. ERGOTININE. with bitter taste.

the seeds of Menispermum cocculus (Boullay, A. Ch. [1] 30, 209; Casaseca, A. Ch. [2] 30, 307; Oppermann, Mag. Pharm. 35, 233; Pelletier a. Couerbe, A. Ch. [2] 54, 181; Liebig, A. 10, 203; Regnault, A. Ch. [2] 68, 160; Barth, J. pr. [1] 91, 155; Paterno a. Oglialoro, G. 6, 521; 7, 193). It is accompanied by anamirtin and picrotin (Barth a. Kretschy, M. 1, 99, 2,796).

Preparation.-1. The grains are exhausted with boiling alcohol, the extract evaporated, and the residue boiled with water. The aqueous solution is ppd. by lead acetate and the filtrate, freed from lead by H₂S, evaporated. The residue is crystallised from benzene and water successively (B. a. K.).-2. The powdered seeds are boiled with water, the filtrate treated with lead acetate and H2S successively, and evaporated to crystallisation. The product is re-crystallised from water and alcohol successively (Schmidt, A. 222, 313).

Properties.—Colourless needles, v. sl. sol. cold water, sl. sol. ether and chloroform, v. sol. alcohol. Very poisonous and very bitter. Sol. alkalis. Reduces Fehling's solution and am-

moniacal AgNO₅.

Reactions.—1. When boiled with benzene for some time it is split up into picrotin and picrotoxinin (Schmidt). Probably the picrotoxin employed was a mixture of these two bodies.—2. HCl passed into an ethereal solution forms picrotoxide.—3. Cold AcCl also forms picrotoxide, but on boiling it yields a crystalline acetyl derivative [185°].-4. NaOAc and Ac2O form C19H22O, [227°]

Picrotoxinin C₁₅H₁₆O₆ aq. [201°]. S. 14 at 15°; S. (benzene) 34 at 21°. Obtained from picrotoxin by treatment with benzene, CHCl, or AcCl. It constitutes about 30 p.c. of crude picrotoxin (Barth a. Kretschy). Colourless picrotoxin (Barth a. Kretschy). Colourless needles or plates. Very bitter and very poisonous. V. sol. hot water. Octoors 11,2004 red. Mixed with dry KNO₃ (3 pts.), moistened it gives a red colour (Langley, Am. S. [2] 34, 109). After heating with milk of magnesia, the cold filtrate is coloured red by FeCl₂, the colour being destroyed by HCl. Bromine forms $C_{1a}H_{1s}BrO_{s}$ [250°-255°]. BzCl gives a crystal-Bromine forms line compound (not the benzoyl derivative) [238°].

Picrotoxide $(C_1, H_{16}O_6)x$. [above 310°]. Formed by the action of AcCl on picrotoxin, and by passing HCl into its ethereal solution (P. a. O.). Crystalline, insol. ordinary solvents. According to Schmidt, picrotoxide obtained by means of AcCl crystallises in needles [225°], sl.

sol. cold water and alcohol.

Picrotin C₂₅H₃₀O₁₂. [c. 247°]. S. 16 at 17°; S. (benzene) 023 at 22° (Schmidt). Constitutes 60 p.c. of crude picrotoxin (B. a. K.). Crystallises with $2\frac{1}{2}$ aq, $3\frac{1}{3}$ aq, and $4\frac{1}{2}$ aq. It is very bitter, but not poisonous. Reduces hot Fehling's solution and ammoniacal AgNO3. H2SO forms a yellow solution. BzCl forms a benzoyl

derivative [245°] (S.).

Anamirtin C₁₀H₂₁O₁₀. Constitutes 2 p.c. of

Wrotechy. M. J. 131). crude picrotoxin (Barth a. Kretschy, M. J, 131). Short needles (from water), v. sl. sol. benzene. Neither bitter nor poisonous. Turns brown at 260° and black at 280° without melting.

Cocculin C₁₀H₂₈O₁₀. Occurs in small quantity in Cocculus Indicus (Löwenhardt, A. 222, 353). Slender needles, sl. sol. hot water, nearly insol. alcohol and ether. Does not give Langley's reaction. Is perhaps identical with anamirtin.

PICRYL. The radicle tri-nitro-phenyl. PICRYL CHLORIDE v. CHLORO-TRI-NITRO-BENZENE.

PIGMENTS, ANIMAL.

BILE PIGMENTS. Bile contains bilirubin, bilifuscin, biliprasin, and probably also biliverdin, which is a product of oxidation of bilirubin. Nitric acid changes the colour of the bile pigments through green, blue, and red to yellow (Gmelin). These colours may be observed by adding nitric acid to a dilute solution of the bile pigments in aqueous alkali, or by gently pouring H₂SO₄ into a solution of the pigments mixed with NaNO₃ (Fleischl, *Fr.* 15, 502). The changes in the absorption spectra produced by nitric acid have been studied by Jaffé (Z. [2] 5, 666). An alcoholic solution of bromine also produces a play of colours with bile pigments (Capranica, G. 11, 430). The absorption spectra of the bile pigments have been studied by Heynsius a. Campbell (Pf. 4, 497) and McMunn (Pr. 35, 388). Bilirubin, hæmoglobin, and chlorophyll all absorb the violet end of the spectrum, giving an abrupt edge: biliverdin transmits more green; biliprasin, bilifuscin, and bilihumin abscrbup to between D and E. The colouring matters of bile are probably got by reduction of homatine.

ttself formed by the action of bile acids on hæmoglobin (McMunn, Pr. 31, 206). According to Latschenberger (M. 9, 52), hæmoglobin yields simultaneously melanin and bile pigments. All the colouring matters of bile, including hæmatine, urobilin (in bile), and bilirubin, are oxidised to choletelin, which body apparently passes into blood serum, and is then excreted by the kidneys. The absorption bands of bile are due to choletelin and urobilin.

Bilirubin C₂₂H₂₆N₄O₆ (Städeler, A. 182, 323; Maly, A. 181, 106), or C_{1e}H_{1e}N₂O₃ by Raoult's method (Nencki a. Rotschy, M. 10, 568). S. (chloroform) ·17 (Thudichum, Z. [2] 4, 554). Ox gall stones are often largely composed of the lime compound of bilirubin (Maly, A. 175, 76).

Preparation.—Brown human gall stones are powdered and extracted with ether; the residue is boiled with water, and treated with dilute HCl. The mass is washed, dried, and extracted with chloroform; the chloroform is distilled off, and the residue treated with absolute alcohol. It is then treated with ether and alcohol repeatedly, again dissolved in chloroform, and precipitated by absolute alcohol (Burdon-Sanderson).

Properties.—Orange powder, insol. water, nearly insol. ether, v. sl. sol. alcohol, sol. benzene The colour of the skin in and chloroform. jaundice is probably due to bilirubin. Bilirubin dissolves in alkalis, forming an orange solution, which gradually absorbs oxygen from the air, and then gives a green pp. of biliverdin on adding an acid. An alkaline solution of bilirubin mixed with an equal bulk of alcohol gives, on adding HNO, containing nitrous acid, a green colour changing through blue to red. Sodiumamalgam forms hydrobilirubin. Bromine-vapour yields various brominated products (Thudichum, C. J. 28, 389; 30, 27). Br in chloroform gives C₅₂H₃₅Br₅N₄O₆, a dark bluish-green powder, forming a dark-blue solution in alcohol or ether (Maly, 4. 181, 106). Chlorine passed into a solution of bilirubin in CHCl, forms several chlorinated bodies (Thudichum). A solution of p-diazobenzene sulphonic acid added to a solution of bilirubin in chloroform mixed with alcohol gives a red colour changing to blue on adding conc. HClAq, and turned red again on adding an alkali (difference from other bile-pigments) (Ehrlich, Fr. 23, 275).—CaC₂₂H₂₄N₄O₆. Occurs in gall stones. Obtained also by ppg. an ammoniacal solution by CaCl₂, as brown flakes drying to a lustrous dark-green mass, which yields a dark-blue powder.

Hydrobilirubin C₁₂H₁₀N₁O₇. Obtained by reducing bilirubin in alkaline solution with sodium-amalgam (Maly, A. 163, 77). According to McMunn it is not, as had been stated, identical with the urobilin which is found in normal urine, in the urine of febrile patients (Jaffé, Virchow's Archiv, 47; Disqué, H. 2, 271), and in excrement (Vaulair a. Masius, Centralbl. f. d. Med. Wissensch. 1871, No. 24). A similar substance is formed by reducing hæmoglobin, hæmatine, or hæmatoporphyrin in alcoholic solution with tin and HCl (Hoppe-Seyler, B. 7, 1065; Le Nobel, C. C. 1887, 538; McMunn, Proc. Physiol. Soc. 1888, 1), and by the action of H₂SO₂ on a solution of albumen in HOAc (Michailoff, J. R. 16, 269). Reddish-brown powder with green lustre, sl. sol. water, v. sol. alcohol, m. sol. ether. Its alkaline

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solutions are brown, its solution in chloroform is yellowish-red. The alcoholic solution shows an absorption-band near F. Does not give Gmelin's reaction with HNO₃. May be reduced by sodium-amalgam or by tin and HCl to a colourless product, which in acid solutions is reoxidised by air to hydrobilirubin.

Choletelin $C_{18}H_{18}N_2O_8$. Occurs in normal urine, and is obtained by passing nitrous vapours into a solution of bilirubin in alcohol (Heynsius a. Campbell; McMunn, J. Th. 1881, 213). Brown amorphous powder, sol. alkalis and alkaline carbonates, alcohol, ether, and chloroform. Ammoniacal AgNO₃ ppts. reddish-brown $C_{18}H_{18}\Delta g_2N_2O_8$. Does not give Gmelin's reaction. Its absorption spectrum contains a broad band from b to F.

Biliverdin $C_{22}H_{36}N_1O_8$. Formed by oxidation of a solution of bilirubin by air or by PbO₂. Dark-green powder, insol. water and chloroform, sl. sol. ether, v. sol. alcohol, CS₂, and benzene. Its solutions are green. Its alcoholic solution gives dark-green pps. with baryta-water, ammoniacal CaCl₂, and AgNO₄. Ag₂O added to its alcoholic solution ppts. bilipurpin, sol. NH₂Aq and reppd. by HCl as a purple powder. Prolonged treatment with Ag₂O converts bilipurpin into yellow biliflavin. HNO₃ added to an alcoholic solution of biliverdin gives a bluish-violet, and finally yellow colour (Thudichum)

red, and, finally, yellow colour (Thudichum). Bilifusoin $O_{18}H_{10}N_2O_4$. Prepared by washing gall stones with ether, hot water, and chloroform, adding dilute HClAq, and extracting with hot chloroform. The extract is evaporated and the bilifusoin dissolved in alcohol, which leaves bilirubin undissolved (Brücke, J. pr. 77, 72; Städeler, A. 132, 325). Almost black mass, which yields a dark-brown powder. Forms a brown solution in alcohol and alkalis, nearly insol. water, ether, and chloroform. Gives Gmelin's colour-reaction. Ppd. by ammoniacal CaCl₂. Simony (Sitz. W. [3] 73, 181) obtained from the bile of a corpse a bilifuscin which did not give Gmelin's reaction, and which formed an olive-brown solution in alcohol, HOAc, and alkalis.

Biliprasin C₁₆H₂₂N₂O₆. Extracted by alcohol from the residue of gall stones from which bilirubin and bilifuscin have been removed by chloroform (Städeler). Nearly black mass, yielding a greenish-black powder, insol. water, ether, and chloroform, v. e. sol. alcohol, forming a green solution which, unlike one of biliverdin, is turned brown by ammonia. Gives Gmelin's reactica.

Bilihumin. Left after extracting biliprasin with alcohol (S.). Black powder, m. sol. warm NaOHAq. Exhibits Gmelin's reaction.

Bilicyanin. A product of partial oxidation of bilirubin and other bile pigments (Maly, Sitz. W. [2] 59, 597). Prepared by adding an alcoholic solution of Br to a solution of bilirubin in chloroform. Occurs in gall stones. Its alcoholic solution is blue, but turned dingy green by alkalis, the blue colour being restored by acids. Bile also contains a blue substance strongly resembling indigo, forming a yellowish solution in alkalis (Ritter, Bl. [2] 18, 212; Andouard, Bl. [2] 31, 139).

URINARY PIGMENTS. Urine usually contains urobilin (v. supra). After urine, containing urobilin, has been mixed with its own bulk of

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HClAq and heated to boiling, and then allowed to cool, the urobilin may be extracted by ether and recognised by its absorption band (Grimbert, J. Ph. [5] 18, 481). According to McMunn (Proc. Physiol. Soc. 1888, 5), there are two kinds of urobilin, one in normal urine and the other in pathological urine, and neither is identical with hydrobilirubin. Normal urobilin is identical with a pigment got from acid hæmatin by successive treatment with H2O2 and sodium-amalgam. By the action of oxidising agents indiga-blue and indirubin can be obtained from urine (v. Indigo). Pigments can also be obtained by boiling urine with HClAq.

Uromelanin. Prepared by evaporating urine to one-sixth of its bulk at 60°, adding 10 p.c. HCl, and, after two days, removing uric acid by filtration. The filtrate is then boiled for 18 hours, and the ppd. pigment washed with water, alcohol, and ether, dissolved in NaOHAq, and reppd. by H₂SO₄ (Udranski, H. 11, 537; 12, 33; cf. Plosz, H. 8, 89). Brownish-black plates, insol. cold water, ether, and chloroform, sl. sol. alcohol and HClAq, v. sol. isoamyl alcohol and alkalis. Not decomposed at 115°. Potash-fusion yields NH₃, formic, acetic, butyric, and protocatechuic acids and pyrocatechin. Urino contains 03 p.c. of this substance, which appears to be a humous body formed by decomposition of the reducing substance of normal urine.

Urofuscohæmatin C₈₄H₅₄N₄O₁₈ (') and urorubrohæmatin C_{si}H_{ss}N_sFeO_{1s} (?) were obtained by Baumstark (B.7, 1170) from the urine sta patient suffering from Lepra. Urofuscohematin is a black pitchy substance, insol. water, alcohol, ether, chloroform, acids, and NaClAq, sol. aikalis, alkaline carbonates, and alkaline phosphates, forming brown solutions. Urorubrohæmatin is a light blue-black mass, which differs from urofuscohæmatin in not being ppd. when HCl is

added to its solution in NaOHAq.

Black pigment in Melanuria, v. vol. iii.

Urorosein. An unstable substance which can sometimes be extracted from acidified pathological urine by isoamyl alcohol, to which it imparts a rose colour. The solution shows an absorption band in the green (Nencki a. Sieber, J. pr. [2] 26, 333). A similar substance (urorubin) was extracted by ether from urine that had been boiled 15 minutes with 7 p.c. HCl, exposed to air (Plosz, H. 8, 85). It was left on evaporation as a dark cherry-red mass. Apparently the same pigment, or rather its leuco-compound, occurs in urine of dogs after administration of skatole (Mester, H. 12, 130). The pigment itself is then got by adding HCl to an alcoholic, ethereal, or aqueous extract of the evaporated urine.

OTHER PIGMENTS. Blood pigments v. HEMOGLOBIN. Muscle pigments v. Muscle.

Eye pigment v. Melanin. A purple pigment occurs in the retina of animals. It is insol. ordinary solvents, but dissolves in bile, and in an aqueous solution of the bile acids. It loses its colour in daylight, but recovers it in the dark (Boll, J. Th. 1877, 818; Kühne, J. Th. 1877, 818; 1878, 279; Ayres, J. Th. 1879, 259).

Yellow pigment of animals v. LUTEIN.

Tetronerythrin. A red pigment extracted by chloroform from the red spot in the eye of the blackcock and red grouse (Wurm, J. 1872, 842; 1875, 885; Merejkowski, J. Th. 1881, 371). H2SO4 gives a blue colour changing to black.

Pyocyanin. Occurs in blue pus (Fordos, J. 1860, 596; Lücke, J. 1863, 658; Gessard, J. Th. 1882, 55). It is formed, together with other pigments, by Bacillus pyocyanicus in peptonised gelatin (Babès, C. R. Soc. Biol. [9] 1, 438). Blue trimetric prisms or needles (from CHCl3) which become green on keeping, v. sol. chloroform, alcohol, and water. Turned red by acids, but becomes blue again on adding alkalis. It shows two absorption bands, one being in the ultraviolet.

, Pigments of Purpura lapillus. The pigments are furnished by a yellowish-white fascia which extends along the rectum (Letellier, C. R. 109, 82). This contains two green substances, applegreen monoclinic crystals turned dark blue by light, and dull-green trimetric crystals turned violet or crimson by light. The green substances are sol. ether and chloroform, but become insoluble on exposure to light. The change to purple (punicin) is accompanied by absorption of oxygen (Schunck, B. 12, 1359).

Pigments of sea-anemones. Actinia mesembryanthemum contains a pigment allied to hemo-chromogen and hemato-porphyrin, and to hæmatine. It also contains biliverdin. Actinia cereus, Bunodes ballii, and Sagartia bellis contain a green pigment resembling chloro-fucin, but not identical with any animal or plant chloro-

phyll (McMunn, Pr. 38, 85).
PIGMENTS, VEGETABLE, v. Chlorophyll, Alkanet, Beth-a-barra colour, Bixin, Brazileïn, CHICA, COLEÏN, CURCUMIN, DRAGON'S BLOOD, HÆ-MATOXYLIN, LITHOSPERMUM ERYTHRORHIZON, LIT-

MUS, LUTRIC ACID, PALMELLIN, and SANTALIN.

Anthocyanin. The blue pigment of flowers is sol. water and alcohol, insol. ether. It is free from N, and is turned red by acids and green by alkalis (Fremy a. Cloez, J. Ph. [8] 25, 249; Filhol, C. R. 39, 194; 50, 345, 1132; Schönn, Fr. 9,

Anthoxanthin. Yellow flowers contain anthoxanthin, which is insol. water, and anthoxanthein, which is sol. water. Both are sol. alcohol and ether. The petals of Rosa gallica contain a pigment sol. alcohol, insol. ether, which forms amorphous $Pb_2C_{21}H_{20}O_{20}$ (?) (H. Senier, Ph. [3] 7, The colouring matter of the berries of 650). Phytolacca decandra has been examined by Hilger and Bischoff, L. V. 23, 456; B. C. 1879, 875).

PILIGANINE. A very poisonous alkaloid occurring in piligan, a Brazilian lycopod (Adrian, C. R. 102, 1322). Soft mass, with alkaline reaction, fuming with HCl. Sol. water, alcohol, and chloroform, sl. sol. ether. Emeto-cathartic in action. Its hydrochloride forms minute deliquescent crystals.

PILOCARPENE $C_{10}H_{16}$. (178°). S.G. 12.852. V.D. 4.0. [α]_D = 1.21. Obtained by steam distillation from jaborandi leaves (Hardy, Bl. [2] 24, 498). Fragrant oil. Dextrorotatory. Yields C, H, 2HO [49.5°

PHOCARPINE $C_{11}H_{16}N_{1}O_{2}$ i.e. $CH \stackrel{N=CH}{<} C.CMe \stackrel{CO}{<} Me_{2} > 0$. [159°] (Blyth). $[a]_{D} = 101.6$ in a 7.24 p.c. solution.

Occurs in the leaves and bark of jaborandi (Pilocarpus) (Gerrard, Ph. [3] 5, 865, 965; Hardy, Bl. [2] 24, 497; Kingzett, C. J. 30, 367; Harnack a. Meyer, A. 204, 67). Prepared synthetically from a-oxy-a-pyridyl-propionic acid CH₃.C(OH)(C₃H₄N).CO₂H by treatment with phosphorus tribromide and heating the resulting CH₃.CBr(C₅H₄N)CO₂H with NMe₂ at 150°. In this way pilocarpidine C₁₀H₁₄N₂O₂ is formed, and this is converted into pilocarpine by heating with MeI and MeOH, and oxidising the product with aqueous KMnO, (Hardy a. Calmels, C. R. 105, 68; Bl. [2] 48, 233).

Preparation.—Jaborandi leaves are digested with 1 p.c. HClAq, the extract treated with Pb(OAc)2, filtered, and the filtrate ppd. by phosphomolybdic acid. The pp. is decomposed by baryta-water at 100° (Pöhl, Bl. [2] 34, 340).

Properties.—Crystalline; begins to sublime at 153°; at 160°-170° the sublimate consists of yellow drops (Blyth). Dextrorotatory. Pilocarpine may be estimated by means of the aurochloride (Christensen, Ph. [3] 12, 400). Poisonous, being diaphoretic. Forms resinous compounds with potash, NaOH, and baryta; these compounds are v. sol. water, v. sl. sol. alcohol, and are decomposed by acids, even by CO2. They may be considered to be salts of

pilocarpic acid C₁₁H₁₈N₂O₃.

Reactions. -1. Fuming HNO_s (300 pts.) converts it into pilocarpidine nitrate (Chastaing, C. R. 94, 968).—2. Boiling with water for twelve hours splits it up into trimethylamine and oxypyridyl-propionic acid (Hardy a. Calmels, C. R. 102, 1562).—3. Boiling HClAq forms MeOH and pilocarpidine.—4. KMnO₄ forms NMe₃, oxypyridyl-malonic acid, and finally pyridine (β)carboxylic acid. -5. Bromine added to a chloroform solution forms C11H15N2O2Br5, crystallising in minute prisms, converted by moist Ag₂O into di-bromo-pilocarpine $C_{11}H_{14}Br_2N_2O_2$ (Chastaing, C. R. 97, 1435). Chlorine forms $C_{11}H_{14}N_2O_2Cl_3$, which is amorphous, and slowly forms crystalline C₁₁H₁₅N₂O₂Cl₅.-6. Yields NMe₃ when distilled with potash (Harnack a. Meyer; cf. Chastaing, C. R. 94, 223).-7. The barium compound on distillation yields jabonine C, H, N, an oil with fœtid odour, yielding the amorphous salts B'AuCl_s, B'HAuCl₄, B'₂PtCl₄, and B'₂H₂PtCl₆,

Salts.—B'HCl: needles, v. sol. alcohol.— B'HNO₂: trimetric lamelle.—B'₂H₂PtCl₆: golden tablets (from hot water).—B'PtCl₄: crystals.—B'HAuCl₄: minute needles.—B'AuCl₃ [88°]: slender needles.—B'2AuCl: small needles. B'HAu2Cl,: minute needles.—B'AgNO: minute radiating needles. - B'2AgNO,: needles. B'CrN₂H₇(SCN)₄: red silky needles, sol. alcohol (Christensen, J. pr. [2] 45, 868).—B'₂CuO₂H₂: green powder, ppd. by adding CuCl₂ to a solution of pilocarpine in baryta-water.—B'AgOH: curdy pp.—B'Me₂PtCl_s: crystals.—B'EtI. [c. 30°].-B'EtBr. [c. 60°]. Very hygroscopic (Chastaing, C. R. 101, 507).

Pilocarpidine C10H14N2O2 i.s. NMe2.CMe(C4H4N).CO2H. Occurs in jaborandi leaves (Harnack, A. 238, 230). Formed by the action of HClAq or fuming HNO, on pilocarpine, by heating dry pilocarpine for twenty-four hours at 120°, by boiling pilocarpine or its baryta compound with water for forty eight hours, and by heating the baryta compound for thirty minutes

at 150°. Prepared synthetically by heating CH₃.CBr(C₅H₄N).CO₂H with trimethylamine at 150° (H. a. C.). Very deliquescent, m. sol. water, v. sol. alcohol. Acts physiologically like pilocarpine, but not so strongly. Its alkaline salts are gummy, sol. water, insol. alcohol, and decomposed by CO2.—B'HCl: radiating needles, v. soluble in water.—B'HAuCl, aq. Rectangular prisms. — B'AuCl₃. [145°]. Yellow plates. — B'H.PtCl₆ aq: small red prisms.—B'MeI.— B' $_{2}$ H. $_{1}$ Pro $_{1}$ Cross aq: sman red parameter $_{2}$ B' $_{2}$ MeAuCl, [153°]. Prismatic needles. Changes when fused into MeCl and B'AuCl, PIMARIC ACID $_{20}$ H $_{30}$ O $_{2}$. [211°]. [α] $_{p}$ = 72·5

in a 3.8 p.c. (saturated) alcoholic solution at 15°. Occurs in galipot, the hardened resin of Pinus maritima (Laurent, A. Ch. [2] 72, 384; [3] 22, 459; Sievert, Z. f. d. g. Naturwiss. 14, 311; Maly, A. 129, 94; 132, 253; Strecker a. Duvernoy, A. 148, 143; 150, 131; Cailliot, Bl. [2] 21, 387; Bruylants, B. 11, 447; Haller, B. 18, 2165; Vesterberg, B. 18, 3331; 19, 2167; 20,

Preparation .- Finely-divided galipot is stirred with half its weight of dilute (70 p.c.) alcohol, left for some days, and squeezed in a cloth. The press-cake is treated several times in this way, finally with 80 p.c. alcohol. The residue is dissolved in somewhat more than the calculated quantity of hot dilute (3 p.c.) NaOHAq. The mixture of Na salts that separates after some days is recrystallised from water, decomposed by HCl, and the free acids crystallised from alcohol or HOAc. (\$)-Pimaric acid remains in the mother-liquor.

Properties.—Rectangular plates, insol. water, v. sl. sol. alcohol, ether, and HOAc, m. sol. hot ligroin, m. sol. hot NaOHAq, sl. sol. NH,Aq. On shaking the ethereal solution with a drop of ammonia the NH, salt separates in slender needles. Not reduced by sodium-amalgam. May be distilled in vacuo. Dextrorotatory. HIAq (S.G. 1.96) forms C₂₀H₂₄ (320°-330° uncor.). By distillation of the (crude) Ca salt Bruylants obtained ethylene, propylene, amylene, acetone, methyl ethyl ketone, di-ethyl ketone, toluene, xylene, ethyl-toluene, terebene, and diterebene.

Salts.—All the salts are insol. ether. KA' (dried at 100°). Soapy mass of pliant needles. NaA' 5aq. Slender needles (from 80 p.c. alcohol), sl. sol. cold water. — CaA', aq: needles. BaA', 9aq: pliant needles. - CuA', - PbA',

-AgA': amorphous pp., becoming crystalline, Ethers.—MeA'. [69°].—EtA. [52°]. Chloride C₂₀H₂₀OCl. [66°]. Got by adding PCl, to a solution of the acid in CS. Small

prisms, v. e. sol. ether and CS2.

(β)-Pimaric acid $C_{\rm m}H_{\rm a}O_{\rm m}$ [140°-150°]. S. 9·26 in 98 p.c. alcohol at 15°. [a]_p = -272° in a 8·17 p.c. alcoholic solution. Obtained as above (Vesterberg, B. 20, 3248). Trimetric prisms; a:b:c = .810:1: .614, insol. water, v. sol. NH₂Aq. Lævorotatory.—NaA': m. sol. ether.—PbA'2: needles, insol. alcohol and ether. Haller (B. 18, 2165) obtained an inactive pimaric acid [c. 149°], which was perhaps a mixture of the dextro- and lævo- varieties.

PIMELIC ACID C,H, O, i.e.
CO.H.CHPr.CH., CO,H. Isopropyl-succinic acid.
Mol. w. 160. [114]. Electrical conductivity:
Walden, B. 24, 2037. Formed by fusing camphoric acid with potash (Hlasiwitz a. Grabowski,

A. 145, 205; Kachler, A. 169, 168). Formed also by heating either of the following isopentane tricarboxylic acids: (CO2H)2CPr.CH2.CO2Hor CO.H.OHPr.OH(CO.H)₂ (Waltz, B. 15, 609; A. 214, 60; Hjelt, B. 16, 2622; Schleicher, A. 267, 123). Obtained also by the action of KOHAq on isopropyl-acetyl-succinic ether (Roser, A. 220, 276). According to Arppe (J. 1864, 377) it is not formed, as stated by Laurent (A. Ch. [2] 66, 163), by oxidation of oleïc acid. Nodules (from water) or triclinic crystals; a:b:c = 497:1: 599; $a=81^{\circ} 50'$; $\beta=100^{\circ} 2'$; $\gamma=85^{\circ} 6'$ (Wreden, A. 163, 323; Von Zepharovitch, Sitz. W. [1] 73, 7). V. sol. water, alcohol, and ether. Yields butyric

acid when fused with potash.
Salts.—(NH₄)₂A". Hygroscopic leaflets.—
Na₂A" (dried at 130°).—CaA"aq: sandy crystalline powder. S. 47 at 9°; 25 at 100° (Bauer a. Schuler, J. 1878, 734; B. 10, 2031; M. 4, 345); ·30 at 13° (Roser).—SrA": nearly insol. hot water, v. e. sol. cold water.—Ag₂A": pp.

Ethyl ether Et.A". (236°-240°).

Chloride $C_1H_{10}O_2Cl_2$. (210°). Anhydride $C_1H_{10}O_3$. (245°-250°). Got by distilling the acid. Reconverted into the acid by alkalis.

Imide C₇H₁₁NO₂. [60°]. Got by heating the ammonium salt. Tables (from water) or small needles (from alcohol-ligroïn). Insol.

ligroin, v. sol. alcohol and ether. n-Pimelic acid $CH_2(CH_2.CH_2.CO_2H)_2$. [103°]. (272° at 100 mm.) (Krafft a. Noerdlinger, B. 22 S. 4.2 at 20°. H.C.p. 828,900. H.F. 818). 243,100 (Stohmann, J. pr. [2] 45, 480).

Formation.—1. By heating suberone with HNO₃ (Dale a. Schorlemmer, C. J. 35, 686; A. 199, 147).—2. By heating furonic acid with HIAq and P at 200° (Baeyer, B. 10, 1358).— 3. By heating pentane w-tetra-carboxylic acid (Perkin, jun., C. J. 51, 242; 59, 825; B. 18, 3249).—4. By oxidation of myristic acid with HNO₃ (Noerdlinger, B. 19, 1898).—5. By reducing OO(CH₂.CH₂.CH₂H)₂ (cf. FURFURYL-ACRYLIG ACID) (Marckwald, B. 21, 1398).

Properties. - Rectangular trimetric plates (from water), v. sol. alcohol, ether, and hot benzene. May be sublimed. NaOEt forms C₁₇H₂₈Na₂O₈, a white powder, insol. ether, sol. water. Not converted into anhydride by AcCl or by PCl, (Volhard, A. 267, 82).

Salts. - BaA"aq. - CaA". Deposited as granular powder when a cold saturated solution

is heated.— Ag₂A": white pp.

Ethyl ether Et₂A". S.G. \$ 1.0080; \(\frac{15}{28} \)

9988; \(\frac{26}{28} \)

9920. M.M. 11.424. Oil with pene-

trating odour (Perkin, C. J. 59, 826).

Pimelic acid $C_1H_{12}O_2$. [103° cor.]. Got by heating chelidonic acid with HIAq at 205° (Haitinger a. Lieben, M. 5, 358). Monoclinic tables, v. sol. hot benzene. Perhaps identical with n-pimelic acid.

(8)-Pimelic acid $C_5H_{10}(CO_2H)_2$. [106°]. Occurs among the products of oxidation of castor oil and earth-nut oil with nitric acid (Gantter a. Hell, B. 17, 2212). Large tables (from water). Readily forms supersaturated solutions.

BaA"aq: plates.—PbA".—CuA".—Ag₂A": pp.
Iso-pimelic soid C₂H₁₀(CO₂H)₂. [104°].
Formed from amylene bromide by successive treatment with alcoholic KOy and HClAq at 170° (Bauer, M. 4, 345; Hell, B. 24, 1889). Prisms, v. sol. water and alcohol. Begins to form an anhydride at 135°.—(NH_a)₂A".—BaA" 1½aq: sandy powder.—CaA". S. 2 at 22°; 14 at 100°.—SrA"4aq.—NiA" 2½aq.—ZnA".—PbA". S. 01 at 100°.—CuA"aq. S. 04 at 16°; 07 at 100°. Bluish-green plates. - CdA"2aq: needles. S.:12 at 15°; 13 at 100°.—Ag₂A".

An amorphous pimelic acid was got by Bauer, as well as the preceding acid, from amylene bromide. Its Ca salt was more soluble

Pimelic acid C₇H₁₂O₄. [87°]. Got by oxidising menthol with KMnO4 in acid solution (Arth, A. Ch. [6] 7, 455; C. R. 107, 107). Needles, v. sol. water. Can be extracted by ether from its aqueous solution.—Ag₂ A''.

Isomerides - v. DI-METHYL-GLUTARIC ACID, METHYL-ETHYL-SUCCINIC ACID, TRI-METHYL-SUC-

CINIC ACID, and PROPYL-SUCCINIC ACID.

PIMENTO. The volatile oil obtained from the pods and seeds of Myrtus Pimenta contains eugenol and a terpene (255°) S.G. 18 98 (Bonastre, J. Ph. 11, 187; Oeser, A. 131, 277).

PINACOLIC ALCOHOL v. Sec - HEXYL ALCOHOL.

PINACOLIN v. METHYL tert - BUTYL KETONE. Benzpinacolin v. vol. i. p. 488.

PINACONE C. H., O. i.e. CMe. (OH). CMe. (OH). Hexylene glycol. [38°]. (170° cor.). S.G. 15 9672; 25 9609. M.M. 7.245 (Perkin, C. J. 45, 506). H.C. 897,697 (Louguinine, A. Ch. [5] 25, 143). Formed by the action of Na or sodiumamalgam on aqueous acetone (Fittig, A. 110, 25; 114, 54; Städeler, A. 111, 277; Friedel, A. 124, 324; Bl. [2] 19, 289; Linnemann, A. Suppl. 3, 374). Formed also from CMe₂Br.CMe₂Br by successive treatment with AgOAc and baryta (Pawloff, A. 196, 122). Small needles (from CS2), v. sol. cold alcohol and ether, sl. sol. cold CS₂. Sl. sol. cold water, but v. sol. hot water, separating on cooling as a hydrate $C_eH_{14}O_26aq$ [46.5°], crystallising in four-sided tables, decomposed by distillation into water and pinacone.

Reactions.—1. Chromic acid mixture reconverts it into acetone.—2. Gaseous HI yields ₽rI and some hexane (Linnemann, Sitz. W. [2] 63, 255). HIAq at 100° forms CaH, I (Bouchardat, Z. 1871, 699).—3. Boiling dilute H2SO4 converts it into pinacolin. Heating with HOAc has the same effect.—4. POCl, forms di-chloro-hexane [160°] (Friedel a. Silva, B. 6, 35).—5. Does not form an acetal when heated with aldehyde (Lochert, A. Ch. [6] 16, 60).—6. When passed through a red-hot tube it yields acetone and isopropyl alcohol (Thörner a. Zincke, B. 13, 645).

PINACONES. Compounds of the form HO.CRR'.CRR'.OH, where R and R' are alkyls. They are obtained by reduction of ketones. On treatment with dehydrating agents they yield (β)-pinacolins CRR'₂.CO.R or isomeric (α)-pinacolins OCRR' or CRR'.O.CRR'. If in these formulæ R = H, the (β)-pinacolin will be an aldehyde; while if at the same time R' = H, we find that glycol is the simplest pinacone, aldehyde the simplest (3)-pinacolin, and ethylene oxide the simplest (a)-pinacolin (Zincke, A. 216, 296). Some (β) -pinacolins are decomposed by heating with soda-lime or alcoholic potash, thus: PhCX_COPh + H_O = PhCX_H + HOBz (Zinckea. Thörner, B. 11, 65; Zagumenny, J. R. 12, 429); but phenyl (α)-naphthyl (β)-pinacolin reacts with alcoholic potash, thus:

 $PhC(C_{10}H_{7})_{2}OOPh+H_{2}O=PhC(C_{10}H_{7})_{2}OH+Ph.CO.H$ (Elbs, J. pr. [2] 35, 506).

PINE APPLE OIL. Artificial pine-apple oil may be made by dissolving butyric ether in alcohol (Hofmann, A. 81, 87).

PINENE v. TERPENES.

PINEY TALLOW. A fat obtained by boiling the fruits of Valeria indica (of Malabar). Melts at 30° to 38°. Its fatty acids consist of palmitic acid (75 p.c.) and oleic acid (25 p.c.) (Dal-Sie, G. 8, 107)

PINIPICRIN C22H36O11. Occurs in the needles and bark of the Scotch fir (Pinus sylvestris) and in the green parts of Thuja occidentale (Kawalier, Sitz. W. 11, 350; 13, 515). Bright-yellow amorphous powder, beginning to soften at 55°, and quite liquid at 100°. Hygroscopic. Tastes bitter. V. sol. water, sol. alcohol, insol. ether. Boiling dilute H₂SO₄ yields glucose (2 mols.) and ericinol C10H16O (1 mol.), v. vol. ii. p. 458.

PINITANNIC ACID C14H16O8? Occurs in the needles of the Scotch fir and in the green parts of the Arbor vitæ (Thuja occidentale) (Kawalier, Sitz. W. 11, 357; 29, 19). Reddish-yellow powder, v. sol. water, alcohol, and ether. FeCl, colours its aqueous solution brown. Gives yellow pps. with lead acetate and subacetate.

Not ppd. by gelatin. PINITE C7H14O6. Methyl ether of dextrorotatory inosite. [186°]. [a]_D = 65° 51′. Supposed to be extracted by water from the hardened sap of Pinus lambertiana of California (Berthelot, A. Ch. [3] 46, 76; Johnson, Am. S. [2] 22, 6; Combes, C. R. 110, 46; Maquenne, A. Ch. [6] 22, 264). Crystalline crusts, v. e. sol. water, almost insol. alcohol. Tastes sweet. Dextrorotatory. Non-fermentable. Does not reduce Fehling's solution. HIAq splits it up into MeI and dextrorotatory inosite [248°]. Pinite is identical with matezite and sennite.

PINOL $C_{10}H_{16}O$ *i.e.* $C_{6}H_{6}OMePr$. (184°). S.G. $^{20}\cdot 953$; $\mu_{D}=1\cdot 469$. Formed, together with pinene nitroso-chloride when oil of turpentine is treated with HOAc, nitrous ether, and HClAq (Wallach, A. 253, 251; 259, 322; 268, 222; B. 24, 1552). Purified by conversion into the dibromide C₁₀H₁₆Br₂O [94°], which is dissolved in dry benzene and treated with sodium-wire.

Liquid, smelling like cineol.

Reactions.—1. Oxidised by KMnO, to CO, oxalic acid, and terebic acid [176°].-2. Br forms C₁₀H₁₆Br₂O and C₁₀H₁₅Br₃O [160°]. -- 3. HBr forms an addition product converted by water into the hydrate C₁₀N₁₀OH₂O [131°] which is not attacked by Ac₂O, but is reconverted into pinol by warming with dilute H2SO4, and yields terpenylic acid [57°] on oxidation by KMnO.

Dibromide C₁₀H₁₆Br₂O. [94°]. (144° at 11 mm.). Trimetric crystals; a:b:c = .570:1:1.555. Insol. water, volatile with steam. Converted by alcoholic potash into 'pinol glycollic ether C₁₀H₁₆(OEt)₂O [53°] (c. 115° at 14 mm.). AgOAc forms C₁₀H₁₆(OAc)₂O [98°] (127° at 13 mm.), which on saponification by hot dilute H₂SO₄ yields 'pinol-glycol' C₁₀H₁₆(OH)₂O [125°], crystallising in matted needles, v. e. sol. chloroform. Boiling with water and Pb(OH), also converts the dibromide into the glycol. Silver propionate yields C10H18(O.COEt)2 [106°]. Formic acid at 100° reduces the dibromide to cymene.

Nitroso-chloride C10H16ONOCl. [103°]. Formed by the action of amyl nitrite and HCl on pinol in HOAc (Wallach, A. 253, 261). Converted by alcoholic NH, into the nitrolamine C₁₀H₁₆ONONH₂ (130° at 14 mm.), a viscid mass which yields a crystalline hydrochloride, B'HCl. Aniline, piperidine, and benzylamine form the three analogous bodies $C_{10}H_{10}O.NO.NHPh~[175°]$, $C_{10}H_{10}O.NO.NHC_3H_{10}~[154°]$, and the compound $C_{10}H_{10}O.NO.NHC_4H$, [136°], each of which forms a crystalline hydrochloride. (β)-Naphthylamine forms, in like manner, pinol-nitrol-naphthylamine $C_{10}H_{14}O.NO.NHC_{10}H_{7}$ [195°].

PINYLAMINE C₁₀H₁₈NH₂. (208°). S.G. 17 943. Formed by reducing with zinc-dust and HOAc nitroso-pinene $C_{10}H_{18}NO$ [132°], which is got from $C_{10}H_{18}NO$ Cl and alcoholic NaOH (Wallach a. Lorenz, A. 268, 197; B. 24, 1550). Oil, turning yellow in air, and giving off NH₂. Absorbs CO₂ from air, forming a solid carbonate Benzoic aldehyde forms C₁₀H₁₈N:CHPh [53°] furfuraldehyde gives C₁₀H₁₈N:CH.C₄H₂O [81°] whilst salicylic aldehyde forms the compound C₁₀H₁₅N:CH.C₅H₄.OH [109°].—B'HCl. [230°]. Needles (from water). Yields cymene on distillation.—B'₂H₂PtCl₆. Yellow plates or needles.— B'HNO₃. Crystals, sl. sol. cold water.—B'₂H₂SO₄.
—B'HCyS. [136°]. Prisms (from water).—

B'₂H₂C₂O₄. [248°]. Scales. Acetyl derivative C₁₀H₁₈NHAo. [109°] Benzoyl derivative. [125°]. Needles.

PINYL-UREA NH2.CO.NHC10H13. Formed from pinylamine hydrochloride and potassium cyanate (Wallach, A. 268, 204). Needles, v. sol. alcohol.

PIPECOLINE v. METHYL-PYRIDINE HEXA-HYDRIDE.

PIPERAZINE v. PYRAZINE HEXAHYDRIDE. PIPERHYDBONIC ACID v. Methylene derivative of DI-OXY-PHENYL-VALERIC ACID.

PIPERIC ACID C12H10O4 i.s.

CH₂<0>C₆H₃.CH:CH.CH:CH.CO₂H. Mol. w. [217°]. S. (alcohol) .37 in the cold; 2 at 78°. Formed, together with piperidine, by boiling piperine with alcoholic potash (Von Babo; Strecker, A. 105, 317; 118, 280; G. C. Foster, C. J. 15, 17; Fittig a. Mielk, A. 152, 25; 172, 134). Yellowish needles (from alcohol). After fusion it melts at 213°. Nearly insol. water, m. sol. hot alcohol and ether.

Reactions.—1. Reduced by sodium-amalgam to two hydropiperic acids, which are the methylene derivatives of di-oxy-phenyl-angelic acids CH₂O₂:C₆H₄.CH₂.CH:CH.CH₂.CO₂H₄ [78°] and CH2O2:C4H2.CH2.CH2.CH:CH.CO2H[131°] (Regel, B. 20, 414). -2. Dilute alkaline potassium permanganate at 4° oxidises piperic acid to piperonal CH2O2:C4H3.CHO and racemic acid (Doebner, B. 23, 2375).-3. Potash-fusion yields protocatechuic, oxalic, and acetic acids.-4. Bromine in CC, forms a tetrabromide which decomposes at 160°-165°.

- NH, A'. Satiny scales. - KA'. -Salts. -BaA'. S. 02 in the cold.—AgA': powder. Ethyl ether EtA'. [78°]. Plates. Reference. - BROMO-PIPERIC ACID.

PIPERIDEINE. A name for Pyridine Tetra-

Dipiperideïne C₁₀H₁₈N₂ i.e. [61°]. Formed by slowly adding v-chloro-piperidine to a boiling 10 p.c. solution of potash in alcohol (Lellmann a. Schwaderer, B. 22, 1000, 1318). Monoclinic crystals, sl. sol. water, v. sol. alcohol and ether. Between 200° and 230° it begins to dissociate into (2 mols. of) pyridine tetrahydride; hence it has no constant boiling-point. Slightly volatile with steam. Ac₂O forms an acetyl derivative (220°). S.G. ^{16:5}/₄ 1·0531. Tin and conc. HClAq reduce it to piperidine. Phenyl thiocarbimide gives $C_{10}H_{11}N_{2}$ CS.NHPh [144°]. CS₂ forms C₁₀H₁₈N₂CS₂, crystallising in needles, and melting at 150° with evolution of gas.

 $Salt.-B''H_2Cl_22aq.$ [150°]. Plates, liquefies c. 80° when quickly heated.

Isodipiperideine C₁₀H₁₈N₂. Formed by heating diazobenzene piperidide at 250° (Heusler, A. 260, 239). Thickish liquid, miscible with water, but can be extracted therefrom by ether. Reduces Fehling's solution and ammoniacal AgNO. Its salts are hygroscopic.

PIPERIDIC ACID C₄H₉NO₂ [184°], which is got by oxidising piperidine v-carboxylic ether with fuming HNO, and heating the product with HClAq at 140°, is γ-amido-butyric acid (Schotten, B. 16, 643; Gabriel, B. 23, 1770).

PIPERIDINE C, H, N i.e.

CH₂CH₂CH₂NH. Pyridine hexahydride. Mol. w. 85. (106°). S.G. 15 ·8664; 25 ·8591. M.M. 5·810 (Perkin, C. J. 55, 700). S.V. 108·76. S.H. ·533 (Colson, Bl. [3] 3, 8). H.F.v. 24,090. H.F.p. 26,990 (Thomsen, Th. 4, 145). Appears to occur in the husks of pepper (Johnstone, C. N. 58, 235; An. 14, 41).

Formation.—1. By distilling piperine with

potash-lime or soda-lime, or by boiling it with alcoholic potash (Wertheim, A. 127, 75; Anderson, A. 75, 82; 84, 345; Cahours, A. Ch. [3] 38, 76; Von Babo a. Keller, J. pr. 72, 53). 2. By reduction of pyridine in alcoholic solution by sodium-amalgam; the yield being 75 p.c. of the theoretical (Ladenburg a. Roth, A. 247, 51). 3. By heating pentamethylene-diamine hydrochloride (Ladenburg, B. 18, 3100).—4. By heating hygric soid C₄H₁₁NO₂ with H₂SO₄ for a few minutes at 300° (Liebermann a. Kühling, B. 24, 413).

Properties.-Liquid, with ammoniacal and peppery smell, miscible with water. Alkaline in reaction. Caustic taste. Ppts. salts of zinc and copper, but does not redissolve the ppd. hydrates. Not attacked by fuming HClAq at 300° or by boiling HNO₃ (S.G. 1.5). Piperidine acts as an ansethetic (B. 14, 713). Oxidised by H_2O_2 to glutaric acid and its imide and δ-amido-valeric aldehyde (Wolffenstein, B. 25, 2777).

Reactions.—1. Bromine and water at 200° yield di-bromo-oxy-pyridine and some CHBr, (Hofmann, B. 12, 984). Piperidine hydrochloride, heated with dry Br at 180, yields dibromo-pyridine (Schotten, B. 15, 427). Bromine and NaOHAq give crystalline C.H.Br.NO (Hofmann, B. 16, 560). Bromine and lime-water yield brome-piperidine C₈H₁₀BrN [234°] (Lell-

mann, B. 22, 1327). -2. Chlorine acts with explosive violence on dry piperidine. In presence of water or chloroform it yields v-chloro-piperidine, which may also be got by using bleachingpowder. It is an unstable, heavy oil (52° at 25 mm.), with pungent odour, and deposits piperidine hydrochloride when kept for some time (Bally, B. 21, 1772).—3. Conc. H₂SO, at 300° forms pyridine (Koenigs, B. 12, 2341). Nitrobenzene at 260° also oxidises it to pyridine (L.). 4. Chloroform, on boiling for some days, yields $CH(C_5H_{10}N)_8$ aq (98° at 15 mm.) (Busz a. Kekulé, B. 20, 3246)...5. CH₂CCl₃ forms, on boiling, ethenyl-tripiperidine CH₂.C(C₅H₁₆N)₃, a liquid (262°) yielding B"H₃Cl₃ and B"'₂3H₂PtCl₅ (B. a. K.)...6. Boiling oxalic ether (1 mol.) yields piperidyl-oxamic ether C, H₁₀N.CO.CO.Et (289°) (Wallach, A. 214, 278; 237, 247). The corresponding acid [129°] splits up on fusion into CO₂ and the formyl derivative of piperidine. PCl, yields CO₂ and C₅H₁₀N.COCl (238°), which acts upon piperidine forming (C₅H₁₀N)₂CO[43°] (298°). NH, Aq converts the ether into piperidyl-oxamide $C_3H_{10}N.CO.CO.NH_2$ [127°], crystallising in monoclinic prisms, converted by P_2O_3 into $C_3H_{10}N.CO.CN$, a heavy oil (264°). When piperidine (2 mols.) is distilled with oxalic ether (1 mol.) there is formed oxalyl-piperidine $C_bH_{10}N.CO.CO.C_bH_{10}N$ [89°] (above 350°). — 7. By heating with phthalic acid (2 mols.) as long as phthalic anhydride sublimes there is formed 'piperilene-amine-phthalein' C18H21N2O2 or $C_6H_4 < \frac{C(NC_6H_{10})_2}{CO} >$, an oil which yields crystalline $C_{18}H_{24}Br_2N_2O_2$ (Piutti, G. 13, 535; A. 227, 197).—8. Phthalic anhydride unites in the cold with piperidine, forming a mixture of ' piperilene-phthalamic acid' C5H10N.CO.C6H1.CO2H and its piperidine salt (P.). On shaking with ether and water the acid goes into ethereal solution, and the salt into aqueous solution. Piper-ilene-phthalamic acid is a heavy oil, v. sol. alcohol. It yields AgA', C₅H₁₁NHA' [c. 150°], and C18H28Br4N2O3, crystallising in long needles. 9. Alloxan and aqueous sulphurous acid give (C₄H₂N₂O₄)C₄H₁,NH₂SO₃, crystallising in plates (Pellizzari, A. 248, 150).—10. On heating with isatin and alcohol on the water-bath there is formed the compound CsH5NO(CsH10N)2 or C₈H₄ C(NC₈H₁₀)₂, crystallising in flat colourless prisms, m. sol. hot alcohol, turned red by HCl, and yielding a blue dye (indigo?) when treated with Ac2O, or when rapidly heated to 125°-160°. Bromo-isatin forms, in like manner, C₈H₄BrNO(C₅H₁₀N)₂, crystallising in needles, while di-bromo-isatin reacts with production of $C_0H_2Br_2(NH_2).CO.CO.NC_0H_{10}$ [1525] (Schotten, B. 24, 1367, 2605).—11. A solution of diazobenzene chloride and sodium acetate forms Ph.N.N.NC, H10, which yields phenol and piperidine when treated with dilute H2SO4, and phenyl-hydrazine and piperidine when reduced by SnCl, and HCl (Nölting a. Binder, B. 20, 3016). Other diazo-compounds act in like manner. Diazobenzene piperidide is decomposed on heating to 250°, yielding benzene, N, and isopiperidein (Heusler, A. 260, 239).—12. Potassium cyanate converts piperidine sulphate into NH₂.CO.NC₅H₁₀, crystallising from alcohol in needles (Cahours). Methyl and ethyl cyanates

yield corresponding ureas. Phenyl cyanate forms $NHPh.CO.NC_5H_{10}$, crystallising from alcohol in prisms [172°]. C,H,NH.COCl gives rise to C,H,NH.CO.NC,H10 [102°], crystallising from dilute alcohol in needles (Kühn a. Riesenfeld, B. 24, 3818).—13. Methyl thio-carbimide forms NHMe.CS.NC₅H₁₀ [129°] (Hecht, B. 23, 287; 25, 815), while ethyl-, propyl-, and phenyl-thiocarbimides form corresponding thio-ureas [46°], [75°], and [99°] respectively. Phenyl-, o-, and p-tolyl-thiocarbimides form C₃H₁₀N.CS.NHPh [98°] (G.); [104°] (Skinner a. Ruhemann, C. J. 53, 558) and C₅H₁₀N.CS.NHC₇H₇ [98°] and [132°] respectively. Potassium sulphocyanide reacts with piperidine sulphate, forming C₅H₁₀N.CS.NH₂ [92⁵] Gebhardt, B. 17, 3039). Allyl-thiocarbimide forms oily C₅H₁₀N.CS.NHC₃H₅, which is converted by conc. HClAq at 100° into $C_sH_{10}N.C \leqslant \stackrel{S.CHMe}{N.CH_2}$, a liquid (277°) yielding $B'C_eH_3N_3O_7$ [112°] and B'MeI [67°] (Avenarius, B. 24, 262).—14. Benzoyl-thiocarbimide added to a solution of piperidine in dry benzene forms C₅H₁₀N.CS.NHBz [123°], crystallising in needles, sol. alcohol and ether (Dixon, C. J. 55, 624). Benzyl-thiocarbimide gives C₅H₁₀N.CS.NHCH₂Ph [88°] (Dixon, C. J. 59, 568). 15. Tri-methyl-trithiocyanurate at 200° forms $C_{14}H_{24}N_{5}S$ [107°], which gives $B'_{2}H_{2}PtCl_{8}$ (Hofmann, B. 18, 2779). 16. Quinone forms red needles [178°], which are probably $C_6H_2O_2(NC_4H_{10})_2$ (Lachovitch, M.~9,~506).—17. Benzoic aldehyde in presence of K_2CO_3 forms CHPh(NC,H10)2 [80°] crystallising from alcohol in flat needles (Ehrenberg, J. pr. [2] 36, 130; Lachovitch, M. 9, 695).—18. Tri-oxy-methylene forms $CH_2(NC_4H_{10})_2$ (230°), which unites with CS_2 giving $C_{11}H_{20}N_2CS_2$ [58°]. — 19. Pinene nitroso-chloride in alcoholic or aqueous solution yields crystalline C10H16NONC5H10 [119°] which forms B'HCl. The corresponding terpine and dipentinene derivatives melt at 154° (Wallach, A. 241, 320; 245, 253). -20. Amylene nitrosonitrate in boiling alcoholic solution forms $CMe_2(NC_6H_{10}).CMe:NOH$, which crystallises from ether in prisms [96°], and is converted by boiling dilute H₂SO₄ into CMe₂(NC₂H₁₀).CO.CH₃, an oil (220°), $\mu_D = 1.934$, volatile with steam, yielding a very hygroscopic hydrochloride (Wallach, A. 248, 172).—21. Fluorescein chloride at 220° forms $\begin{array}{c} C_{6}H_{4}:C_{2}O_{2} < \begin{array}{c} C_{6}H_{4}(NC_{0}H_{10}) \\ C_{6}H_{3}(NC_{0}H_{10}) \end{array} > O \quad \text{ which } \quad \text{yields} \\ B''H_{2}PtCl_{6} \text{ and } B''H_{2}Cl_{2} \text{ which } \text{ is purple in dilute} \end{array}$ solution and yellow in conc. HClAq (Lellmann a: Büttner, B. 23, 1387).—22. Bromo-phenanthrene at 260° forms crystalline C₁₄H,NC₅H₁₀ [113°] which yields B'₂H₂PtCl₅ 6aq (L. a. B.).—23. Bromo-anthracene at 260° reacts forming C, HoNC, H, crystallising from ether in yellow prisms and giving B'2H2PtCl6 2aq (L. a. B.).-24. Chloro-acetic acid forms C, H, N.CH, CO, H, which forms hemihedral prisms (containing aq) and yields CuA', 4aq, HA'HCl, HA'HBiI,, (HA'), 3HAuCl, and HA'BaCl, (Kraut, A. 157, - 25. a-Chloro-propionic ether produces C.H. N.CHMe.CO.H crystallising from water and alcohol in prisms and yielding HA'HAuCl₄ (Brühl, B. 9, 34).—26. Benzene sulphochloride and NaOHAq form C₅H₅.SO₂.NC₅H₁₀ [93°] (Hinsberg, A. 265, 182; Schotten a. Schlömann, B. 24, 3689). This body is oxidised by KMnO. C.H.SO.NH.CH.CH.CH.CH.CO.H.

27. Picryl chloride forms C₅H₁₀N.C₆H₂(NO₂), [106°] (S. a. S.).—28. Glycerin dichlorhydrin forms liquid $C_{10}H_{22}N_2O$ (280°-290°) which yields B"H₂PtCl₆ (Ladenburg, B. 14, 1879).—29. CS₂ forms (C,H1,N)2CS2 or C,H10N.CS.SNC4H12 crystallising in slender monoclinic needles [174°], converted by an alcoholic solution of iodine to $(C_5H_{10}N.CS)_2S_2[130^\circ].-30$. Dicyandiamide reacts on 'piperyl-biguanide' with formation of C,H,,N,, i.e. C. H₁₀N.C(NH).NH.C(NH).NH₂ [163°?] crystallising in silky needles, and yielding the salts B'H₂Cl₂ [217°], B'H₂SO₄ [173°], B'₂H₂SO₄ [219°], B'H₂PtCl₅ [252°], B'H₂AuCl₅, and the copper compound Cu(C,H₁,N₅)₂H₂SO₄ crystallising in rose-red needles (Bamberger, B. 24, 605, 904). Piperyl-biguanide is converted by CHCl, and caustic potash into 'piperyl-formoguanamine' the salts B'HCl [201°], B'₂H₃SO₄ aq [222°], B'2C₆H₂N₃O₇ [188°], B'₄H₂PtCl₆ [219°], B'HAuCl₄ [90°] and [158°], and B'₂AgNO₅ [229°-238°] (Hjelt, B. 25, 529). Piperyl-biguanide sulphate heated with NaOAc at 200° yields piperyl-aceto- $C_sH_{10}N.C \leqslant_{N:CMe}^{N.\tilde{C}(NH_2)} \searrow N$ guanamine [179°] (Hjelt, B. 25, 533). The acetyl derivative of piperyl-biguanideC₃H₁₀N.C(NAc).NH.C(NH).NH₂ [193°] is got by heating the copper salt with HOAc.

Salts.—B'HCl. [237°]. M.M. 10.034 in a 53 p.c. solution (Perkin, C. J. 55, 716). Needles, v. sol. water and alcohol.—B'HAuCl, [206°]. Not decomposed by boiling water (De Coninck, Bl. [2] 45, 131).—B'2H2PtCl3. [196°] (Ladenburg, B. 18, 3100); [200°] (Wallach a. Lehmann, A. 237, 241). Red needles. Not decomposed by boiling water (De Coninck, Bl. [2] 45, 131). Crystallises also with EtOH in orange needles [191°].—B'2PtCl2.— B'Zn₂OCl₂: amorphous pp. got by adding piperidine to a solution of ZnCl2 (Lachovitch, M. 9, 517).—B'HBr. Plates (Lellmann, B. 20, 680).-B'HI. Long needles.—(B'HI), 2BiL. Scarl (from alcohol) (Kraut, A. 210, 319) .plates B'ICl. [148°]. White needles. — B'ICIHCI. Yellow crystals (from water) (Pictet a. Krafft, Bl. [3] 7, 72). -B'HNO3. Small needles. -B'₂H₂C₂O₄: needles.—B'₂H₄FeCy₆ 3aq. Yellow triclinic crystals.—Piperate B' $C_{p}H_{p}O_{s}$.
laminæ [100°] (Babo a. Keller).—B'₂H₂SO_s.

Nitrosamine C₄H₁₀N.NO. (218°). Silkv

Nitrosamine C₅H₁₀N.NO. (218°). S.G. 19:5 1-066. Formed from piperidine and nitrous acid (Wertheim, A. 127, 75; Schotten, B. 15, 425; Knorr, A. 221, 298). Pale-yellow liquid, sl. sol. water, v. sol. conc. HClAq, but reppd. on dilution. Zinc and HClAq reduce it to piperidine and NH₈. Heated in a current of HCl at 100° it gives piperidine and NOCl. Sodium-amalgam forms C₅H₁₀N.NH₂ and piperidine.—B"H₂Cl₂: syrupy.—*B"HCl: crystalline mass.

Hydrazine C.H.₁₀N.NH.. 'Piperyl-hydrazine.' (146°) at 728 mm. S.G. 146° 928. Got by reducing the nitrosamine with zinc-dust and HOAc (Knorr, B. 15, 859; A. 221, 299). Liquid, with ammoniacal odour, miscible with water, alcohol, and ether. Volatile with steam. Oxidised by HgO in the cold to the tetrazone C₁₀H₂₀N. [45°], which yields B'₂H₂PtCl₆. Reduces cold ammoniacal AgNO₃ and hot Fehling's solution. Nitrous acid converts it into piperidine. CS₂ gives rise to (C₆H₁₀N.NH)₂CS [181°]. The compound

 $C_3H_{10}N.NH.CS.NC_3H_{10}[86^\circ]$ may also be obtained. BzOl in ether forms $C_3H_{10}N.NHBz$ [196°]. Benzoic aldehyde forms C₅H₁₀N.N:CHPh [63°]. Salt.—B'HCl. [162°]. Tables (from alcohol). Reacts with potassium cyanate forming $C_sH_{10}N.NH.CO.NH_2$ [136°] and with potassium sulphocyanide forming $C_sH_{10}N.NH.CS.NH_2$ [167°].—Methylo-iodideB'MeI.[215°].Begins to decompose at 150°.

Formyl derivative C, H10 N.CHO. (222°). S.G. ²³ 1-0193. Formed by the distillation of C.H., N.CO.CO.H (v. Reaction 6) (Wallach a. Lehmann, A. 228, 251; 237, 252). Got also by heating piperidine with formamide (Lachovitch, M. 9, 699). Liquid, miscible with water, alcohol, and ether. PCl₅ yields a base C₁₁H₂₀N₂.—B'HCl: deliquescent needles. — B'4H2PtCle. [172°].

Plates.—B'HgCl₂. [149°]. Needles.

Acetyl derivative C₅H₁₀NAc. (227°). S.G. 2 1.011 (Wallach a. Kamensky, A. 214, 238; cf. Schotten, B. 15, 426). Liquid, miscible with water. When heated with BzCl it gives AcCl and benzoyl-piperidine (Pictet, B. 23, 3014). On heating with bromine it gives pyridine, bromo-pyridine, and di-bromo-pyridine (Hofmann, B. 16, 587). When Cl is passed into cooled acetyl-piperidine and the product is warmed with water, there is formed $C_rH_{11}Cl_2NO_2$ [122°] (Bally, B. 21, 1772).

Benzoyl derivative C_aH₁₀NBz. [48°]. (above 360°). Formed from piperidine, BzCl, and NaOHAq (Cahours; Schotten, B. 17, 2544; 21, 2238). Triclinic prisms (from alcohol). Oxidised by KMnO, to benzoyl-5-amido-valeric acid [94°]. Benzoic aldehyde forms the compound NH C(CHPh).CH₂ CH₂ [89°], which yields B'HCl [166°] and B'HNO, [98°], both being crystalline (Rügheimer, B. 24, 2186).

Bromo-bensoyl derivative C.H.o.N.CO.C.H.Br. The o-compound is an oil, while the p-compound crystallises from alcohol in rectangular monoclinic tables [95°] (Schotten, B. 21, 2248).

m-Nitro-bensoyl derivative C₅H₁₆N.CO.C₅H₄NO₂. [34°]. (184° at 54 mm.). S. (alcohol) 33 in the cold, 50 at 78°. Darkyellow monoclinic crystals. Crystallises from water with about 5aq, and then melts at 84°. Yields, on reduction, the m-amido-benzoyl derivative [125°].

Oxy-bensoyl derivative v. Piperidide

of OXY-BENZOIC ACID.

Cuminyl derivative C_{1b}H₂₁NO. Tables. Cinnamyl derivative C₆H₇CO.NC₅H₁₀. [122°]. Formed from the anhydride and piperidine (Herstein, B. 22, 2265). Stellate needles.

Alkyl-piperidines. On heating the alkyloiodides of pyridine, a pair of alkyl-pyridines is produced, that of lower boiling-point being (a)-alkyl-pyridine, and the other (γ) -alkyl-pyridine. On reduction these give the corresponding alkyl-piperidines. The same isopropyl-pyridines are obtained both from pyridine isopropylo-iodide, and pyridine n-propylo-iodide, an intramolecular change taking place in the propyl group in the latter case. Conyrine is (a)-propyl-pyridine, coniine is (a)-propyl-piperidine (Ladenburg, B. 18, 1587). By adding potash to piperidine alkylo-iodides, *-alkylpiperidines can be obtained.

Methyl-, Ethyl-, Propyl-, Phenyl-, and Tolyl-Piperidines v. METHYL, ETHYL, PROPYL, PHENYL, and Tolyl Pyridine HEXAHYDRIDES.

PIPERIDINE y-CARBOXYLIC ACID

CH2CH2N.CO2H. Methyl ether MeA'. (201°). Formed from piperidine, ClCO2Me, and KOHAq (Schotten, B. 15, 425; 16, 647). Heavy oil. Converted by HNO, containing urea, into C₃H₇(NO₂)N.CO₂Me [103°], whence Br forms a compound [180°].

Ethyl ether EtA'. Piperyl - urethane. (211°). Formed from piperidine and ClCO₂Et. Oil. Br in HOAc forms C₅H₇BrN.CO₂Et [140°], while HNO, free from NO, forms 'nitrodehydropiperyl urethane 'C,H,(NO2)N.CO2Et [52°],

whence Br in HOAc forms C₃H_{1,3}BrN₂O₅ [157°].

Chloride C₅H₁₀N.COCl. (238°). Formed from C₅H₁₀N.CO.CO₂H and PCl₅ (v. PIPERIDINE, Reaction 6). Liquid, slowly decomposed by cold water into piperidine, CO2, and HCl.

Amide C₅H₁₀N.CO.NH₂. [106°]. Got from piperidine sulphate and potassium cyanate. Needles. Yields B'HNO₃. [67°] (Franchimont a. Klobbie, R. T. C. 8, 302).

Anilide. [172°]. Formed from piperidine and phenyl cyanate, or from the chloride and aniline (Gebhardt, B. 17, 3040; Wallach, A. 228, 250; 237, 250).

Piperidide (C,H₁₀N)₂CO. [43°]. Isomeride v. Hexahydride of Pyridine

CARBOXYLIC ACID.

DI-PIPERIDYL v. DIPYRIDYL dodecahydride.

PIPERIDYL-CYANURAMIDE v. Cyanuramide in article Cyanic acids

PIPERIDYL-MELAMINE v. CYANIC ACIDS. PIPERILENE v. vol. iii. p. 807.

PIPERINE C, H, NO, i.e.

C, H, N.CO.CH:CH.CH.CH.C. H, O.CH. Piperylpiperidine. Mol. w. 285. [128°]. Occurs in black pepper (Piper nigrum), long pepper (P. longum), and in the black pepper of Western Mrica (Cubeba Clusii) (Oersted, S. 29, 80; Pelletier, A. Ch. [2] 16, 344; 51, 199; Merck, N. J. T. 20, 1, 34; Wackenroder, Br. Arch. 37, 347; Duflos, S. 61, 22; Warrentrapp a. Will, A. 39, 283; Wertheim, A. 70, 58; Gerhardt, Compt. Chim. 1849, 375; A. Ch. [3] 7, 253; Anderson. 4 75, 82; 84, 345; Cabourg A. Ch. [8] 88, 76. A. 75, 82; 84, 345; Cahours, A. Ch. [3] 88, 76; Stenhouse, A. 95, 106; Von Babo a. Keller, J. pr. 72, 53; Strecker, A. 105, 317).

Formation.—By heating piperidine with the chloride of piperic acid (Rügheimer, B. 15, 1390).

Preparation.—Ground pepper (1 pt.) is boiled with slaked lime (2 pts.) and water, the filtrate evaporated to dryness at 100°, and the piperine extracted with ether and recrystallised from alcohol (Cazeneuve a. Caillot, Bl. [2] 27, 290).

Properties.—Monoclinic prisms, v. sl. sol. hot water, m. sol. alcohol and ether. Inactive to light. Insol. dilute acids and alkalis. Decomposed by alcoholic potash into piperic acid and piperidine. Conc. H₂SO₄ forms a blood-red solution. HNO₃ gives a greenish-yellow colour changing to red. Phosphomolybdic acid gives a flocculent pp.

Salts .- The hydrochloride is crystalline, but decomposed by water.—B',H,PtCl: roseate monoclinic crystals. Not decomposed by hot water (De Coninck, Bl. [2] 45, 131).—B'2HHgCl3: triclinic crystals. $-B'_2HI_2$. [145°]. needles (Jörgensen, J. pr. [2] 3, 328). Steel-blue

PIPEROKETONIC ACID v. DI-OXY-BENZYL

ETHYL KETONE CARBOXYLIC ACID.

PIPERONAL v. Methylene derivative of PROTOCATECHUIC ALDEHYDE.

PIPERONYL-ACRYLIC ACID v. Methylene ether of CAFFEIG ACID.

PIPERONYL ALCOHOL v. Methylene deri-

vative of DI-OXY-BENZYL ALCOHOL. PIPERONYLIC ACID v. Methylene deriva-

tive of Protocatechuic acid. PIPEROPROPIONIC ACID v. DI-OXY-PHENYL-PROPIONIC ACID.

PIPER-PROPYL-ALKINE v. OXY-PROPYL-

PIPERIDINE

PIPERYL. This name is given to radicle CH₂O₂:C₆H₃.CH:CH.CH:CH.CO. Piperyl has also been used to denote C₅H₁₀, the divalent radicle which is united to NH in piperidine; many of the derivatives of this divalent 'piperyl' are described under PIPERIDINE.

PIPERYLENE v. PENTINENE.
PIPERYLENE TETRABROMIDE v. Tetra-

BROMO-PENTANE

PIPITZAHOÏC ACID C15H20O3. Perezone. [104°] (A. a. L.); [107°] (Mylius, B. 18, 480). May be extracted by alcohol from Pipitzahuac root or Radix Perezia (De la Sagra, C. R. 42, 873, 1072; Weldt, A. 95, 188; Anschütz a. Leather, C. J. 49, 715; B. 18, 709, 715). Flat golden plates (from alcohol), nearly insol. water, v. sol. alcohol and ether. May be sublimed. Volatile with steam. Aqueous alkalis form a violet solution. Aqueous methylamine forms methyl - amido - pipitzahoïc acid C₁₅H₁₉(NHMe)O₃ [114°], crystallising in blue needles, while aniline forms C18H19(NHPh)O3 [139°], o-toluidine gives $C_{13}H_{19}(NHC_7H_7)O_6$ [110°], p-toluidine yields p-tolyl-amido-pipitza-hoic acid [134°]. Bromine forms C₁₅H₂₀Br₂O₅ [109°].

Salts.-PbC₁₅H₁₈O₂.-CuA'₂.-AgA':purple

Ethyl ether EtA'. [141°].

Acetyl derivative C16H19AcO8. [115°].

Trimetric plates; a:b:c = 629:1: 845.

Oxim? C15H21NO3. Amidopipitzahorc acid. [154°]. Flat violet-brown needles. Formed from the acid and alcoholic hydroxylamine (Mylius, B. 18, 936). The same body is got by boiling phenyl-amido-pipitzahoïc acid (v. supra) with NH₂Aq (A. a. L.).

Oxypipitzahoïc acid $C_{15}H_{18}(OH)_2O_2$. [129°] (A. a. L.); [134°] (M.). Formed by warming phenyl-amido-pipitzahoïc acid (v. supra) with alcoholic H₂SO₄. Orange plates, nearly insol. water. Its alkaline solutions are violet. Bromine forms an unstable dibromide [140°-146°]. On warming with H2SO4 it is converted into perezinone C₁₅H₁₈O₈ [144°], which crystallises in yellow needles or prisms, and yields C₁₅H₁₇NaO₃, crystallising in easily soluble yellow tables.

PIRYLENE C.H. i.e. CH CH.CH. Formed by distilling with solid NaOH the methylo-iodide of the base obtained by the action of Ag₂O on the iodide formed from di-methylpyridine hexahydride and iodine (Ladenburg, A. 247, 60). Oil. Does not ppt. ammoniacal Cu₂Cl₂

PISCIDIN C₂₈ H₂₄O₈. [192°]. The poisonous principle of Jamaica dogwood (*Piscidia* Erythrina), from which it can be extracted by lime-water (Hart, Am. 5, 39). Six-sided prisms (from alcohol), insol. water, sl. sol. cold alcohol, sol. benzene, CHOl, and conc. HClAq. Sedative and narcotic.

PITTAKAL v. EUPITTONIC ACID.

PITURINE is NICOTINE.

PLASMINE v. PROTEÏDS.

PLATINAMMINES v. PLATINUM-AMMONIUM COMPOUNDS, p. 292.

PLATINATES. Pto, H, acts as an acidic hydroxide towards strong bases, forming compounds of the type xPtO.yMO. These platinates are generally obtained by reacting on PtCl, Aq with caustic or carbonated alkalis, and sometimes by fusing PtCl, with bases and then

washing with water.

Barium platinates. \mathbf{The} compound 2PtO2.3BaO was obtained, in hexagonal crystals, by Rousseau (C. R. 109, 144) by heating PtCl4 with BaO for some time, then adding BaCl, equal to the quantity of BaO used, and heating to c. 1100° (m.p. of Cu) in a Pt dish for some hours, and washing with water. The crystals are insol. acetic acid, but sol. HClAq. The salt decomposes at an orange-red heat, in presence of BaCl₂, with separation of Pt. Topsöe (B. 3, 462) obtained PtBaO₂.4H₂O by decomposing H2PtCl8Aq by excess of BaO,H2 in sunlight; to the pp. thus obtained Johannsen (A. 155, 204) gave the composition

3PtBaO₃.BaCl₂.PtOCl₂.11H₂O.

Calcium platinates. According to Herschel (A. 3, 317), CaOAq added to PtCl₄Aq in sunlight produces a white pp. of

PtCaO₃.CaO.PtOCl₂.7H₂O.

Sodium platinates. When a mixture of Na₂CO₃Aq and H₂PtCl₆Aq stands for some days in a warm place, a pp. of 3PtO₂.Na₂O.6H₂O separates (Döbereiner a. Weiss, A. 14, 21). Rousseau (C. R. 109, 144) obtained crystals of Na platinate by heating a mixture of equal weights of NaOH and NaCl with some Pt black, in a Pt crucible, to c. 1100 for two or three hours.

For descriptions of the bromoplatinates, chloroplatinates, iodoplatinates, &c., v. Platini-BROMIDES, PLATINI-CHLORIDES, PLATINI-IODIDES,

THIOPLATINATES. PtS2 combines with some more basic sulphides to form salts which may be called thioplatinates; some salts are also known containing Pt and Sn combined with alkali metal and S, these may be called thiostannoplatinates. The alkali thioplatinates are obtained by fusing together spongy Pt, S, and alkali carbonate, and washing with water, wherein the thioplatinates are insoluble. Other thioplatinates-of Cd, Cu, Fe, Pb, Mn, Hg, Ag, Tl, Sr, and Zn—are obtained by double decomposition from the alkali salts. The thioplatinates belong to the forms M¹₂Pt₄S₆ and M¹₁Pt₄S₆; the corresponding thioplatinic acids H₂Pt₈S₆ and H₄Pt₈S₆ are obtained by decomposing K₂Pt₈S₆ and Na, Pt, Se respectively by dilute HClAq.

Potassium thioplatinate $K_2Pt_4S_6 = K_2S.3PtS.PtS_2$ (Potassium platin-thioplatinate). Blue-grey, metal-like crystals; S.G. 644 at 15°. Glows like tinder when heated in air, forming K2SO4 and Pt; reduced in H with

evolution of H_2S and formation of Pt. Obtained by fusing an intimate mixture of 1-2 parts Pt black with 12 parts of a mixture of equal parts of S and K_2CO_2 in a porcelain crucible, keeping the molten mass over the blowpipe for a few minutes and extracting with water when cold.

Sodium thioplatinate

Na, Pt, Se = 2Na, \$\bar{S}\$. 2PtS. PtS. (Disodium platinthioplatinate). Copper-red, rhombic needles; decomposed in air. Obtained similarly to the K compound mentioned above.

For details regarding thioplatinates, and also thiostannoplatinates, v. Schneider, P. 136, 105; 138, 604; 139, 661; 141, 519; 148, 633; 149, 881 (cf. E. von Meyer, J. pr. [2] 15, 1).

M. M. P. M. PLATINITES, derivatives of; v. Platino-bromides, Platino-chlorides, Platino-nitrites, &c.

PLATINI- and PLATINO- COMPOUNDS. Such compounds as platini-bromides, called also bromoplatinates, and platino-nitrites will be described here.

PLATINI- COMPOUNDS. These compounds are derived from platinic compounds, especially from PtCl₄, PtBr₄, and PtI₄; they generally react as salts of acids containing tetravalent atoms of Pt in their acidic radicles.

Platini-bromhydric acid H₂PtBr₆·9H₂O (Bromoplatinic acid). Formed by heating spongy Pt with Br, and HBrAq (b.p. 126°), in a sealed tube, to 180° (Meyer a. Züblin, B. 13, 404; Halberstadt, B. 17, 2962). Also by dissolving Pt black in HNO₃Aq mixed with HBrAq, heating with repeated additions of HBrAq, evaporating over CaO, and washing the crystals, on an asbestos filter, with CS₂ (H., l.c.). Large, clear, carmine-red, monoclinic crystals; v. sol. water, alcohol, ether, acetic acid, and CHCl₂. Melts at 100° with partial decomposition (Topsöe, Ar. Sc. 35, 58; 45, 223).

Platini-bromides M¹₂PtBr₆ (Bromoplatin-

Platini-bromides M¹-PtBr_e (Bromoplatinates). Thomsen (Th. 8, 430) gives [Pt, Br¹, 2RBrAq] = 57,160 and [Pt,O², 6RBrAq] = 80,360, where R = H, Na, K, Am, ½Ca, ½Ba, ½Sr, ½Mg. These salts are generally formed by evaporating H₂PtBr_eAq, or a solution of Pt in Br and HBrAq, with metallic bromides. They are red crystalline solids, isomorphous with the corresponding Cl salts. As solutions of these salts give a pp. of Ag₂PtBr_e, and not AgBr, with a limited quantity of cold AgNO₃Aq, and on electrolysis of their solutions the Pt goes with the Br to the positive electrode, they are better regarded as ordinary salts than as double salts 2MBr.PtBr_e.

Ammonium Platini-Bromide Am_PtBr_e. Red octahedra; S.G. 4.2; sl. sol. cold water. S. at 20°=.59 (Halberstadt, B. 17, 2962). Formed

by evaporating H₂PtBr₆Aq + NH₄Br.

POTASSIUM PLATINI-BROMIDE K₂PtBr₆. Red octahedra; S.G. 4·54 (Topsöe, Ar. Sc. 35, 58; 45, 223). S. 2·07 at 20°, 10 at 100° (H., l.c.). Obtained by ppg. H₂PtBr₆Aq by KBr, or by evaporating H₂PtCl₆Aq with KBr. Thomsen (Th. 8, 480) gives

[Pt,Br 4 ,2KBr] = 59,260; [K 2 PtBr 4 ,Aq] = -12,260. Double compounds with K $_2$ PtCl $_2$ are

described by Pitkin (C. N. 41, 118).

SODRUM PLATINI-BROMIDE Na₂PtBr_c.6H₂O. Dark-red, triclinic prisms; S.G. 3·328; very sol.

water and alcohol. Obtained by evaporating PtCl,Aq with HBr till the Cl is turned out, adding NaBrAq, evaporating to dryness, dissolving in a little water, and crystallising (Thomsen, J. pr. [2] 15, 294). Thomsen (Th. 3, 430) gives [Pt,Br*,2NaBr,6H*O] = 65,330; [Na²PtBr*,6H²O] = 18,540; [Na²PtBr*,6H²O,Aq] = -8,550. Platini-bromides of Ba with 10aq, Ca with 12aq, Co with 12aq, Cu with 8aq, Pb, Mg with 12aq, Mn with 12aq, Ni with 6aq, Sr with 10aq, and Zn with 12aq, are described by Topsöe (Ar. Sc. 35, 58; 45, 223); von Bonsdorff (P. 19, 343) describes a Mn salt with 6aq.

Platini-bromonitrites (Nitro-bromoplatinates. Platini-nitrobromides). Salts derived from the platini-bromides by replacing Br by NO₂. Bygently heating K₂Pt(NO₂)₄(v. Platino-nitrites, p. 284) with Br, polassium platini-dibromonitrite, K₂PtBr₂(NO₂)₄, is obtained as a yellow powder, sl. sol. cold, more sol. hot, water (Vèzes, C. R. 112, 616). When an aqueous solution of this salt is concentrated at a gentle heat, it yields crystals of potassium platini-tribromonitrite, K₂PtBr₃(NO₂)₃ (V., C. R. 115, 44). Potassium platini-tetrabromonitrite, K₂PtBr₄(NO₂)₂, is obtained by the regulated action of Br on K₂Pt(NO₂)₄; it forms red prisms, sol. water with partial decomposition (V., C. R. 115, 44).

Platini-chlorhydric acid H₂PtCl_a.6H₂O (Chloroplatinic acid). A solution of this compound is the starting-point for the preparation of very many Pt compounds. Formed by dissolving Pt in aqua regia, repeatedly evaporating with conc. HClAq till every trace of HNOs is expelled, and allowing to crystallise (Weber, P. 131, 441; Jörgensen, J. pr. [2] 16, 345; Topsöe, Ar. Sc. 35, 58). Red-brown, very deliquescent crystals; S.G. 2.431. Easily sol. alcohol, forming H_PtCl_(OEt)₂ (Schützenberger, A. Ch. [4] 21, 362). Heated to 230°, gives PtCl₂ (v. Platinum dichloride, Preparation, p. 289). Heated in Cl to above 350° gives PtCl₂ (v. Platinum tetra-CHLORIDE, Formation, p. 289). By adding much H2SO4 to H2PtCl6.6H2O in a little water, Pigeon (C. R. 112, 1218) obtained the hydrate with 4H₂O; and by heating in vacuo at 100°, over fused KOH, he obtained HPtCl₅.2H₂O. Thomsen (Th. 3, 430) gives $[2HClAq,Pt,Cl^2] = 84,620$; $[6HClAq,Pt,O^2] = 64,060$. Pigeon (C. R. 110, 77) gives $[H^2PtCl^6.6H^2O,Aq] = 4,340$. H_2PtCl_6Aq reacts as a dibasic acid; the platini-chlorides are numerous and important; the Am and K salts are only sl. sol. water and insol. alcohol, and are much used as forms for estimating K and ammonia. Very many organic bases replace H and form salts analogous with the metallic platinichlorides.

Platini-chlorides M¹₂PtCl_e (Chloroplatinates). Thomsen (Th. 3, 430) gives [Pt,Cl¹,2RClAq] = 84,620; and [Pt,O²,6RClAq] = 64,660; where R=H, Am, K, Na, ½Ba, ½Ca, ½Sr, ½Mg. Pigeon (C. R. 110, 77) gives [PtCl¹,2HClAq] = 24,800. These salts are generally obtained by evaporating H₂PtCl_eAq with metallic chlorides. Most of them are yellow-red, crystalline solids; usually e. sol. water and alcohol. They are better regarded as ordinary salts than as double chlorides (cf. Platin-Bromder, supra). The platinichlorides were investigated by von Bonsdorff (P. 17, 250); later by Cleve (Bl. [2] 21, 118, 197, 247, 345); also by Topsöe (Ar. Sc. 35, 58); and

by Nilson (B. 9, 1056, 1142). Crystallographically considered, these salts fall into four groups:—(1) M₂PtCl₈, where M = Am, Cs, K, Rb, Tl (and Pt may be replaced by Pd, Ir, or Sn); regular, isomorphous with Am₂(or. K₂)PtBr₆, Am₂(or K₂)PtI₈, fluosilicates, fluostannates, and fluoziroonates of similar composition. (2) MPtCl₈-6aq, where M = Cd, Co, Cu, Fe, Mg, Mn, Ni, Zn (and Pt may be replaced by Pd or Sn); hexagonal. (3) MPtCl₈-12aq, where M = Mg or Mn; hexagonal; isomorphous with corresponding platini-bromides. (4) M(PtCl₈)₂-24aq, where M = Ce₂ or La₂; hexagonal.

AMMONIUM PLATINI-CHLORIDE Am₂PtCl₆ (Ammonium chloroplatinate. Platinsalammoniac). Ppd. on adding NH₄Ol to H₂PtCl₆Aq, PtCl₄Aq, or a solution of Pt in conc. HClAq with a little HNO₂. Yellow, regular, octahedral crystals. S.G. 8·065 (Topsöe, l.c.). S. ·665 at 20°, 1·25 at 100° (Michaelis, G.-O. 1, 1187). Scarcely sol. alcohol or ether. Decomposed at red heat, leaving Pt black. For reactions with NH₃ v. Platinum-ammonium compounds, p. 292.

K.PtCl. POTASSIUM PLATINI - CHLORIDE (Potassium chloroplatinate). Ppd. by adding KCl, or other K salt, to conc. H2PtCl6Aq, or to a solution containing Pt which has been evaporated with excess of HÖlAq. Reddish-yellow regular octahedra. S.G. 3.586 (Bödeker). S. 74 at 0°, 1.12 at 20°, 2.17 at 50°, 3.79 at 80°, 5.13 at 100° (Michaelis, I.c.). Almost insol. alcohol, or alcohol and ether. Slightly sol. dilute acids; sol. KOHAq. Reduced to Pt and KCl by heating to redness; more quickly by heating with reducing agents such as oxalic acid or sulphurous acid, or by heating in a stream of hydrogen. Vèzes (C. R. 110, 757) describes a nitroso-derivative, K,Pt(NO)Cl₃. Pigeon (C. R. 110, 77; 112, 791) gives [Pt,Ol⁴,2KClAq] = 83,330; [PtCl⁴,2KClAq] = 25,330; [PtCl⁴,2KCl] = 29,700. Thomsen (Th. 3, 430) gives $[Pt,Cl^4,2KCl] = 89,500$.

Potassium platini-bromochloride K₂PtBr₂Cl₄; by ppg. H₂PtCl₂Aq by KBr (Pitkin,

C. N. 41, 118).

The following platini-chlorides have also been isolated: AlCl, PtCl, 15aq (Welkow, B. 7, 304; Salm-Horstmar, P. 99, 638); Ba, with 4aq (Topsõe; Bornsdorff, Precht, Fr. 1879. 509); Be, with 8aq (Welkow, B. 6, 1288; Thomsen, B. 3, 827); Cd, with 6aq (Topsõe); Cs (Crookes, C. N. 9, 37; Bunsen, P. 119, 371); Ca, with 9aq (Topsõe; Precht); Ce, various salts (Cleve, Marignac, Holzmann, J. pr. 84, 80); CrCl, PtCl, 10aq (Nilson, B. 9, 1056, 1142); Co, with 6aq (Topsõe); Cu, with 6aq (T.); Di, various salts (Frerichs a. Smith, A. 191, 331); 2InCl, 5PtCl, 36aq (Crookes, J. 1864, 256); Fe, various salts (T.; Bonsdorff; Nilson); La, various salts (Cleve, Marignac, Jolin, B. 11, 910); Pb, with 3 or 4aq (Birnbaum, J. 1867, 319; Topsõe); Li, with 6aq (Scheibler, J. pr. 67, 485; Jörgensen a. Topsõe, Gm.-K. 3, 1174); Mg, with 6aq (T.); Mn, with 6aq (T.); Ni, with 6aq (T.); Rb (Crookes; Bunsen); Ag (Birnbaum); Na, with 6aq (Marignac, Topsõe, Precht; for thermal data v. Thomsen, and Pigeon, Lc.); Sr, with 8aq (Bonsdorff); Tl (Crookes); Th, with 12aq (Cleve); Sn, with 12aq (Nilson). Compounds of PtCl, with chlorides of Er (Nilson), Hg (Birnbaum), VO (Brauner, M. 3, 58), and ZrO (Nilson) probably exist. No com-

pounds are formed with PtCl₄ and chloride of Sb, As, or Bi.

Platini-chlorenitrites (Platini-nitrochlorides. Nitrochloroplatinates). Salts derived from the platini-chlorides by replacing Cl by NO₂. A few of these salts are described by Vèzes (C. R. 115, 44); the principal are dichloronitrite K₂PtCl₂(NO₂), trichloronitrite K₂PtCl₃(NO₂), and pentachloronitrite K₂PtCl₃(NO₂). L₂O (v. also Blomstrand, J. pr. [2] 3, 214).

Platini-iodhydric acid H₂PtI₆.9H₂O (Iodoplatinic acid). Brown deliquescent crystals; probably monoclinic; by dissolving PtI₄ in HIAq, and evaporating. Easily decomposed, even in solution, rapidly at 100°, with separation of PtI₄ (Topsöe, Ar. Sc. 38, 297; Clementi, J. 1855. 420; Lassaigne, A. Ch. [2] 51, 113).

Platini-iodides M¹₂PtI₆ (Iodoplatinates). Reddish-brown, metal-like salts; generally formed by evaporating H₂PtCl₈Aq with excess of metallic iodides, or by adding iodides to PtI₄Aq. The following have been described (v. Topsöe, Clementi, Lassaigne, l.c.): Am; Ca, with 12aq; Co; Fe; Mg, with 9aq; Mn; Ni, with 6aq; K; Na, with 6aq; Zn, with 9aq.

Platini-iodonitrites (Platini-nitro-iodides. Nitro-iodoplatinates). M¹₂PtI_{3-x}(NO₂)_x. A few of these salts are described by Vèxes (C. R. 113, 696). A nitroso-platini-iodide, K₂Pt(NO)I₃, is also described.

Platini-molybdates. By boiling Pt(OH,) with an acidified solution of Na molybdate, Gibbs (Am. S. [3] 14, 61) obtained a complex compound which may provisionally be classed as a platini-molybdate, 4Na₂O.10MoO₃.PtO_x.29H₂O.

Platini-nitrobromides; v. Platini-bromo-NITRITES, p. 282.

Platini-nitrochlorides; v. Platini-chloro-NITRITES, supra.

Platini-nitro-iodides; v. Platini-1000-

NITRITES, supra.

Platini-tungstates. Complex compounds of PtO₂, WO₃, and strong bases (v. Gibbs, Am. S. [3] 14, 61; Rosenheim, B. 24, 2397).

PLATINO- COMPOUNDS. These compounds are derived from platinous compounds, especially from PtCl₂, PtBr₂, and PtI₂; they generally react as salts of acids of the form H₂PtX₄, where X is a monovalent negative radicle, generally Cl, Br, or I.

Platino-bromhydric acid H₂PtBr₄ (Bromoplatinous acid). This compound is probably contained in a solution of PtBr₂ in HBrAq.

Platino-bromides M¹₂PtBr₄ (Bromoplatinites). Only one of these salts, K₂PtBr₄, has been isolated. Potassium platino-bromide is obtained by adding a very little water to a mixture of the corresponding Cl salt and NaBr, in the ratio K₂PtCl₄: 4NaBr, boiling, sucking up the clear liquid from ppd. NaCl, and allowing to cool, when the salt crystallises in dark-brown octahedra, or brown-red needles, which are very sol. water (Thomsen, J. pr. [2] 15, 294). Thomsen (Th. 3, 430) gives [2KBr,Pt,Br²] = 32,310.

Platino-bromonitrites v. Platino-nitrites, p. 284.

Platino-chlorhydric acid H.PtCl₄ (Chloroplatinous acid). This acid has not been isolated; but it almost certainly exists in a solution of PtCl, in HClAq, in the liquid obtained by decomposing BaPtOl, by the equivalent quantity of H₂SO₄Aq (Nilson, J. pr. [2] 15, 260), and in the solution formed when conc. H2PtCl2Aq is added to hot conc. K₂PtCl₄Aq (Thomsen, J. pr. [2] 15, 294). When these liquids are evaporated in vacuo, a residue is obtained agreeing in composition with the formula HPtCl₂.2H₂O (= HCl.PtCl₂.2H₂O = H₂PtCl₈(OH).H₂O); at 100° this loses H₂O and HCl and leaves PtCl₂ (Nilson, *l.c.*).

Platino-chlorides M12PtCl, (Chloroplatinites). These salts are obtained by evaporating metallic chlorides with PtCl2Aq, or in many cases by reducing platini-chlorides. Most of the platinochlorides are very soluble in water, and crystallise only from very conc. solutions, generally forming dark-red crystals. Many of them are described by Nilson (J. pr. [2] 15, 260). Thomsen (Th. 3, 430) gives $[Pt,Cl^2,2MClAq] = 41,830$; and [Pt,O,4MClAq] = 31,550; where M = H, K, Na, Am, $\frac{1}{2}$ Ba, $\frac{1}{2}$ Ca, $\frac{1}{2}$ Sr, or $\frac{1}{2}$ Mg.

AMMONIUM PLATINO-CHLORIDE. Am₂PtCl₄ Obtained, in (Ammonium chloroplatinite). four-sided prisms, by adding AmCl to PtCl2 in HClAq, and evaporating; also by reducing hot Am2PtClaAq by SO2, or hot H2PtClaAq by SO2 and then adding AmCl (Peyrone, A. 55, 205; Thomsen, B. 2, 668; Grimm, A. 99, 95). [Pt,Cl²,2AmCl] = 43,550 (Th. 3, 430).

Potassium platino-chloride K₂PtCl₄ (Potassium chloroplatinite). Large, ruby-red, foursided prisms; S.G. 3.2909 at 21°; easily sol. water, insol. alcohol (Nilson). An aqueous solution is not ppd. by soda or potash when cold; on boiling, all the Pt is thrown down as Pt(OH)2 (Thomsen, J. pr. [2] 15, 295). [Pt,Cl²,2KCl] = 45,170 (Th. 3, 430). Obtained by adding KCl to PtCl2 in HClAq, and evaporating (Magnus, P. 14, 241); also by reducing K2PtCl6Aq by H.S (Böttger, J. pr. 91, 251), or better by Cu₂Cl₂ (Thomsen, J. pr. [2] 15, 294). K₂PtCl₃ is made into a paste with water, warmed, and moist Cu₂Cl₂ is added, little by little, until a small excess is present; the liquid, which is nearly black, is filtered, the red crystals which separate on cooling are washed with alcohol and re-crystallised.

The following platino-chlorides have been isolated and described: Ba, with 3aq; Be, with 5aq; Ca, with 8aq; Cs (Böttger, J. pr. 91, 251); Co, with 6aq; Cu, with 6aq (Thomsen, l.c.; Millon a. Commaille, C. R. 57, 822); Feous, with 6aq; Pb; Li, with 6aq; Mg, with 6aq; Mn, with 6aq; Ni, with 6aq; Rb; Ag; Na, with 4aq; Sr, with 6aq; Tl; Zn, with 6aq (Hünefeld, S. 60, 197). Compounds of PtCl₂ with the chlorides of the following metals are also described by Nilson (l.c.); Al, Ce, Cr, Di, Er, La, Th, and Y.

Platino-chloronitrites; v. Platino-nitrites, infra.

Platino-chlorophosphates (Phospho-platino-chlorides). Several salts, and a few acids, containing Pt, Cl, and P have been isolated by Schützenberger (Bl. [2] 17, 482; 18, 101, 148). The classification of these compounds is very incomplete. The following scheme is accepted as provisional (cf. Seubert, Ladenburg's Handwörterbuch der Chemie, 9, 314):

Platino-chlorophosphoric acid Cl.Pt:P(OH);; by dissolving Cl.Pt.PCl, in water and crystal-

lising; (Cl2Pt:POs)2Pbs.8aq, and various esters, isolated.

Platino-chlorodiphosphoric acid

P(OH)3 Cl2Pt: ; by the action of moist air on P(OH), Cl₂Pt:PCl₃.PCl₃; ethyl ester also isolated.

Platino-chloropyrophosphoric acid P(OH)2

ClPt: Ò ; by warming a solution of Ý(OH)

Cl₂Pt:PCl₃.PCl₃ Platino-chloro-anhydropyrophosphoric acid $P(OH)_2$

ClPt: O ; by heating the foregoing acid to P(OH) 150°

Ethyl diplatino-chloro-phosphate

 $\frac{\text{Cl}_2\text{Pt}}{\text{Cl}_2\text{Pt}} > \text{P(OEt)}_s$; by dissolving $\frac{\text{Cl}_2\text{Pt}}{\text{Cl}_2\text{Pt}} > \text{PCl}_s$ in alcohol; corresponding acid not isolated (Cochin, C. R. 86, 1402).

Platino-cyanides and derivatives; v. vol. ii.

p. 344.

Platino-iodhydric acid H2PtI4 (Iodoplatinous acid). This acid probably exists in a solution of PtI, in HIAq.

Platino-iodonitrites: v. Platino-nitrites. Platino-nitrites, and derivatives. Platinous nitrite [Pt(NO2)2] has not been isolated; but several compounds are known in which the group Pt(NO_o), forms part of the acidic radicle. These Pt(NO₂)₂ forms part of the acidic radicle. platino-nitrites belong to the form M2Pt(NO2)4; they are not to be regarded as double salts, but as derivatives of the acid H2Pt(NO2)4; the ordinary reagents for Pt do not show the presence of this metal in solutions of these salts, nor do these solutions give the reactions of nitrites; their solutions doubtless contain the ions M and Pt(NO₂)₄. Most of the platino-nitrites are obtained from the potassium salt K2Pt(NO2)4, which is formed by the reaction of equivalent weights

of K₂PtCl₄ and KNO₂.

PLATINO NITROUS ACID H₂Pt(NO₂), has not been obtained pure, but it doubtless exists in the solution obtained by decomposing BaPt(NO2)4 by H2SO4Aq; on evaporation, this solution gives red crystals (Lang, J. pr. 83, 415), but the process causes partial decomposition to triplatino-octonitrous acid H4Pt3O(NO2)8.2H4O

(Nilson, B. 10, 934). POTASSIUM PLATINO-NITRITE K₂Pt(NO₂) Small, lustrous, (Potassium nitroplatinite). colourless, monoclinic prisms; by mixing solutions of equivalent weights of K2PtCl, and KNO2, and evaporating (Topsöe, J. 1879. 307). S. 3.8 at 15°; more sol. hot water. Combines with Br and Cl to form potassium platini-dibromo-[dichloro-] nitrites, K₂PtBr₂[Cl₂](NO₂). Most of the other platino-nitrites are obtained from this salt, by adding AgNO.Aq, separating Ag.Pt(NO2)4, and decomposing this by metallic chlorides; or making BaPt(NO2), by the action of BaCl,Aq on the silver salt, and decomposing this by sulphates (Nilson, B. 9, 1722; 10, 930; 11, 879; v. also Blomstrand, J. pr. [2] 3, 186).

The other platino nitrites which have been

isolated are: Al, Am, Cd, Ce, Cr, Co, Di, Sr, Fe, La, Pb, Li, Mg, Mn, Hg, Ni, Ag, Na, Tl, Y, and Zn.

PLATINO-BROMONITRITES M¹₂Pt(NO₂)_{4-x}Br₁. A few of these salts have been obtained by Vèzes (C. R. 113, 696; 115, 44). Potassium platino-bromonitrite K₂PtBr(NO₂)₃ and dibromo-nitrite K₂PtBr₂(NO₂)₂ are described.

PLATINO-CHLORONITRITES M¹₂Pt(NO₂)_{4-x}Cl_x. Vèzes (l.c.) has described the potassium monochloro-salt K₂PtCl(NO₂)₃, and the dichloro-salt

 $\mathbf{K_2PtCl_2(NO_2)_2}$.

PLATINO-IODO-NITRITES M₂Pt(NO₂)₄₋₁I_x (Nilson, B. 10, 930; 11, 879; Groth, Z. K. 4, 469; Vèzes, C. R. 115, 44). Potassium platino-di-iodo-nitrite K₂PtI₂(NO₂)₂.2H₂O is obtained, in small, black crystals, by the action of an alcoholic solution of I on K₂Pt(NO₂)₄. The other salts isolated are those of Al, Am, Ba, Be, Cd, Cs, Ca, Ce, Co, Cu, Di, Fe, La, Pb, Li, Mg, Mn, Ni, Rb, Ag, Na, Sr, Tl, Y, and Zn.

Platino-nitrobromides; v. PLATINO-BROMO-

NITRITES, supra.

Platino-nitrochlorides; v. Platino-chloro-NITRITES, supra.

Platino-nitro-iodides; v. Platino-iodo-

NITRITES, supra.

Platino-oxalic acid and platino-oxalates $H_2Pt(C_2O_4)_2$ and $M^1_2Pt(C_2O_4)_2$ (Söderbaum, Bl. [2] 45, 188). The sodium salt $Na_2Pt(C_2O_4)_2$, $4H_2O$ is obtained, in copper-coloured crystals, by heating $Na_2O.3PtO_2.6H_2O$, with $1\frac{1}{2}$ crystallised oxalic acid, cooling the blue-coloured liquid, treating the brown needles which separate with hot water, and crystallising; sometimes the salt separates with 5H₂O as golden-coloured crystals. The other salts are obtained by double decomposition from the Na salt; they seem to exist in two forms corresponding with the two sodium salts. Platino-oxalic acid H₂Pt(C₂O₄)₂·2H₂O is obtained as red, lustrous, metal-like orystals by decomposing the silver salt with the equivalent weight of HClAq, filtering, and concentrating the blue solution in vacuo. Salts of the following metals are described by Söderbaum (l.c.): Am, Ba, Ca, Mg, Mn, Ni, K, Ag, Na, Sr, and Zn.

Platino-phosphochlorides v. PLATINO-CHLORO-

РНОЅРНАТЕS, р. 284.

Platino-seleno-stanuates. Schneider (J. pr. [2] 44,507) has described two salts, K₂Pt₃SnSe₆ and Na₂Pt₃SnSe₆, which may be called platino-seleno-stanuates. They are formed by heating together Pt black, SnSe₄, K₂CO₃ or Na₂CO₃, and Se.

Platino-stannates. This name may be given to some compounds derived from the acids $H_2Pt_2Sn_2O_4$ and $H_2Pt_2Sn_2O_4$, described by Schneider (P. 136, 105) and Schützenberger (C. R. 98, 985); cf. also Lévy a. Bourgeois (C. R. 94, 1365).

Platino-sulphocyanhydric acid, and salts v.

vol. ii. p. 357.

Platino-sulphonates, and derivatives (Liebig, A. 23, 23; Litton a. Schnedermann, A. 42, 316; Birnbaum, A. 189, 164; 152, 137; 159, 116; Döbereiner, J. pr. 15, 315; Lang, J. pr. 83, 415). (Platino-sulphites. Sulpho-platinites.) When K₂PtCl₄ is heated with KHSO₂Aq, or SO₂ is passed into K₂SO₂Aq holding Pt(OH)₄ in suspension, the liquid is neutralised by K₂CO₂ and

evaporated, yellow, microscopic crystals are obtained of K₂Pt(KSO₃)₄.2H₂O. From this potassium platino-sulphonate other similar salts are obtained. These salts are better regarded as derived from platino-sulphonic acid H₂Pt(HSO₃)₄ than as double compounds of PtSO₃ and K₂SO₄[K₂Pt(KSO₄)₄ = PtSO₃.3K₂SO₃]; they are analogous to the platino-chlorides M₂PtCl₄ and the platino-nitrites M₂Pt(NO₂)₄. The ammonium, sodium, and silver salts have been isolated. Sodium-platino-oxy-sulphonic acid Na₂Pt(HSO₃)₂O and ammonium-platino-oxy-sulphonic acid Am₂Pt(HSO₃)₂O have also been isolated.

PLATINO - CHLOROSULPHONATES (Platino-chlorosulphites. Sulpho - chloroplatinites).

M¹-yPt.Cl₃.M¹SO₃. The potassium salt K_2Pt.Cl₃.KSO₃ is obtained by warming K_2PtCl₄ with SO₂Aq, evaporating, and adding KCl. Ammonium platino - chlorosulphonic acid Am₂Pt.Cl₃.HSO₃ crystallises from a solution of Am₂PtCl₅, or Am₂PtCl₄, in warm conc. SO₂Aq; the potassium and sodium salts of this acid are known. A complex acid, AmH.Pt.Cl.SO₃.HSO₃, and the salts AmK.Pt.Cl.SO₃.KSO₃ and Am₂.Pt.Cl.SO₃.SO₄Am have been isolated.

Platino-thiosulphates. A few salts have been obtained which may be regarded as compounds of hypothetical platinous thiosulphate and sodium thiosulphate PtS₂O₃.xNa₂S₂O₃.yH₂O, where x is 3, 4, 6, and 7. They may also be looked on as derivatives of the hypothetical acid H₂Pt(S₂O₃H)_x, where H is replaced by Na (Schottländer, A. 140, 200; Jochum, C. C. 185.

642). M. M. Pt. At. w. 194.3. Mol. w. unknown. Melts at c. 1775° (Violle, C. R. 89, 702); older determinations generally gave m.p. c. 2000° (for references v. Carnelley's Melting and Boiling Point Tables, p. 10), although Becquerel (C. R. 57, 855) gave 1460°-1480°. S.G. 21·48 to 21·5 at 17·6°, after melting in H (Deville a. Debray, C. R. 81, 839; for other values v. Clarke's Specific Gravity Tables, 2nd edit. p. 15). S.H. ·03248 (Regnault); ·0314 (Dulong a. Petit); ·0317 + ·000012t at t° (Violle, P. M. [5] 4, 318). C.E. (linear at 40°) ·00000899 (Fizeau, C. R. 68, 1125, for Pt that had been melted); from 0° to 100°, .00000881 (Calvert a. Johnson, B. A. 1858. for hammered Pt). T.C. (Ag = 1000) 379 (C. a. J., l.c.); c. 84 (Wiedemann a. Franz, J. 1855. 91). Heat of fusion c. 5276 (for 194 g.) (Violle, l.c.). E.C. (Ag=100) 8.042 at 12°-18° (Becquerel, A. Ch. [8] 17,242); 10.53 at 20.7° (Matthiessen, P. 103, 428). For electrical resistance of Pt wire v. Arndtsen (P.104,1). Crystallises in the regular system (v. infra, Properties). For chief lines in emission-spectrum v. B. A. 1884. 436.

Occurrence.—Alloyed with Ir, Pd, Rh, Au, Cu, and Fe in the sands of many rivers, often associated with titaniferous iron and chromeiron, and in the older rocks in different parts of the world. The quantity of Pt in 'platinum ores,' varies from c. 50 to c. 90 p.c. A Canadian Ni ore was found by Clarke a. Catlett to contain from '008 to '02 p.c. Pt (Am. S. 37, 572). In 1741 Watson received some particles of a metallike substance from the gold-bearing sand of the river Pinto in New Granada; the shining silvery particles were called in the district where they occurred platina del Pinto (platina = diminutive

of Spanish plata = silver). Watson extracted a new metal from the substance sent him, and called it platinum (T. 1750. 585). For a short account of the history of the Pt metals, v. NOBLE METALS, iv. 628.

Formation.—1. By decomposition of PtCl₄ or (NH₄)₂PtCl₅ by heat.—2. By heating an alloy of Pt and Pb in a current of air, whereby the Pb is oxidised and the Pt remains.—3. By reducing many Pt salts by Zn, Fe, NaOHAq and FeSO₄Aq, Na₂CO₂Aq and sugar, alcohol added to an alkaline solution, &c.

an alkaline solution, &c.

Preparation.—1. The Pt ore is obtained as a fine powder by fusing with 2 or 3 times its weight of Zn, powdering the brittle alloy so formed, and removing the Zn by dilute H₂SO₄Aq (Descotils, G. A. 27, 231; Hess, J. pr. 40, 498). The residue is heated to redness, then warmed with HClAq (to remove portions of the baser metals), and then treated with cold aqua regia which dissolves Au. The residue is heated, in retorts, with aqua regia; the solution contains most of the Pt, along with Rh, Pd, and a little Ir; in the residue are found osm-iridium (v. vol. iii. p. 47), Ru, and a little Pt. The acid is distilled off, carrying with it most of the Os as OsO4; the concentrated solution is neutralised by Na₂CO₃, and the Pd is ppd. as PdCy₂ by HgCy₂; NH₄Cl is added to the filtrate, and (NH₄)₂PtCl₆ is thus ppd. mixed with some (NH₄)₂IrCl₆. The Ir salt may be removed by digesting with slightly warm KCNAq until the undissolved portion is light yellow; the salt IrCl, 3NH, Cl is thus formed and dissolved, while (NH₄)₂PtCl₅ remains (Wöhler a. Mucklé, A. 104, 368). Or the NH₄.Pt and NH₄.Ir chlorides may be dissolved in hot water, and the Ir salt reduced to IrCl, 3NH, Cl, which dissolves, by SO, (cf. IRIDIUM, vol. iii. p. 46). The residual ammonium platinichloride is then heated until all NH, and Cl are volatilised.—2. Finely-divided Pt ore is dissolved in aqua regia diluted with 2 pts. water, under increased pressure (which aids the solution); the solution is evaporated, and the residue is heated to 125°, whereby the chlorides of Pd and Ir are reduced to the lower chlorides; treatment with HClAq dissolves PtCl₄, which is ppd. as $(NH_4)_2$ PtCl₂ by addition of NH_4 Cl (Heræus, v. Hofmann's Chem. Industrie auf der Wiener Weltausstellung, 2, 999; v. also Dullo, J. pr. 78, 369).—3. Commercial Pt is melted with 6 times its weight of Pb, the granulated alloy is treated with dilute HNO, Aq (1:8 by volume), which dissolves most of the Pb, Cu, Pd, and Rh; the black residue is dissolved in diluted aqua regia-Ir remains undissolved—the solution is evaporated with H2SO4Aq, whereby Pb is removed as PbSO, the filtrate and washings are mixed with excess of NH₂Cl and some NaCl, heated to 80°, and allowed to stand for some days; the pp. is washed repeatedly with saturated NH_ClAq, and then with very dilute HClAq; it is then dried, heated to dull redness in a Pt vessel with KHSO, and a little NH, HSO, and washed with boiling water-Rh goes into solution as Rh-K disulphate, and finely-divided Pt remains (Matthey, Pr. 28, 463). For other methods v. Deville a. Debray, A. Ch. [3] 56, 385; 61, 5.— 4. Pure Pt is obtained by evaporating a solution of the ore in aqua regia with HClAq till HNO, is nearly removed, adding NaOHAq till strongly

alkaline, boiling for a long time (the alkalinity gradually disappears, owing to formation of NaClOAq), adding alcohol, and then making the turbid liquid acid by HClAq, filtering from olive-green IrCl₃, and ppg. by conc. NH₄ClAq (W. von Schneider, A. Suppl. 5, 261). The Pt obtained by strongly heating the ppd. (NH₄)₂PtCl₃ should be dissolved in diluted aqua regia, and re-ppd. by addition of NH₄Cl; on heating this pp. pure Pt remains (Seubert, A. 207, 8). For another method of obtaining pure Pt, v. Mylius a. Förster. B. 25, 665.

ter, B. 25, 665.

The finely-divided Pt obtained by the above methods may be fused into a lump by the use of the O-H flame. The metal is placed in a cavity made in a block of lime (made by strongly heating marble), which is covered by another block, through the top of which a hole is pierced to admit the flame (v. Deville a. Debray, A. Ch. [3] 56, 385; cf. Platinum in Dictionary of Applied Chemistry).

Platinum black, which is extremely finelydivided Pt, is obtained by reducing solutions of some Pt salts by certain organic reducing compounds, or by Zn or Mg, &c. PtCl, Aq may be reduced by Mg (Böttger, J. pr. [2] 2, 137), zincdust or Fe in powder (Brunner, A. 109, 253), FeSO, Aq and NaOHAq (Hempel, A. 107, 97), sugar and Na₂CO₃Aq, or by alcohol. Pt black is also formed by fusing Pt with twice its weight of Zn, powdering the alloy, dissolving out Zn by H₂SO₄Aq, and washing with very dilute HNO₃Aq (Döbereiner, A. 17, 67). Loew (B. 23, 289) recommends to dissolve 50 g. PtCl, in 50-60 c.c. water, to add c. 70 c.c. formic aldehyde solution of 40 to 45 p.c., to cool, and then to add gradually 50 g. NaOH in 50 c.c. water, to set aside for 12 hours, and then to filter and wash until the washings pass through black, when the washing should be stopped for some hours, and continued when the washings again pass through colourless (O is absorbed and temperature rises); the residue is washed until quite free from NaCl, and dried over H₂SO₄. Liebig (P. 17, 101) dissolved PtCl₂ in warm conc. KOHAq, added alcohol to the hot liquid till CO2 came off freely, decanted, washed the ppd. Pt, successively, with alcohol, HClAq, KOHAq, and finally, several times, with boiling water.

Crystals of platinum are obtained by heating Pt to redness, in a glass or porcelain tube, in a stream of dry Cl, for twenty-four hours (Hodgkinson a. Lowndes, C. N. 58, 158, 223; Seelheim, B. 12, 2066; Joly, N. 43, 541; cf. Troost a. Hautefeuille, C. R. 84, 94). Joly (N. 43, 541) obtained cubical crystals of Pt, about-1 mm. in length, by sprinkling some finely-powdered topaz on a ribbon of Pt, and heating to bright redness by an electric current for c. two hours. Moissan (C. R. 109, 807) obtained crystalline Pt by heating dry PtF, in a Pt tube, to bright redness. Pt that has been melted shows a crystalline structure when touched with aquaregia (cf. Koettig, J. pr. 71, 190; Noguès, C. R. 47, 832; Kalischer, B. 15, 706; Mallet, Am. S. [2] 20, 340; Phipson, C. N. 5, 144).

Treatment of platinum residues.—Pt residues accumulated in the laboratory generally contain Pt as (NH,),PtCl_e; the liquid portion usually contains alcohol. The solid part is warmed, in a water-bath, with K₂CO₂Aq, KOHAq.

or NaOHAq, and the alcoholic liquid is added little by little, or HCO₂Na is added (Duvillier, C. R. 84, 444), till the salt is reduced to Pt. The black residue is washed, dried, boiled with HClAq, and again washed and dried (Knösel, B. 6, 1159).

Properties.—Compact platinum is white metal with a greyish tinge; it is easily polished, thereby acquiring a very high lustre. Much softer than Ag, rather softer than Cu; thin plates or wire can be cut easily with scissors. Without taste or smell. Very malleable and ductile; a wire 2 mm. diameter breaks with a weight of 124 kilos; tenacity is, therefore, c. same as Fe. Pure Pt can be drawn into very thin wire; by fusing into Ag wire, drawing out, and dissolving away the Ag, a wire of Pt .0009 mm. diameter is said to have been obtained (Gaiffe, C. R. 1877. 625). Most easily fused of Pt metals except Pd; very thin wire melts in the outer blow-pipe flame; larger masses require the O-H flame. Becomes soft and workable much below its m.p. Melts in an ordinary fire, owing to combination with C or Si (Heraeus, D. P. J. 167, 182; Deville, A. Ch. [3]46,182; Boussingault, C. R. 82,591; Schützenberger a. Colson, C. R. 94, 1710). Crystallises in the regular system, chiefly in octahedral and dodecahedral forms (v. Seelheim, B. 12, 2066). Said to volatilise to some considerable extent when kept molten (Deville a. Debray, C. R. 44, 1101). By heating in certain gases, especially in Cl, volatile compounds are formed and again decomposed. Molten Pt absorbs O, and 'spits'

when cooled rapidly.

Pt occludes H. According to Graham (P. M. [4] 36, 63), Pt foil absorbs five to six times its own volume of H at a dull-red heat; Berthelot (C. R. 94, 1383) says that Pt absorbs from 80 to 120 times its volume of H, probably with formation of definite compounds. Berliner (W. 35, 791; cf. Neumann, M. 13, 40) found that ordinary Pt foil absorbed 127 volumes of H; foil that had been quite freed from gases and cleaned absorbed 200 vols. of H and 80 vols. of O or CO. The whole of the occluded gas is removed, with difficulty, by strongly heating in vacuo; under ordinary conditions about 80 vols. of H are retained. O is also occluded by Pt foil, and the condensed O brings about oxidations, e.g. electrolytic gas (H_2+O) explodes when exposed to Pt foil at c. 180° (Berliner, l.c.). Pt is pervious to H at a bright-red heat; Graham (P. M. [4] 32, 401, 503) found that 489.2 c.c. H passed, per minute, through Pt foil 1.1 mm. thick and having a surface of 1 square metre. Gases other than H scarcely diffused through at all.

Pt is not acted on by pure HClAq, HNO₃Aq, or H₂SO₄Aq, nor is it attacked by O; various mixtures of acids dissolve it. Pt is acted on by Cl and by substances which evolve Cl; also, at high temperatures, by molten alkalis, nitrates, KCN, a mixture of alkali carbonates with S, by P and Si, and by most of the metals.

Platinum black is a porous, heavy, very finely-divided black powder, which becomes metal-like, grey, and lustrous when rubbed; S.G. 15-8 to 17-6. Pt black absorbs large quantities of certain gases, especially O; according to Döbereiner (A. 17, 67), from 178 to 258 vols.

O are occluded by 1 vol. Pt black. The O condensed in Pt black is able to bring about many processes of rapid oxidation: H, CO, C,H, C₂H₆O gas, Et₂O vapour, &c., are rapidly oxidised, generally with ignition (v. von Mulder, R. T. C. 2, 44; Schönbein, J. pr. 98, 76; P. 105, 258; von Mulder a. van der Meulen, R. T.C. 1, 167); alcohol is oxidised to acetic acid, formic and oxalic acids to CO2 and H2O, As2O3 to As2O3 (von Mulder, R. T. C. 2, 44). Pt black charged with H acts as a reducer; water is formed when the hydrogenised Pt black is brought into O (Wilm, B. 14, 878; Berthelot, C. R. 94, 1377); aqueous solutions of KClO₃, KNO₃, K₄FeCy₆, also C₆H₆NO₂, &c., are reduced (Gladstone a. Tribe, C. N. 37. 245); ozone is reduced to O (von M. a. van der M., l.c.). One cause of these actions is probably the heat produced during the condensation of the gases in the porous Pt (cf. Berthelot, A Ch. [5] 30, 519); it is not probable that an oxide of Pt is formed and then reduced (v. V. Meyer,

J. pr. [2] 14, 124; cf. Tommasi, B. 11, 811).

The at. w. of Pt has been determined (1) by strongly heating PtCl₂ (Berzelius, P. 8, 179 [1813]); (2) by determining Pt in K₂PtCl₆, and by finding the ratio of Pt:KCl in the same salt (B., P. 13, 469 [1826]; Andrews, Chem. Gazette, 1852. 379); (3) by analyses of K.PtCl_a and (NH₄)₂PtCl_a (Scubert, A. 207, 29 [1800]; Halberstadt, B. 17, 2962 [1884]; cf. Dittmar a. McArthur, E. Tr. 33, 561 [1888], and criticism thereon by Seubert, B. 21, 2179; (4) by determining S.H. of solid Pt (Violle, P. M. [5] 4, 318). As no V.D. of a Pt compound has yet been determined, the at. w. cannot be found by the direct application of the law of Avogadro. The older determinations gave 196 to 197; Seubert proved that the true value is c. 194.5, which is between the values for Ir (192.5) and Au (196.8). No compound of Pt has been gasified from which the valency of the atom of Pt in gaseous molecules can be determined. Pt is closely allied to Ir, and less closely to Os, in its chemical properties. It is distinctly metallic physically, and, on the whole, chemically also. PtO₂ is both basic and acidic; Pt forms numerous acids by combining with H and negative radicles such as Cl, (NO2)4Cl2, (CN)4, (SCN)6, &c. For a fuller account of the chemical relations of Pt v. Noble metals, vol. iii. p. 628, and cf. Ibon GROUP OF ELEMENTS, vol. iii. p. 67.

Pt is used for making crucibles &c. for laboratory use, and vessels for evaporating conc. oil of vitriol, &c. Apparatus is sometimes platinised by placing in hot PtCl₄Aq containing KOH and some organic reducing compound.

Reactions and Combinations.—1. Is not acted on by oxygen; concerning absorption of O by Pt v. Properties.—2. Heated to redness with sulphur, in presence of borax, PtS is formed (Deville a. Debray, C. R. 89, 587).—3. Said to form a compound by heating Pt black with selenion.—4. With chlorine, bromine, and iodine Pt reacts at temperatures above c. 300° (Langer a. Meyer, Pyrochemische Untersuchungen [Braunschweig, 1885] 44, 57). Cl and Br attack Pt in presence of water.—5. Scarcely acted on by fluorine below 100° (Moissan, A. Ch. 1892, 125). 6. Hydrogen is occluded by Pt; a considerable quantity of heat is produced, and perhaps compounds are formed (v. Platinuk hydride, p.

290) .- 7. Heated with arsenic, a compound PtAs, is said to be formed (v. Gm.-K. 3, 1192-1193).—8. Phosphorus combines when heated with finely-divided Pt (Schrötter, J. 1849. 246). 9. A boride is formed by heating Pt black with amorphous boron (Descotils, A. Ch. [3] 67, 88; Deville a. Wöhler, J. 1856. 279).—10. Heated to whiteness with silicon, Pt forms several compounds; also combines with Si when heated with silica and carbon (v. Platinum silicides. p. 291).-11. Softens when heated with carbon (v. Platinum carbide, p. 289).—12. Alloys with many of the more easily fusible metals (v. PLATINUM ALLOYS, infra).-13. Pt is not acted on by pure hydrochloric, nitric, or sulphuric acid (Scheurer-Kestner, C. R. 86, 1082; 91, 59). It dissolves in warm aqua regia, forming PtCl, (v. Dullo, J. pr. 78, 369). Alloying modifies the solubility in acids; alloys of Pt with small quantities of Ir or Rh are scarcely sol. in aqua regia, while an alloy with much Ag dissolves in conc. HNO₈Aq; Pt ppd. by zinc dissolves in conc. HNO Aq (v. Winkler, Fr. 13, 369; van Riemsdyk, B. 16, 387; Wilm, B. 13, 1198).—14. Conc. sulphuric acid containing a little nitrous acid dissolves small quantities of Pt (Scheurer-Kestner, C. R. 86, 1082; 91, 59). Nitric acid, containing bromine or bromhydric acid, and also hydrochloric acid, into which chlorine is passed, dissolve Pt, forming PtBr. and PtCl, respectively (Wagner, W. J. 1876. 149 Seubert, A. 207, 16).-15. Pt is acted on, at red heat, by molten alkalis, baryta, nitrates, potassium cyanide, and by a mixture of alkali car-bonate and sulphur.—16. PtCl₂ or PtBr₂ and PtI₂ are formed when Pt wire is heated in iodine mono- or tri-chloride, or in chlorine mixed with iodine or iodine bromide; with phosphorus pentachloride Pt phosphide is formed; with carbon tetrachloride O, C₂Ol_e, and Cl are produced; hydrogen chloride forms PtCl2; hydrogen fluoride produces a soluble Pt salt; and with mercurous chloride Hg and PtCl2 are obtained (Hodgkinson a. Lowndes, C. N. 58, 223).
Platinum, acids of. The hydroxide Pt(OH).

reacts with strong bases as an acid. Several acids have been isolated containing Pt in combination with more or less complex negative radicles; these are H.PtCl., H.PtCl., H.PtBr_e, H.PtI_e; H.Pt(NO₂)₄, H.Pt₃O(NO₂)₈; H.PtCy₄, H.PtCy₄Cl₂, H.Pt₅(SCN)_e; H.Pt₄Se; Pt(NH₂)₂Cl.SO₂H; PtCl₂P(OH)₂. These acids are described as platino- and platini-chlorhydric acid, &c. (pp. 283 and 282); platino-nitrous acid, &c. (p. 284); platino-cyanhydric acid, &c. (vol. ii. pp. 844-5); thioplatinic acid (p. 281); platos-ammine-chloro-sulphonic acid (p. 293); platino-

chloro-phosphoric acid (p. 284).

Platinum, alloys of. Pt alloys with many metals by fusing it with them; with Sb, As, Pb, Sn, and Zn heat and light are produced. Alloys which seem to be definite compounds have been obtained with arsenic (PtAs, Gehlen, Gm.-K. 3, 1192); antimony (PtSb, Christofle); cadmium (PtCd2, Deville a. Debray, A. Ch. [3] 56, 385); lead (PtPb, Bauer, B. 3, 836; 4, 449; D. a. D., C. R. 90, 1195); tin (Pt₂Sn₂, PtSn₄, and Pt₂Sn₃, D. a. D., A. Ch. [3] 56, 385; Debray, C. R. 104, 1470); and zinc (Pt₂Zn₂, D. a. D., l.c.). Alloys have also been formed with bismuth (Gehlen, Gm.-K. 3, 1192); copper (D. a. D., A. Oh. 1859. 611; Hélonis, B. 6, 42); gold (Dodé, B. 6, 1273); iridium (D. a. D., A. Ch. [3] 56, 385; Pelouze, C. R. 49, 896; Matthey, Pr. 28, 463; D. a. D., C. R. 81, 839; Morin, C. R. 78, 1502); iron (D. a. D., C. R. 89, 587; Daubrée, C. R. 80, 526); nickel (Hélonis, B. 6, 42); potassium (V. Meyer, B. 13, 392); silver (D. a. D., A. Ch. 1859. 611; H., l.c.); sodium (V. M., l.c.).

Amalgams of platinum. According to Crafts (Bl. [2] 49, 856), Hg has no action on Pt at the ordinary temperature, but Pt dissolves to a very small extent in boiling Hg, air being excluded. When acid is present Hg alloys with Pt (Casamajor, Am. 6, 540; Skey, C. N. 22, 282; Krouchkoll, J. de Ph. [3] 3, 139). The amalgams are most easily formed by adding sodium-amalgam to PtCl, Aq; with 13.5 p.c. Pt the amalgam is a thick liquid; with 25.8 p.c. Pt it is solid; an amalgam with c. 30 p.c. Pt has also been obtained (Joule, C. J. [2] 1, 378).

Platinum, antimonide of. An alloy of Pt and Sb, approximating to the composition PtSb, is obtained by melting the elements together in the ratio Pt: 2Sb; also by passing SbH, into H₂PtCl₈Aq, and removing ppd. Sb and PtCl₂ by washing with K₂SAq and KCNAq mixed (Christofle, Recherches sur les Combin. de l'Antimoine,

Göttingen, 1863).

Platinum, arsenides of. An alloy PtAs, is said to be formed by heating Pt black with As (Gehlen, Gm.-K. 3, 1192). By passing AsH. into PtCl,Aq, and heating the pp. in dry CO., Pt,As, is said to be formed (Tivoli, G. 1885.

487)

Platinum-arsenic hydroxide. According to Tivoli (G. 1885, 487) a compound PtAsOH is produced by passing AsH,, containing H but no other impurity, into PtCl,Aq (corresponding with 2 g. Pt in 60 c.c. water), and drying the black flocculent pp. at 120°-130°. The compound is decomposed by washing with alcohol or water; hot conc. H₂SO₄ at once separates Pt; heating in dry CO₂ produces As₂O₃, Pt₃As₂, and H₂O. Gibbs (Am. 8, 289) described some compounds of Pt and As,Os.

Platinum, boride of. Pt and B combine when melted together (Descotils, A. Ch. [3] 67, 88). It is best to heat Pt black and amorphous B under borax (Wöhler a. Debray, A. 101, 113). When excess of B is used, a crystalline compound containing 91.8 p.c. Pt., S.G. 17.3, is formed (Martius, A. 109, 79).

Platinum, bromides of. Pt and Br com-bine when heated together to above 300° (Langer a. Meyer, Pyrochemische Untersuchungen [Braunschweig, 1885], pp. 44, 57). Two-bromides are known, PtBr₂ and PtBr₄; the V.D.

of neither has been determined.

PLATINUM DIBROMIDE PtBr₂ (Platinous bromide.) Formed by heating HBrAq, b.p. (Platinous 126°, with Br and spongy Pt to 180° in a sealed tube-H2PtBrg.xH2O is thus formed-evaporating to dryness, heating to c. 200°, and washing the residue with boiling water (Topsoë, J. 1868. 273). Pullinger (C. J. 59, 602) heats spongy Pt with Br and HBrAq in a flask with a refluxcondenser, evaporates to dryness, and heats the residue to c. 280°. PtBr, is a brown powder: decomposes slowly at c. 800° in an air-current (P., l.c.); insol. water, sol. HBrAq and KBrAq. With KBr forms K2PtBr. (v. PLATINO-BROMIDES_

p. 282). Combines with CO at c. 180° to form PtBr2.CO, a bright red, crystalline solid: melts at 177.7°; not very hygroscopic; sol. alcohol; decomposed by water (Pullinger, l.c.). This com-

pound is called by P. carbonyl-bromoplatinite.
PLATINUM TETRABROMIDE PtBr. (Platinic bromide). Formed by heating spongy Pt with Br and HBrAq in a sealed tube to 180°—or by boiling Br, Pt, and HBrAq in flask with reflux-condenser (Pullinger, C. J. 59, 602)filtering, evaporating, heating residue at 180° till HBr ceases to come off; treating with boiling water, filtering from traces of PtBr2, evaporating, and drying at 180° (H. Meyer a. Züblin, B. 13, 404; Halberstadt, B. 17, 2963). A dark-brown, non-hygroscopic powder; sol. aqueous alcohol, more sol. absolute alcohol or ether, v. sl. sol. water (S. at $20^{\circ} = 41$) (H. l.c.); somewhat sol. glycerin. Pt black separates when solution in alcohol, ether, or glycerin is heated. With HBr forms H₂PtBr₆ (v. Platini-bromhydric acid, p. 282). With many metallic bromides forms salts M₂¹PtBr₆ (v. Platini-bromides, p. 282). When spongy Pt is heated with HBrAq and excess of HNO3, small, dark-brown, very hygroscopic crystals of platinum nitrosyl bromide, PtBr. 2NOBr, are obtained (Topsoë, J. 1868. 273).

Platinum, carbide of. Pt becomes brittle and more friable when heated to redness with charcoal. By heating the compound which PtCl. forms with acetone, Zeise (J. pr. 20, 209) obtained a black solid, to which he gave the com-

position PtC.

Two chlorides of Platinum, chlorides of. Pt are known, PtCl₂ and PtCl₄. When Pt black is heated in dry Cl at 240°-250°, PtCl₃ is formed, according to Schützenberger (C. R. 70, 1134, Pigeon (C. R. 108, 1009) says that at 360° PtCl, and PtCl, are formed, but the action is incomplete. Troost a. Hautefeuille (C. R. 84, 94) found that PtCl₂ was formed when Pt was heated in Cl to 1400°, and the tube was suddenly

PLATINUM DICHLORIDE PtCl. (Platinous V.D. not deterchloride. Platinochloride).

mined.

Formation.—1. By heating H2PtCls to c. 230° Berzelius, Gm.-K. 3, 1081).—2. By heating Pt black in Cl at 240°-250° till no more Cl is taken up (Schützenberger, C. R. 70, 1134, 1287).—
3. By decomposing Na,Pt(NaSO₃), by HClAq (Liebig, A. 23, 23; for formation of Na,Pt(NaSO₂), v. Platinosulphonates, p. 285).—4. A solution of PtCl₂ is obtained by passing SO₂ into PtCl₄Aq till

the liquid becomes red.

Preparation. -1. H2PtCl6Aq is evaporated to dryness, and the residue is heated in a bath of molten tin, i.e. at c. 230°, with constant stirring as long as Cl is given off (Berzelius, Gm.-K. 3, Or the heating is stopped before much of the solid is decomposed, the residue is dissolved in water, the dark-brown opaque liquid is evaporated, and the PtCl₂ which separates is dried at c. 150° (cf. Magnus, P. 14, 239).— 2. H.PtCls.6H2O is heated in vacuo, in presence of molten KOH, at 100° for 2 or 3 days; the temperature is then raised to 360°, and maintained thereat so long as Cl is given off (Pigeon, C. R. 112, 1218). Shenstone a. Beck (C. J. 61, 445) say that PtCl₂, prepared as directed above, Vol. IV.

always contains small quantities of some basic compound; and that when the salt is strongly heated small quantities of HCl and O are obtained, as well as Cl. Shenstone (C. J. Proc. 1892 93. 38) finds that PtCl₂ almost free from basic compounds can be prepared by heating PtCl, in a stream of dry HCl; a sample prepared in this way gave only 15 p.c. of HCl and O when decomposed by heat.

Properties and Reactions .- A brown powder; S.G. 5.87 at 11° (Bödeker). H.F. [Pt,Cl²] = 22,600 (Berthelot, C. R. 87,615). Insol. water, Insol. water, sol. HClAq in absence of air. Decomposed by heating to redness, giving off all Cl, and leaving Pt. According to Shenstone a. Beck (v. Preparation), the Cl thus obtained contains a little HCl and O, and H2O is also given off. Not acted on by HNO, Aq or dilute H, SO, Aq; decomposed by KOHAq, giving Pt(OH)2; by heating with conc. H2SO4, and then with water, Kane obtained a black powder to which he gave the composition PtCl₂.3PtO (B. J. 24, 238).

Combinations.—PtCl2 combines with many metallic chlorides; the compounds are described as platino-chlorides (p. 284). With HCl an acid H.PtCl, is formed (v. Platino-chlorhydric acid, p. 283). With CO the compounds PtCl, CO, PtCl, 2CO, and 2PtCl, 3CO are formed. These compounds are produced by passing alternate currents of Cl and CO over spongy Pt at 250° (Schützenberger, A. Ch. [4] 21, 350). Pullinger (C. J. 59, 598) found that a fourth compound is also formed—PtCl₂·2COCl₂ (or PtCl₃·2CO). Carbonyl chloroplatinite (PtCl₂·CO) combines with HCl, and with various hydrochlorides of organic bases (v. Mylius a. Förster, B. 24, 2424). PtCl₂ combines with PCl₃ to form PtCl₂.PCl₃ and PtCl₂.2PCl₃; and from these is derived a number of complex bodies. By treating PtCl, PCl, with H,O an acid PtCl, P(OH), is obtained, from which various salt-like compounds are derived; similarly PtCl₂.2PCl, yields an acid PtCl₂.P₂(OH)₀, and this gives many salt-like derivatives (Schützenberger, A. Ch. [4] 15, 100; 21, 350; Baudrimont, C. R. 53, 637; cf. PLATINO-CHLOROPHOSPHATES, p. 284). By heating PtCl_x.PCl_y with PtCl₂ Colchin (C. R. 86, 1402) obtained 2PtCl_x.PCl_y. For ammoniacal derivatives of the platino-phosphorus chlorides v. Colchin (l.c.) and Quesneville (M.S. [3] 6, 659), also Schützenberger (Bl. [2] 14, 97; 17, 482; 18, 101, 148).

PLATINUM TETRACHLORIDE PtCl. (Platinic chloride. Platinichloride). V.D. not determined.

Formation.-1. By heating Pt in Cl to c. 1700° (Langer a. Meyer, l.c. 44, 57); if a thin wire of Pt is heated by an electric current nearly to melting, in a stream of Cl, PtCl, is formed (Hodgkinson a. Lowndes, C. N. 58, 158, 223).— 2. By allowing H2PtCle.6H2O to remain over KOH, when it loses 4H₂O, and then heating in Cl to c. 350° (Pigeon, C. R. 110, 77).—3. By heating Pt and Se, mixed with AsCl, in Cl (v. Preparation).—4. PtCl. 5H2O is obtained by adding AgNO, Aq to H2PtCl, Aq in the ratio of H.PtCla:2AgNO, heating, filtering from AgCl, and evaporating finally over H.SO, (Norton, J. pr. [2] 2, 469; 5, 865; Engel, Bl. [2] 50, 100; Quesneville, M. S. [3] 6, 659).

Preparation.-1. Spongy Pt is mixed with rather less than its own weight of Se, the mix-

ture is placed in a tube of hard glass filled to one-third with AsCl,, and heated in a stream of Cl till the mass liquefies and boils, when the tube is sealed and heated for some hours at 250°; on cooling, the colourless crystals are separated from the pale-yellow crystals and the yellow liquid, and are heated in vacuo at 110°, whereby PtCl, and SeCl, remain; this residue is then heated at 360° in a stream of Cl, when SeCl, sublimes and PtCl, remains (Pigeon, C. R. 108, 1009).—2. Dry H₂PtCl₆ is placed in a porcelain boat which is heated, in a tube of hard glass, to 165° for about 15 hours in a stream of

dry HCl (Pullinger, C. J. 61, 422). Properties and Reactions. — H.F. [Pt.Cl] = 59,800; [Pt,Cl⁴,Aq] = 79,400 (Pigeon, C. R. 110, 77; 112, 791). A brown solid, said by Pigeon to be deliquescent; described by Pullinger as very soluble in water, but not deliquescent. The hydrate (v. Formation, No. 4) forms large, red, monoclinic crystals; according to Norton this compound has 5H2O, according to Engel 4H₂O; all H₂O except one molecule is removed at 100°, but the last molecule is not removable without decomposition. Decomposed by heating to dull redness (Pigeon, C. R. 110, 77); in presence of Cl, may be heated to c. 360°; in dry HCl slight decomposition occurs at c. 200° Rullinger, l.c.). HClAq produces H.PtCl₈Aq; NH,Cl ppts. (NH₄).PtCl₈. AgNO₃Aq ppts. Ag₂PtCl₈ from cold PtCl₄Aq, but on heating AgCl is formed and PtCl₄ goes into solution (Jörgensen, J. pr. [2] 16, 345). Pigeon (C. R. 112, 791) gives [Ag2PtCl4,Aq] = 12,160, to form PtCl,Aq and 2AgCl; PtCl,Aq is decomposed by light, for some measurements v. Foussereau (C. R. 103, 248).

Combinations.—1. With water (v. Formation, No. 4, and Properties). - 2. With hydrogen chloride to form H,PtCl, 6H,O; the compound HPtCl, 2H,O was obtained by Pigeon by heating H₂PtCl_s.6H₂O in vacuo at 100° (v. Platinichlorhydric acid, p. 282). Pigeon (C. R. 110, 77) gives [PtCl',2HClAq] = 24,800. — 3. With many metallic chlorides to form platini-chlorides (q. v., p. 282).—4. A compound with phosphorous chloride PtCl_{*}.PCl_{*} is formed by treating PtCl. PCl, with Cl (Schützenberger, Bl. [2] 14, 97; 17, 482; 18, 101, 148; cf. Baudrimont, C. R. 53, 637).—5. A compound with nitrosyl chloride, PtCl, 2NOCl.H,O, is one of the products of the action of fuming HNO, on H.PtCl.Aq (Weber, P. 131, 441).—6. Combines with alcohol, and with ethyl sulphide, to form PtCl, 2C2H6O and PtCl, 2(Et)2S respectively (Schützenberger, C. R. 70, 1134; Blomstrand, J. pr. [2] 17, 189).—7. Compounds with various organic bases and metallic chlorides, EtNH₂, &c., are described by Jörgensen (J. pr. [2] 83, 409).

Platinum, chloro-iodides of; v. PLATINUM IODOCHLORIDES, infra.

Platinum, cyanides of; and derivatives:

v. vol. ii. p. 343.

Platinum, fluoride of. Only one fluoride of Pt has been isolated, PtF4. Prepared by heating Pt wire in F, in a Pt or fluorspar tube, to c. 500°; if HF is mixed with the F combination occurs at the ordinary temperature (Moissan, C. R. 109, 807). Small buff-coloured crystals, or fused masses of a deep red colour. Dissolves in a little water, forming fawn-coloured solution, which almost at once decomposes, with rise of temperature and production of PtO, xH,O and HFAq. Heated to bright redness in Pt tube, PtF, is decomposed to F, and Pt which separates in crystals. A compound with PF, viz. PtF₄.2PF₅, is formed by passing PF₅ over spongy Pt heated to dull redness (M., Bl. [3] 5, 454). The decomposing action of HO on PtF, explains the failure of former attempts to prepare a fluoride of Pt in the wet way.

Platinum-iridium, or platin-iridium. An alloy of Pt and Ir which remains, mixed with osm-iridium, when many samples of Pt ore are heated with aqua regia (cf. IRIDIUM, vol. iii. p.

Platinum, hydride of. Pt black absorbs considerable quantities of H. According to Berthelot (A. Ch. [5] 30, 519), the relative weights of Pt and H are approximately 15:1 and 10:1; a considerable quantity of heat is produced. No certain evidence of the formation of hydrides has been obtained; but Thoma's result (Z. P. C. 3, 69), that the excess of H, above a definite quantity, absorbed by Pt when Pt is the negative electrode during the electrolysis of dilute sulphuric acid, is not easily given up, indicates the probable existence of a compound or compounds (cf. Palladium, vol. ii. p. 720).
Platinum, hydroxides of; v. Platinum,

OXIDES AND HYDROXIDES OF, p. 291.

Platinum, iodides of. Pt and I combine directly to form PtI,; the lower iodide PtI, has

also been obtained, but not pure.

PLATINUM DI-IODIDE PtI₂ (Platinous iodide. Platino-iodide). This compound is described as a black, heavy powder, insol. water, alcohol, and acids; giving Pt and I at 300°-350°; decomposed by NaOHAq to PtO₂H₂. It is produced by boiling PtCl₂ with fairly conc. KIAq (Lassaigne, A. Ch. [2] 51, 113); but it has not been obtained pure (Clementi, J. 1858, 420; Topsos, Ar. Sc. 38, 297). The compound PtI, CO, carbonyl iodoplatinite, is described by Mylius a. Förster (B. 24, 2424).

PLATINUM TETRA-IODIDE PtI, (Platinic iodide. Platini-iodide). V.D. not determined. Prepared by heating PtO₂ with HIAq (Clementi, J. 1855. 420); also by the action of cold HIAq on H₂PtCl₆.6H₂O (Topsos, Ar. Sc. 38, 297), and evaporating. Pullinger (C. J. 59, 602) obtained PtI, by dissolving spongy Pt in a hot solution of I in HIAq, evaporating to dryness, heating to 180°, and washing with water. A brown-black amorphous powder, insol. water; sol. HIAq and metallic iodide solutions (Topsoë, l.c.); not wholly decomposed by heating with Na, CO, to the m.p. of the carbonate (Pullinger, l.c.). Combines with HI to form H_PtI_0.9H_O (v. Plating-IODHYDRIC ACID, p. 283), and with metallic iodides to form salts (v. Platini-iodides, p. 283).

Platinum, iodochlorides of. A compound PtI2Cl2 is said to be formed, in large red plates, by dissolving Pt and I in aqua regia and evaporating at 100° (Kämmerer, A. 148, 329). Another compound PtI Cl was obtained by Mather, as a black powder, by evaporating H.PtCl, with a small excess of HIAq, and heating the residue

to 150° (Am. S. 27, 257).

Platinum-nitrosyl bromide and Platinumnitrosyl chloride, v. Platinum Tetrabeomide. p. 289; and Platinum Tetrachloride, Combina-

tions, No. 5, p. 290.

Platinum, oxides and hydroxides of. Three oxides have been isolated; PtO, Pt₃O₄, and PtO₂; hydrates of all are known. PtO forms a few salts; it also dissolves in molten KOH and NaOH. PtO, forms corresponding salts, and also combines with alkaline oxides to form

platinates (q. v. p. 281).

PLATINUM MONOXIDE PtO (Platinous oxide). Obtained as a violet powder, by strongly heating the pp. formed by CaOAq acting on H2PtCl,Aq in sunlight, and washing with water and then with HNO₃Aq (Döbereiner, P. 28, 181). Also by carefully heating PtO₂H₂. Reduced to Pt at red heat, also by action of reducers such as HCO2HAq. Dissolves in molten KOH and NaOH, but the compounds thus formed have not been examined. A compound which is perhaps Pto.co is described by Mylius a. Förster (B. 24, 2440).

PLATINOUS HYDROXIDE PtO, H, was obtained by Thomsen (J. pr. [2] 16, 294) by adding NaOHAq to K₂PtCl₄Aq (1:12) in the ratio 2NaOH:K₂PtCl₄ and heating to boiling. A black powder, sol. HClAq, HBrAq, and SO₂Aq, forming corresponding salts. Freshly ppd. PtO₂H₂ dis-

solves in excess of NaOHAq.

TRI-PLATINUM TETROXIDE Pt.O. (Platinoplatinic oxide). A black powder, insol. acids; obtained by heating dehydrated Na₂PtCl₂ with 4 parts dry Na₂CO₃ till fusion begins, boiling with water, then with dilute HNO₃Aq, and finally with aqua regia (Jörgensen, J. pr. [2] 16, 844). Loses its O at red heat; easily reduced to Pt by H or coal-gas, even without heating. Prost (Bl. [2] 46, 156) describes a hydrate Pt, O. 9H, O.

PLATINUM DIOXIDE PtO₂ (Platinic oxide). A black powder, obtained by gently heating PtO₂H₄. PtO₂ is insol. acids, but salts of this

oxide are formed from the hydroxide.

PLATINIC HYDROXIDE PtO, H,. Formed by boiling PtCl4Aq for a considerable time with a large excess of NaOHAq, acidifying by acetic acid, drying the white flocculent pp. in the air, and heating the yellow hydrate, PtO,H,2H,O, to 100° (Topsoë, B. 3, 462; Fremy, A. Ch. [3] 31, 478). Also obtained by evaporating H.PtCl.Aq with excess of Na.CO., rubbing up the residue in water, and treating with acetic acid (Döbereiner, P. 28, 181; Topsoë, l.c.; cf. Wittstein, R. P. 74, 43). A hydrate PtO₄H₄.H₂O was obtained by Prost (Bl. [2] 46, 156) by adding water to a solution of PtO₄H₄ in conc. HNO, Aq. A reddish-brown powder; dehydrated by heating gently; at a higher temperature gives Pt and O. Heated in H gives Pt. Sol. NaOHAq, HClAq, HNO₂Aq, and H₂SO₄Aq. With acids forms platinic salts, PtX₄, where $X = \frac{1}{2}SO_4$, $\frac{1}{2}CO_2$, With acids NO, &c.; with strongly basic oxides forms platinates, PtO2.xMO (v. PLATINATES, p. 281). Compounds with SnO and SnO, are described by Delachanel a. Mermet (C. R. 81, 370), Schneider (P. 136, 105), Lévy a. Bourgeois (C. R. 94, 1365), and Schützenberger (C. R. 98, 985). PtO4H dissolves in solutions of certain molybdates and tungstates, forming platini-molybdates and platini-tungstates (q. v. p. 283).

Platinum, oxychlorides of. When AgNO, Aq

is added to PtCl,Aq in the ratio PtCl,:2AgNO, or when AgNO, Aq and H.PtCl.Aq are heated tion of an alkali hydrosulphide. Also formed

together in the ratio H2PtCls:4AgNO, a pp. is obtained which has the empirical composition PtCl₂(OH)₂.2AgCl; this substance may be regarded as a compound of the hydrated oxychloride PtOCl₂.H₂O (Jörgensen, J. pr. [2] 16, 345). The pps. formed by CaOAq and BaOAq in PtCl,Aq are supposed to contain oxychlorides of Pt (v. Johannsen, A. 155, 204).

Platinum, oxysulphide of, 2PtOS.H.O. PtS. slowly oxidises by exposure to air; by washing the product (to dissolve H_aSO_4 and SO_2) and drying in a stream of CO_2 , a black powder is obtained having the composition 2PtOS.H2O (perhaps HO.PtS.O.SPt.OH). This substance is an energetic oxidiser; H, CO, H,S, SO, take fire when brought into contact with it; NH, is absorbed and oxidised to HNO2 and HNO3 (E. von

Meyer, J. pr. [2] 15, 1).

Platinum, phosphides of. Pt and P combine when strongly heated together. Schrötter obtained a grey metal-like mass, S.G. 8.77, to which he gave the composition PtP₂, by heating Pt black in vapour of P (J. 1849. 246). By heating Pt and P to white heat, Clarke a. Joslin (Am. 5, 231) obtained Pt₃P₅; and by heating in a mussle they got Pt.P. sol. aqua regia but changed to insoluble PtP.

Platinum, salts of. But few salts, excepting the halides, are known wherein Pt replaces the H of acids; Pt more frequently enters into the acid radicle, than the positive radicle, of salts.

Platinum, selenide of. By heating Pt and Se, Berzelius obtained a grey, infusible powder; perhaps PtSe (Gm.-K. 3, 1081).

Platinum, selenocyanides of; for a double

salt v. vol. ii. p. 348.

Platinum, silicides of. A white, crystalline, brittle solid, Pt.Si., was obtained by Guyard (Bl. [2] 25, 510) by heating together equal parts of Pt black and powdered crystalline Si (cf. Winkler, J. pr. 91, 203). According to Miles (Am. 8, 428), the conditions favourable to the formation of compounds of Pt and Si are a high temperature, the presence of a reducer such as CO, a basic substance such as MgO, and SiF,, in contact with Pt (v. also Brown, Am. 7, 173).

Platinum, sulphides of. Pt and S combine

when heated together. Two sulphides have been isolated, PtS and PtS₂; there is also evidence

of the existence of Pt.S.

PLATINUM MONOSULPHIDE PtS (Platinous sulphide). Obtained by heating Pt black and S, and volatilising excess of S, in absence of air; also by heating (NH4), PtCl, with S, out of contact with air, till NH,Cl and excess of S are volatilised; also by passing H2S into PtCl,Aq (Berzelius). Prepared in crystals by heating to redness a mixture of 1 pt. Pt, 1 pt. borax, and 10 pts. pyrites (Deville a. Debray, A. Ch. [2] 55, 215). A grey, lustrous solid; S.G. 6.2. PtS prepared in the wet way is a black powder. changed in air, even when moist; unacted on by boiling acids; loses S when heated in air, leaving Pt; reduction in H begins at 19°. A compound with CO-PtS.CO-was obtained by Mylius a. Förster (B. 24, 2438) by the action of H₂S on PtCl_CO.

PLATINUM DISULPHIDE PtS. (Platinic sul-Obtained by passing H.S into phide). Obtained by passing H.S into Na.PtCl.Aq, or by dropping H.PtCl.Aq into solu-

by digesting, in a strong closed flask, 1 pt. PtCl4, 4 pts. alcohol, and 5 pts. CS₂ (Böttger, J. pr. 3, 274); and by decolourising H₂PtCl₈Aq by SO₂, and then heating in a sealed tube at 200° (Geitner, A. 129, 358). PtS2 is also a product of the action of HClAq in air on Na,Pt,S_a or H,Pt,S_a (v. Thioplatinates, p. 281). Colloidal soluble PtS₂ was obtained by Winssinger (Bl. [2] 49, 452) by passing H2S into very dilute PtCl4Aq, and dialysing. A black solid; as obtained from Na,Pt,S, forms steel-grey needles, S.G. 5.27. PtS, prepared by one of the wet methods must be dried in absence of air, else it is partially decomposed. Heated out of air forms PtS, and Pt and S. Oxidised to sulphate by strong oxidisers, such as aqua regia, Cl, fuming HNOs, molten KClO₃ or KNO₃. Freshly ppd. PtS₂ dissolves in alkali sulphide solutions forming thioplatinates (q. v.), from which solutions it is reppd. by acids. Classon (J. pr. [2] 15, 193) describes a compound with EtSH.

PLATINUM SESQUISULPHIDE, Pt_2S_3 . This compound is said to be formed, as a steel-grey, metal-like solid, S.G. 5·52, by the oxidation in air of $H_2Pt_4S_3$ (Schneider, P. 136, 105; 138, 604; 139, 661; 141, 519; 148, 633; 149, 381). Schneider also describes a compound $4PtS.PtS_2$

 $= Pt_5S_6$.

Platinum, sulphocyanides of; v. vol. ii. p. 357; v. also Guareschi, Giorn. della R. Accad. di Med. 1891. No. 5; abstract in B. 25, Ref. 7.

Platinum, thicoarbide of. Pt₂CS₂. This compound is described by Schützenberger (C. R. 111, 391). It is a black powder; obtained by passing a stream of N, or H, charged with CS₂, over spongy Pt heated to 400°-450°.

M. M. P. M.

PLATINUM - AMMONIUM COMPOUNDS (Ammonio-platinum compounds. Ammoniacal

(Ammono-platinum compounds. Ammoniacat platinum bases. Platinammines). When a solution of PtCl₂ in HClAq reacts with NH₃, more than one compound of the form PtCl₂.2NH₃ is obtained. These compounds react with acids, oxidisers, &c., to form many derivatives containing Pt, N, H, and negative radicles, in which the reactions of Pt and NH₂ are more or less modified.

The platin-ammonium compounds may be regarded as salts of bases derived from two or more NH₄ groups, in which part of the H is replaced by Pt; thus PtCl₂·2NH₃ may be called platoso-diammonium chloride, and formulated as N₂H₆Pt.Cl₂. Similarly PtCl₂·4NH₃ may be

regarded as Pt < $NH_2(NH_4)Cl$, and called ammo-

uium platoso-diammonium chloride. The compounds derived from PtCl₄ are regarded, in this scheme, as platini-ammonium compounds; thus, PtCl₄:2NH₃ is Cl₂Pt.NH₂.NH₄.Cl₂ (v. Hofmann, T. 1851. [2] 357; Weltzien, A. 97, 19; Kolbe, J. pr. [2] 2, 217; Grimm, A. 99, 67).

This view of the constitution of the compounds in question was opposed by Claus (*J. pr.* 63, 99), Blomstrand (*B.* 4, 40, 639, 673; 6, 1469), Jörgensen (*J. pr.* [2] 33, 489), and others. Jörgensen found that pyridine, N.C., H., forms a compound with platinous chloride similar to PtCl., 4NH.; in this pyridine compound, which has the composition PtCl., 4NC, H., the atoms of hydrogen cannot be in direct union with N; for

the constitution of pyridine is known to be

 $HC \bigvee_{N} CH$. Now if the ammoniacal platinum

bases are all to be derived from NH₄, by substituting H by NH₄ groups, it is evident that the substituted H atoms must always be directly bound to N atoms. But the existence of the pyridine compound shows that this is not necessary. Hence PtCl₂.4NH₂ may be regarded as NH₂.NH₂.NH₂.NH₂.NH₂.NH₂.NH₃.NH₂.NH₃.NH₂.NH₃

Pt<NH3.NH3Cl NH3.NH3Cl

The classification and nomenclature of the platin-ammonium compounds proposed by Blomstrand (Chemie der Jetztzeit, Heidelberg, 1869; B. 2, 202; 4, 40, 639, 673; 6, 1469) is now usually adopted. In the following formulæ Ristands for a monovalent negative radicle. Blomstrand uses the term ammine to distinguish these compounds from the amines, which contain the group NH₂.

Class I. PLATO- or PLATOSO- COMPOUNDS.

Series 1. Platosemiammines, RPtNH₃R.
, 2. Platosammines, Pt(NH₃R)₂.

3. Platodiammines, Pt(NH₂.NH₃R)₂.

, 4. Platosemidiammines, Pt(NH₃.NH₃R)R.

,, 5. Platomonodiammines, Pt(NH₃.NH₃R)(NH₃R).

Series 2 and 4 are isomeric. Cleve has isolated two distinct aniline compounds, Pt(NH_{*})₂·(C₆H₅NH₂)₂·R₂ (cf. Jörgensen, J. pr. [2] 33, 489).

Class II. PLATINI- or PLATIN- COMPOUNDS. Series 1. Platinammines, R₂Pt(NH₂R)₂.

,, 2. Platinidiammines, R₂Pt(NH₃:NH₃R)₂.

3. Platinisemidiammines, R₂Pt(NH₃.NH₃R)R.

, 4. Platinimonodiammines, R,Pt(NH₃.NH₃)R(NH₃R). 5. Platinitriammines,

 $R_2Pt(NH_3.NH_3.NH_3R)_2$

Series 1 and 3 are isomeric.

The prefixes mono, di, &c., are used to denote the number of NH_3 groups in direct union with one another, and not the number of NH_3 groups in union with the atom of Pt. As the compounds in Series 3 contain one diammine chain, (NH_3,NH_3) , they are called semidiammines; and as those in Series 4 contain a diammine and a monammine chain, they are called monodiammines.

Class III. DIPLATINUM COMPOUNDS.

These contain 2 Pt atoms; the group Pt, may be divalent, tetravalent, or hexavalent.

Series 1.

Diplatodiammines, Pt(NH, NH, R) | Pt(NH, NH, R)

Series 2.

Pt(NH₂.NH₂R)

RPt(NH₂.NH₃R)

RPt(NH₃.NH₄R)

or

Pt(NH₃.NH₄R)

R.Pt(NH,.NH,R)

Series 3.

Diplatinammines,

RPt(NH₃R)₂ | RPt(NH₃R)₂

Series 4.

RPt(NH₈.NH₃R)₂

Diplatinidiammines,

RPt(NH3.NH3R)2

Iodides of platinum bases with 4 and 8 atoms Pt are also known; I₂Pt₄.8NH₃.I₈, and I₂Pt₄.16NH₃.I₁₆.

Many organic bases, such as aniline, pyridene, &c., can take the place of NH₃ in these compounds, and the N of NH₃ is often replaceable by P or As.

Class I. Plato- or Platoso- compounds.

Series 1. Platosemiammines, RPtNH₃R. None of these has been isolated, but Cossa (B. 23, 2503) has obtained a compound which probably contains platosemiammine chloride; the compound is 2PtNH₃Cl₂.Pt(NH₃)₄Cl₂.

Series 2. Platosammines, Pt(NH₂R)₂. These compounds are generally obtained by heating the corresponding platodiammines, Pt(NH₂,NH₂R)₂, which thus lose 2NH₃; by heating with NH₂Aq, platodiammines are re-formed; with oxidisers, platini- compounds are obtained.

PLATOSAMMINE CHLORIDE Pt.NH₃Cl.NH₄Cl (Peyrone, A. 51, 1; 55, 205; 61, 178). Obtained by heating Pt(NH₃),Cl₂ (plato-diammine chloride) to 250°, or by evaporating that compound with HClAq at 100° and dissolving out AmCl in water; also by evaporating with HCl the solution of the corresponding nitrate or sulphate. Microscopic, sulphur-yellow crystals. S. 022 at 0°, '77 at 100°. Sol. NH₃Aq, forming Pt(NH₃,NH₃Cl)₂ (platodiammine chloride); in aqua regia forms Cl₂Pt(NH₃Cl)₂ (platinammine chloride). With AgNO₃Aq, in the ratio 2Pt(NH₃Cl)₂:AgNO₃, one half of the chloride remains unchanged and one half gives Pt(NH₃,NO₃)₂. Heated to 270°, the chloride is decomposed thus:

 $3Pt(NH_3Cl)_2 = 3Pt + 4NH_4Cl + 2HOl + N_2$. Platosemidiammine chloride

Pt(NH₃.NH₃Cl)Cl, and platodiammine chloride platinous chloride Pt(NH₃.NH₂Cl)₂.PtCl₂(Magnus' green salt), are isomeric or polymeric with platosammine chloride. Other compounds, perhaps also isomeric, were obtained by Peyrone (l.c.) and by Cossa (B. 23, 2503).

Grimm (A. 99, 67) obtained a double compound with AmCl; Pt(NH₃Cl)₂2AmCl. Jörgensen (J. pr. [2] 33, 489) obtained the pyridine compound Pt(NC₃H₄Cl.)₂; and also the mixed compound Pt(NC₃H₄Cl.NH₃Cl (v. also Hedin, Dissertation, Lund; B. 20, Ref. 108).

PLATOSAMMINE HYDROXIDE

Pt.NH₁OH.NH₂OH. Said to be obtained, as a crystalline, very soluble solid, by decomposing the sulphate, Pt(NH₂.NH₂)SO₄, by BaOAq (Oding, C. N. 21, 269, 289). The compound thus formed may have been the isomeric platosemidiammine hydroxide, OH.Pt(NH₂.NH₂OH). A solution of Odling's hydroxide reacts strongly alkaline, absorbs CO₂ from the air, neutralises acids, decomposes salts of NH₄ giving off NH₄, and ppts. many metallic hydroxides from salts of the metals.

PLATOSAMMINE OXIDE Pt(NH, NH,)O.

grey solid, insol. water or NH₃Aq; heated in a retort to 195° gives N, NH₃, H₂O and Pt. Obtained by heating platodiammine hydroxide (Pt[NH₃NH₃.OH]₂) to 110° (Reiset, A. Ch. [3] 11, 417).

PLATOSAMMINE SULPHONATES and CHLORO-SULPHONATES Pt.NH₃SO₃M.NH₃SO₃M and Pt.NH₄Cl.NH₈SO₃M. Ammonium platosammine sulphonate, Pt(NH₃SO₃Am)₂, is formed by the reaction of excess of Am₂SO₃Aq on Pt(NH₅Cl)₂ (Peyrone, A. 51, 1; 55, 205; 61, 178). The salts of Ba, Co, Cu, Pb, Mn, Ni, Ag, Na, UO₂, and Zn have been obtained. Ammonium platosammine chlorosulphonate, Pt.NH₅Cl.NH₃SO₃Am.H₂O, is obtained, in colourless rhombic tablets, by leading SO₂ into a boiling solution of Pt(NH₃Cl)₂, and neutralising with NH₃Aq. If the neutralisation is omitted, platosammine chlorosulphonic acid, Pt.NH₂Cl.NH₃SO₃H, is obtained (Peyrone, Lc.).

Lc.).

The other salts of the platosammine series which have been isolated are the following; \$M = Pt(NH_3)_2: -bromide, MBr_2 (Jörgensen, J.pr. [2] 33, 521, 531, 535); cyanide, MCy_2 (Buckton, A. 92, 280); iodide, MI_2 (Cleve, Bl. [2] 7, 12; 17, 482; Reiset, A. Ch. [3] 11, 417); nitrate and nitrite, M(NO_3)_2 and M(NO_2)_2 (Cleve, l.c.); sulphate and sulphite, MSO_4.H_2O and MSO_8.H O (Cleve, l.c.); and sulphocyanide, M(CyS)_2 (Buckton, l.c.). Substituted derivatives of the bromide, of the form Pt(NH_2RBr)_2 and Pt.NH_3Br.NH_RBr, where R = Et, Me, or Pr, have been obtained (Jörgensen,

l.c.). Series 3.—Platodiammines

Pt(NH₃.NH₃R)₂. Formed by action of NH₃Aq on platinous salts, platosammine or platosemidiammine compounds. Easily lose NH₃ on heating, giving platosammine compounds. Give platinidiammine compounds by action of oxidisers.

PLATODIAMMINE CHLORIDE

Pt(NH₃,NH₂Cl)₂,H₂O (Reiset, A. Ch. [8] 11, 417). Sometimes called Reiset's first chloride. Obtained by the prolonged action of boiling NH, Aq on PtCl₂, Magnus's green salt (v. infra), platosammine chloride Pt(NH,Cl)2, or platosemidiammine chloride Pt(NH, NH, Cl)Cl, precipitation by alcohol, and crystallisation from water. Also formed by reduction of platinidiammine chloride $Cl_2Pt(NH_3NH_3Cl)_2$, by H_2S (Thomsen, J. 1868. Large, colourless, tetragonal crystals; S. 25 at 16.5°, more sol. hot water; insol. absolute alcohol and ether. Addition of H₂SO₄Aq, HNO₂Aq, or H₂C₂O₄Aq ppts. the corresponding sulphate, nitrate, or oxalate. By gently warming with HNO, Aq the salt (NO,)2Pt(NH,.NH,Cl)2 is formed; Cl, Br, and FeCl, Aq react similarly. 240°-270° NH, is given off, and Pt(NH,Cl)2 is formed (v. Platosammine chloride, supra). Combines with PtCl2, and PtCl4, to form M.PtCl2 and M.PtCl₄ respectively [M = Pt(NH₃.NH₃Cl)₂]. Compounds are known in which NH, is wholly or partly replaced by pyridine, also by alkyl radicles (Jörgensen, J. pr. [2] 33, 489), and also by PEt₂ (Cahours a. Gal, C. R. 70, 1381).

Double compounds. Combines with various metallic chlorides to form compounds of the form M.XCl₂, where M = Pt(NH₂,NH₂Cl)₂, and X = Pt, Cu, Pb, Hg, Sn, and Zn (v. Thomsen, J. 1868. 278; Buckton, C. J. 5, 213; Millon a. Commaille, C. R. 57, 822).

The most important of these compounds is PLATODIAMMINE CHLORIDE PLATINOUS CHLORIDE,

known as Magnus's green salt,

the action of NH₂Aq on PtCl₂ in HClAq, or on PtCl₄ in HClAq after passing in SO₂ till AmCl ceases to give a pp. (Magnus, P. 14, 242; Gros, A. 27, 241). Also by adding Pt(NH₃.NH₂Cl), to PtCl₂ in HClAq. It is best prepared by making Am₂PtCl₄, from PtCl₄ in HClAq treated with SO₂ and crystallised from AmClAq, and heating this with HClAq and NH₂Aq (Claus, A. 107, 138). Green, microscopic prisms or needles; insolwater, alcohol, and dilute HClAq. Digestion with hot NH₂Aq forms Pt(NH₃.NH₂Cl)₂; Cl, Br, or HNO₂Aq produces platinidiammines (Gros, Lc.; Raewsky, A. Ch. [3] 22, 278). Boiling with AgNO₂Aq produces PtCl₂ and Pt(NH₃.NH₂NO₃)₂. If the empirical formula nPt(NH₃)₂Cl₂ is given to Magnus's green salt, it is seen to be isomeric, or polymeric, with platosammine chloride Pt(NH₂Cl)₂, platosemidiammine chloride Cl.Pt.NH₄.NH₄Cl,

platomonodiammine chloride platinous chloride 2[Pt(NH, Cl)(NH, NH, Cl)].PtCl₂,diplatosindiammine chloride Cl₂Pt₂(NH, NH₃Cl)₂, and with a double salt obtained by Cossa (B. 23, 2503),

2[Pt.NH₃.Cl₂].Pt(NH₃)₄Cl₂.

PLATODIAMMINE HYDROXIDE Pt(NH₂NH₂OH)₂. Sometimes called Reiset's first base (Reiset, A. Ch. [8] 11, 417). White, crystalline, deliquescent needles; by decomposing the sulphate by the proper quantity of BaOAq, filtering, and evaporating in vacuo. Solution in water is strongly alkaline, absorbs CO₂ from the air, and drives out NH₂ from NH₄ salts. NH₃ is not given off when an aqueous solution is boiled alone or with potash. Melts at c. 110°; at a higher temperature gives off NH₃ and H₂O and leaves platosammine oxide Pt(NH₃)₂O (q. v. p. 293).

The other chief compounds of the platodiammine series are the following; $M = Pt(NH_3)_4$:—carbonates, MCO_2 .4aq and $M(CO_2H)_2$ (Peyrone, A. 51, 14; Reiset, A. Ch. [3] 11, 417); chromates $MCrO_4$ and MCr_2O_7 ; bromide, MBr_2 .3aq (Cleve, J. 1867, 321); iodide, MI_2 .xaq (Cleve); nitrate and nitrite, $M(NO_3)_2$ and $M(NO_2)_2$.2aq (Peyrone, l.c.); phosphate, $MHPO_4$.aq (Cleve, Bl. [2] 7, 12; 17, 482); sulphate and sulphites, MSO_4 ,

MSO₂, M(SO₃H)₂.2aq (Cleve, l.c.).

Series 4. Platosemidiammines, Pt(NH₂,NH₂R)R. Generally formed from the chloride by double decomposition: the chloride is a product of the action of NH₂Aq or (NH₄)₂CO₂Aq on PtCl₂. Prolonged treatment of the platosemidiammines with NH₃Aq produces diammine compounds. Jörgensen (J. pr. [2] 33, 489) has replaced NH₃ in these compounds by alkyl radicles and by pyridine.

PLATOSEMIDIAMMINE CHLORIDE Pt. NH₂.NH₂.NH₂.Cl
Sometimes called *Peyrone's chloride*. PtCl₂ is formed by the action of SO₂ on PtCl₄ in HClAq, and the solution is heated to boiling with excess of (NH₂)₂CO₂ until the red solution has become light-yellow and a greenish-brown pp. forms, when the liquid is rapidly filtered. The chloride separates as the liquid cools; it must be filtered

off at once (elge it may re-dissolve), and crystallised from boiling water (Peyrone, A. 51, 14; 55, 209; v. also Thomsen, Gm.-K. 8, 1115; Cleve, Bl. [2] 7, 12, 17, 482). Small yellow crystals. S. ·26 at 0°, 3·6 at 100°. Decomposes at 270°, giving off NH, and HCl. Boiling NH, Aq forms platodiammine chloride, $Pt(NH_x.NH_xCl)_2$; Cl, Br, I, and some other oxidisers produce platinisemidiammine compounds, $R_xPt(NH_x.NH_xR)R_x$. With AgNO_x, and Ag,SO₄, the corresponding nitrate and sulphate are formed.

It is doubtful whether the hydroxide Pt(NH,NH,OH)OH has been prepared; the compound obtained by Odling (C. N. 21, 269, 289) was this or the isomeric platosammine hydroxide, Pt(NH,OH)₂ (v. Platosammine Hydroxide, Pt(NH,OH)₂ (v. Platosammine Hydroxide, Pt(NH,OH)₂ (v. Platosammine Hydroxide, Pt(NH,OH)₂ (v. Platosammine Hydroxide, p. 293). The other compounds of this series which have been obtained are the following; M = Pt.NH, NH,: -bromide. MBr₂; chloroxide, MGl.SO₃H (Cleve. l.c.); iodide, MI₂; nitrate and nitrite, M(NO₃)₂ and M(NO₂)₂; sulphate, MSO₄; double sulphites, MSO₂-2X₂SO₃, X₂=2NH₄, Ba, Co, Ag₂; these may be regarded as compounds of sulphites with salts of the hypothetical platosemiammine sulphonic acid Pt(NH₂,NH₃,SO₂H)SO₃H.

Series 5. Platomonodiammines Pt(NH,NH,R)(NH,R). Most of the salts of this series are prepared from the double salt of the chloride with PtCl₂.

PLATOMONODIAMMINE CHLORIDE PLATINOUS CHLORIDE 2Pt
NH₃·NH₃·Cl
.PtCl₂. This compound is one product of the action of NH₄Aq on PtCl₂ in HClAq; it is prepared by neutralising PtCl₂ in HClAq by (NH₄)₂CO₃Aq, heating to boiling, and adding (NH₄)₂CO₃Aq drop by drop, when it separates in small red tablets (Peyrone, A. 55, 209). Fairly sol. cold water, more sol. hot water. Boiled with NH₃Aq forms Magnus's green salt, Pt(NH₃.NH₃Cl)₂.PtCl₂. Isomeric, or polymeric, with Magnus's green salt, platosammine chloride, and platosemidiammine chloride.

PLATOMONODIAMMINE CHLORIDE

 $Pt < NH_aNH_aCl.$ Obtained, in lustrous colourless crystals, by adding HClAq to an ice-cold solution of the nitrate.

PLATOMONODIAMMINE NITRATE

Pt<\NH,NH,NO. Formed by decomposing the chloride in solution by AgNO.Aq, filtering, and evaporating. The sulphate

Pt<\(\frac{\text{NH}_2\text{NH}_3}{\text{NH}_2}\)\SO_4.H_2O is also known (v. Cleve, l.c.; Blomstrand, B. 4, 40, 639, 673; 6, 1469).

Class II. Platini- or Platin- compounds. Series 1. Platinammines, R.Pt(NH₂R)₂. These compounds are formed by the addition of Cl, Br, or I to platosammines, or by the action of such oxidisers as HNO₂Aq on platosammines; they are generally changed to platinidiammines by heating with NH₂Aq, and are reduced to platosammines by SO₂Aq. The radicles in direct union with Pt are replaced with more difficulty than those in union with NH₂. The nomenclature is arranged to indicate, first, the nature of the radicles in direct union with Pt, and then those in direct union with NH₂.

I. Chloro- compounds Cl.Pt(NH.B),
CHLOROPLATINAMMINE CHLORIDE Cl.Pt(NH.Cl),
Formed by suspending platosamr-ine chloride,

Pt(NH,Cl), in boiling water, and passing in Cl until the original citron-yellow particles are changed to microscopic yellow octahedra (Gerhardt a. Laurent, C. R. 30, 273), or until the liquid, which is yellow at first, begins to turn reddish. Aqua regia, or KMnO Aq and HClAq, may be used instead of Cl (Odling, C. N. 21, 269, 289). A heavy, yellow, crystalline powder. S. 1 at 0°; 3 at 100°. Unacted on by boiling conc. HNO3 or H2SO4; sol. boiling KOHAq without evolution of NH, sol. NH, Aq, with formation of chloroplatinidiammine chloride, Cl.Pt(NH₃.NH₃Cl)₂. The Cl is ppd. as AgCl only after long boiling with AgNO₃Aq (Grimm, A.

CHLOROPLATINAMMINE NITRITE Cl₂Pt(NH₃,NO₂)₂. Ppd. in colourless rhombic plates by addition of excess of HClAq to solution of the nitrato-nitrite, (NO_s)₂Pt(NH_s.NO₂)₂, which is formed by boiling platosammine nitrite, Pt(NH3.NO2)2, with HNO3Aq.

CHLOROPLATINAMMINE CHLORONITRITE Cl₂Pt(NH₃Cl)(NH₃·NO₂), is produced by the reaction of K₂PtCl₄ and (NO₃)₂Pt(NH₂NO₂)₂.

II. Hydroxyl compounds

(OH)2Pt(NH3R)2.

HYDROXYL PLATINAMMINE HYDROXIDE

(OH),Pt(NH3.OH)2. Yellow, lustrous crystals; by adding excess of NH₈Aq to a boiling solution of the nitrato-nitrate (NO₃)₂Pt(NH₃.NO₃)₂, and allowing to cool. Scarcely sol. water; easily sol. dilute acids. Boiling KOHAq does not evolve NH₃; decomposes above 130°, giving off NH₃ and H₂O, and leaving Pt (Gerhardt a. Laurent, C. R. 30, 273).

HYDROXYL PLATINAMMINE NITRATE (OH)₂Pt(NH₃.NO₃)₂. 2aq. Crystalline pp. by decomposing Cl₂Pt(NH₃Cl)₂ by AgNO₃Aq. Not acted on by cold HClAq; evaporation with HNO₂Aq produces (NO₃)₂Pt(NH₃,NO₃)₃ (G. a. L., l.c.; Cleve, Bl. [2] 7, 12; 17, 482).

HYDROXYL PLATINAMMINE SULPHATE $(OH)_2Pt(NH_3)_2SO_4$ aq. Formed as a pale-yellow, hard crust of needle-shaped crystals, by decomposing Cl₂Pt(NH₂.Cl)₂ by Ag₂SO₄Aq. Evaporation with H₂SO₄Aq produces the sulphato-sulphate, SO₄.Pt.(NH₂)₂SO₄.3aq; when HClAq is used, Cl₂Pt(NH₂)₂SO₄ and Cl₂Pt(NH₂Cl)₂ are produced (Cleve, l.c.).

The following salts of this series have also been isolated: Bromo-bromide, Br₂Pt(NH₂Br)₂ (Cleve, 1801sted: Bromo-oromias, Br₂Pt(NH₂Br)₂ (Cleve, l.c.); bromo-nitrite, Br₂Pt(NH₂NO₂)₂ (Cleve, l.c.); iodo-iodide, I₂Pt(NH₂I)₂ (Cleve, l.c.); nitrato-nitrate, (NO₃)₂Pt(NH₂NO₃)₂; nitrato-nitrite, (NO₃)₂Pt(NH₂NO₃)₂; nitrato-chloronitrite, NO₃Cl.Pt(NH₂NO₃)₃ (Cleve, l.c.); sulphato-sulphate, SO₄.Pt(NH₂)₂SO₄.3aq (Cleve, l.c.)

Series 2. Platinidiammines, R₂Pt(NH₃NH₃R)₂. Generally formed by action of Cl, Br, I, and other oxidisers on platodiammines. A large number of these compounds is known. R attached directly to Pt may be the same as, or different from, R united directly to NH₂. The nomenclature is similar to that of the platinammine series. The radioles in direct union with Pt are removed with more difficulty than those united with NH.

I. Chloro-compounds ClR.Pt(NH,.NH,R)(NH,NH,R); the R's may be the same or different.

CHLOROPLATINIDIAMMINE CHLORIDE Cl₂Pt(NH₃,NH₂Cl)₂. By passing Cl into a fairly conc. boiling solution of platodiammine chloride, Pt(NH, NH,Cl), till the liquid begins to turn red (Raewsky, A. Ch. [3] 22, 278); also by dissolving chloroplatinammine chloride, Cl₂Pt(NH₂Cl)₂, in NH₃Aq (Gerhardt a. Laurent, C. R. 80, 273). Pale-yellow, crystalline powder; scarcely sol. cold water, sl. sol. boiling water. HNO,Aq produces the chloro-nitrate Cl₂Pt(NH₂.NH₃.NO₃)₂; a little AgNO₃Aq produces the chloro-hydroxyl-nitrate Cl(OH)Pt(NH, NH, NO₃)₂; long boiling with AgNO₂Aq is needed to remove all Cl as AgCl.

Reduced by H2S, or action of Ag, Pt(NH₂NH₃Cl)₂ (Thomsen, J. 1868. 278; Jörgensen, J. pr. [2] 33, 489). Forms double compounds with PtCl₂ and PtCl₄ (Reiset, A. Ch. [3] 11, 417; G. a. L., l.c.; Cleve, Bl. [2]

7, 12; 17, 482).

The other chloro- compounds which are known are the following: Chloro-nitrate, Cl.Pt(NH₃.NH₃.NO₃)₂ (Gros, A. Ch. [2] 69, 204; Raewsky, A. Ch. [3] 22, 278; Hadow, C. J. [2] 4, 345); chloro-chloro-nitrite, Cl.Pt(NH₃.NH₃.NO₂)(NH₃.NH₃Cl); chloro-hydroxyl-chloride, Cl(OH)Pt(NH₃.NH₃Cl)₂ (Cleve, l.c.; Hadow, l.c.); chloro-hydroxyl-carbonate, Cl(OH)Pt(NH₂.NH₃)₂OO₃.H₂O (Raewsky, l.c.); chloro-hydroxyl-chromats and dichromate, Cl(OH)Pt(NH₃.NH₃)₂CrO₄ and Cl(OH)Pt(NH₂.NH₃)₂Cr₂O₇ (Cleve, l.c.); chloro-hydroxyl-nitrate, Cl(OH)Pt(NH₂.NH₃.NO₃)₂, (Raewsky, l.c.; G. a. L., l.c.; Cleve, l.c.); chloro-chromate and dichromate, and chloro-sulphate, MCrO₄, MCr₂O₇, and MSO₄, where M = Cl₂Pt(NH₂,NH₃)₂ (Cleve, l.c.).

II. Bromo-compounds. The following bromo-bromide, MBr₂; bromo-dichromate, MCr₂O₇; members of this class have been isolated (Cleve, l.c.); (Cleve, l.c.); bromo-bromide, MBr₂; bromo-chloride, MCl₂; bromo-dichromate, MCr₂O₇; bromo - nitrate, M(NO₃)₂; bromo - phosphate, M(PO₄H₂)₂.2aq (in these formulæ $M = Br_2Pt(NH_3.NH_3)_2$); bromo-chloro-chloride, BrClPt(NH₃.NH₃Cl)₂; bromo-hydroxyl-chloride and nitrate, MCl₂ and M(NO₃)₂, where

 $M = Br(OH)Pt(NH_3.NH_3)_2$

III. Hydroxyl compounds. Hydroxylnitrate and sulphate, M(NO₃)₂ and MSO₄, where $\mathbf{M} = (\mathbf{OH})_2 \mathbf{Pt} (\mathbf{NH_3.NH_2})_2.$

IV. Iodo-compounds. Iodo-iodide, MI2; iodo-nitrate, $M(NO_s)_2$; iodo-sulphate, MSO_4 ; where $M = I_2Pt(NH_s, NH_s)_2$.

IV. Carbonato-compounds. Carbonatobromo - carbonate, carbonato - chloro - carbonate. and carbonato - nitrato - carbonate, Br. (CO.)R,

Cl₂(CO₃)R, and (NO₃)₂O₃R, where $R = 2[Pt(NH_4,NH_4)_2].(CO₃)_2$. V. Nitrato-compounds (Cleve, l.c.; Gerhardt a. Laurent, C. R. 30, 273): (NO₃)₂Pt(NH₃.NH₃Cl)₂.2aq ;

VI. Nitrito- compounds (Cleve, la.; Hadow, C. J. [2] 4, 345): (NO₂)₂Pt(NH₂,NH₂,NO₅)₂; I(NO₂)Pt(NH₂,NH₂,NO₅)₂.

VII. Sulphato-compounds (Cleve, l.c.): $(SO_4)Pt(NH_3.NH_3)_2SO_4.aq$; OH.Pt(NH, NH,)2Cl.2aq;

,so, OH(SO,)Pt(NH,NH,)2SO,3aq;

Br₂(SO₄)Pt(NH₃.NH₃), (SO₄) aq. Series 3. Platinisemidiammines, R₂Pt(NH₃.NH₃R)R. These compounds are isomeric with the platinammines, R₂Pt(NH₂R)₂. They are generally formed by oxidising the platosemidiammines, Pt(NH₃.NH₄R)R.

CHLOROPLATINISEMIDIAMMINE CHLORIDE Cl₂Pt(NH₃,NH₃Cl)Cl. By chlorinating platosemidiammine chloride, Pt(NH₃,NH₃Cl)Cl, by Cl or aqua regia (Cleve, l.c.). Orange, crystalline powder; S. 33 at 0°, 1.52 at 100°. Not acted on by conc. H2SO4 or alkali solutions.

The following salts of this series are known

(Cleve, l.c.):

bromo-bromide, Br.Pt(NH, NH, Br)Br; bromo-nitrite, Br₂Pt(NH₃.NH₃.NO₂)NO₂; chloro-nitrite, Cl2Pt(NH3.NH3.NO2)NO2; chloro-hydroxyl-nitro-nitrite, Cl(OH)Pt(NH3.NH3.NO2)NO2; trihydroxyl-nitrate, (OH),Pt(NH, NH, NO,)OH; hydroxyl-sulphate, (OH),Pt/NH, NH, NO,

Series 4. Platinimonodiammines, R,Pt(NHg,NHgR)(NHgR). Of this series the following have been isolated (Cleve, l.c.): bromonitrate, Br₂Pt(NH₃.NH₃.NO₃)(NH₃.NO₃); bromosulphate, Br_Pt(NH_3,NH_3,SO_,NH_3).aq; chloro-chloride, Cl_Pt(NH_3,NH_3Cl)(NH_2Cl); hydroxyl-nitrate, (OH)_Pt(NH_3,NH_3,NO_3)(NH_3,NO_3).aq; hydroxyl-bromo-nitrate,

(OH)BrPt(NH,.NH,.NO,)(NH,.NO,).aq; iodobromide, I2Pt(NH3.NH3Br)(NH3Br).

Series 5. Platinitriammines,

R,Pt(NH3.NH3.NH3R),. This constitution is probably to be given to salts formed by electrolysing ammonium carbamate or carbonate solution, using electrodes of Pt (Drechsel, J. pr. [2] 20, 378; 26, 277; Gerdes, J. pr. [2] 26, 257). The salt formed as described is probably carbonato-platinitriammine carbonate, (CO_s).Pt \(\bigcap of HNO, H,SO, and HCl, respectively, on this compound are obtained the nitratonitrate, sulphato-sulphate, and chloro-chloride, $(NO_s)_2$.M. $(NO_s)_2$, SO_4 .M. SO_4 .aq, and Cl_2 .M. Cl_2 , where $M = Pt(NH_2.NH_2.NH_3)_2$. Class III. Diplatinum compounds.

Pt(NH3.NH,R)

Series 1. Diplatodiammines, | Pt(NH, NH, R)

(Blomstrand, J. pr. [2] 3, 207). Cleve (l.c.) gives to these compounds the formula

 $Pt_2(NH_3)_2(NH_2)_2R_2$

The following members of this series are known: chloride, MCl, hydroxide, M(OH), nitrate, M(NO_s)₂; and sulphate, MSO₄; where $\mathbf{M} = \mathbf{Pt}_2(\mathbf{NH}_3)_4.$

Diplatosindiammines, Series 2.

RPt(NH,NH,R)

The only compound of this $\begin{array}{l} \mathrm{RPt}(\mathrm{NH_2N}) \\ \mathrm{series} \ \mathrm{is} \ \mathrm{the} \ chloride \ \mathrm{Cl_2Pt_2(\mathrm{NH_2)_4Cl_2}} \\ \mathrm{RPt}(\mathrm{NH_2R})_2 \end{array}$ RPt(NH, NH,R)

Series 3. Diplatinammines, RPt(NH,R), The only compound of this series is the iodide I₂Pt₂(NH₃)₄I₄ (Cleve, l.c.); by treatment with HIAq, the compound Pt₄I₁₀(NH₃)₈ and $Pt_sI_{1s}(NH_s)_{16}$ are produced.

Series 4. Diplatinidiammines,

RPt(NH,NH,R)2

(Cleve, l.c.). The iodo-nitrate, RPt(NH3.NH3R)2 I₂Pt₂(NH₂)₈(NO₃)₄, is formed by reacting on I,Pt(NH₃)4(NO₃)2 with NH,Aq and treating the product with HNO,Aq; most of the other compounds of the series are formed from the iodonitrate. Various classes of compounds of this series are formed by varying the composition of the radicles, R; the nomenclature is similar to that used for the platinidiammines. The compounds which have been isolated are bromonitrate, bromo-anhydronitrate, and bromosulphate, $M(NO_s)_4$, $M(NO_s)_4$, and $M(SO_s)_2$ -2aq, where $M = Br_2Pt_2(NH_s)_s$; hydroxyl-chloride, dichromate, phosphate, and sulphate, MCl, aq, $M(Cr_2O_7)_2$, $M(\hat{H}PO_4)_2$, and $M(\hat{S}O_4)_2$.2aq, where $M=(OH)_2Pt_2(NH_3)_3$; iodo-iodide, anhydro-iodide, nitrate, anhydronitrate, phosphate, and sulphate, MI, MI,O, M(NO₃),4aq, M(NO₃),2O, M(HPO₄),2 and M(SO₄),3 where M = I₂Pt₂(NH₃)₈; nitrato-nitrate, (NO₃)₂Pt₂(NH₃)₈(NO₃),4aq.

PLATINUMS, FULMINATING. Various

compounds of Pt, N, Cl, O, and H, all of which explode when heated, are formed by the action of KOHAq on (NH₄) PtCl₆, or by ppg. Pt(SO₄), with NH, Aq and boiling the pp. with KOHAq (Proust, Gehlen's J. 1, 348; Döbereiner, G. A. 72, 194; Fourcroy a. Vauquelin, Gehlen's J. 1, 348; Davy, S. 19, 91). These compounds are classed together as fulminating platinums. E. von Meyer (J. pr. [2] 18, 305) classifies according to the quantities of Cl they contain; all contain Pt and N in the ratio of equal numbers of atoms: tetrachloro-compound Pt,N,Cl,O₁,H₂, trichloroxy-Pt,N,Cl,O(H)O₁,H₂, dichloro-Pt,N,Cl,O₁,H₂, chloroxy-Pt,N,Cl,O(H)O₁,H₂. Tetrachloro-fulchloroxy- Pt.N.Cl(OH)O12H22. Tetrachloro-ful-minating platinum is formed by adding KOHAq drop by drop to Am, PtCl, in the ratio 4.6KOH: Am, PtCl, and washing the pp. by repeatedly boiling with acetic acid solution and then with water. At 150° 4H₂O is given off; explosion occurs at a higher temperature. Dilute H₂SO₄Aq produces H₁C₂O₄ and evolves CO₂. By treatment with KOHAq, or NH₂Aq, Cl, is easily removed, a third Cl is removed with difficulty, but the fourth Cl remains unattacked. M. M. P. M.

PLUMBAGO. A name sometimes given to graphite; v. Carbon, vol. i. p. 686.

PLUMBATES. Salts wherein PbO2 acts as acidic radicle, v. vol. iii. p. 132.

PLUMBITES. Salts wherein PbO acts as

acidic radicle, v. vol. iii. p. 129.

PLUMERIC ACID C₁₀H₁₀O₄. [139]. Occurs as calcium salt in the milky juice of *Plumeria* acutifolia (Oudemans, A. 181, 154). crystals (from water), v. sol. alcohol and ether, sl. sol. cold water. Yields salicylic acid on fusion with KOH. Reduced by sodium amalgam to crystalline 'hydroplumeric' acid C10H12O3.-K,A'' 3aq. — Ca(H,A''), 4aq. S. 5 at 20°. — CaH,A'' 5aq. S. 25 at 20°. — Ca,(HA''), 8aq. — Ag, H, A'v aq. — Ag, HA'v 1 aq: needles (from water).

PODOCARPIC ACID C17H22O, i.e.

 $C_bH_{1s}.C_aH_2Me(OH).CO_2H$. Mol. w. 274. [188°]. [a]_D = 136°. Constitutes at least 90 p.c. of the resin found by de Vrij in the trunk of an old tree, *Podocarpus cupressina*, var. imbricata (Oudemans, Å. 170, 213). Plates (from dilute alcohol), insol. water, v. sl. sol. benzene, v. e. sol. alcohol and ether. Dextrorotatory. Resolved at 300° into water and anhydride $C_{34}H_{42}O_s$. Yields methanthrene $C_{15}H_{12}$ on distillation with zinc-dust. The Ca salt on distillation yields p-cresol, hydrocarpol, carpene C_bH_{14} , and methanthrol $C_{15}H_{12}O_s$. Br added to its alcoholic solution forms $C_{17}H_{20}\text{BrEtO}_3(\text{EtOH})$ [above 80°], which gives off alcohol when heated for some time, leaving $C_{17}H_{20}\text{BrEtO}_3$ [158°].

AgA' 2 laq.

Methyl ether MeA'. [174°]. Grains. Ethyl ether EtA'. [146°]. Needles. Acetyl derivative C₁,H₂₁AcO₂. [152°]. Small needles (from dilute alcohol).

Nitropodocarpic acid C₁₇H₂₁(NÓ₂)O₃. [205°]. Formed, as well as the di-nitro-acids, by the action of HNO₃ (S.G. 1·34). Small crystals, insol. water, m. sol. hot alcohol. May be reduced to unstable amido-podocarpic acid, which forms

C1, H23NO3HCl ;aq.

Salts.—(NH₄)₂C₁,H₁₉NO₅4aq.— K₂C₁,H₁₉NO₅5¹₂aq. Red needles with green lustre, v. sol. water.—Na₂C₁,H₁₉NO₅9aq. Scarlet plates.—BaC₁,H₁₉NO₅7aq. Red needles.— BaC₁,H₁₉NO₅3aq.—BaA'₂4aq: yellow needles, sl. sol. water.—CaC₁,H₁₉NO₅4aq: orange needles.

Di-nitro-podocarpic acid $C_{17}H_{20}(NO_2)_2O_3$. [203°]. Pale-yellow crystals, insol. water, m. sol. alcohol. — Salts: $K_2C_{17}H_{18}N_2O_7$ 5aq. Crimson needles, with green reflex. —Ba $C_{17}H_{18}N_2O_7$ 4aq. —

 $Ag_2C_1, H_{18}N_2O_7$ 4aq: orange pp.

Sulpho-podocarpic acid $C_1, H_{21}(SO_3H)O_3$ 8aq. Got by warming with H_2SO_4 . Amorphous mass. Salts: Na₂A'' 7aq: easily soluble rosettes.— Ba(HA'')₂ 6aq. — BaA'' 8aq: white lamins. — CaA'' 7aq: thin lamins.

Hydrocarpol $C_{1a}H_{20}O$. (220° in vacuo). A product of the distillation of calcium podocarpate. Viscid oil, v. sol. alcohol and ether. Yields p-cresol, carpene, and methanthrol $C_{1a}H_{12}O$ [122°] when distilled. Methanthrol is crystalline, sol. KOHAq.

PODOPHYLLIN. The resin extracted from the root of Podophyllum pellatum (Guareschi, B. 12, 683; G. 10, 16; Busch, Ph. [3] 8, 443; Senier a. Lowe, Ph. [3] 8, 445). It contains bitter and physiologically active 'picropodophyllin' crystallising in needles [200°-210°], and several resins, e.g. 'podophyllo-quercetin' [247°-250°] and 'podophyllic acid' (Podwissotzky, Ph. [3] 12, 1011).

POLYCHROÏTE v. SAFFRON.

POLYMERISM; a term generally applied to the phenomena of the existence of more than one substance having the same composition, and related in properties, but with different molecular weights; v. Isomerism, vol. iii. pp. 79-81.

M. M. P. M.

POLYPORIC ACID C₁₈H₁₄O₄. [above 300°]. Occurs in a fungus belonging to the genus Polyporus, which grows on the bark of diseased or dead oaks. The fungus is yellow, but is turned violet by ammonia (Stahlschmidt, A. 187, 177; 195, 365). Bronzed tables (from alcohol), insol. water, ether, benzene, CS₂, and HOAc, sl. sol. chloroform and alcohol. Alkalis form purple solutions, decolourised by zinc-dust. Yields a tetraliydride when boiled with conc. KOHAq. Yields benzene on distillation with zinc-dust. Conc. HNO₃ forms C₁₈H₁₂(NO₂)₂O₄ [230°]. KClO₃ and HCl give C₁₈H₁₄Cl₄O₄ [108°] and C₁₈H₁₂Cl₄O₂ [110°].

Salts.—K₂A" 2aq: purple monoclinic crystals.—Na₂A" 2aq: violet needles.—(NH₄)₂A" 2aq.—BaA" 4aq.—BaA" 2aq.—SrA" 4aq.—SrA" aq.—CaA" 3aq: pale-violet needles.—MgA" 3aq.—

Ag A": insoluble pp.

Methylether Me₂A". [187°]. Ethylether Et₂A". [134°]. Needles.

Ethyl ether $\text{Et}_2\text{A}''$. [134°]. Needles. Diacetyl derivative $C_{18}\text{II}_{12}\text{Ac}_2\text{O}_4$. [205°].

Yellow needles, sl. sol. alcohol.

Hydride C₁₈H₁₈O₄. [163°]. Formed, together with the acid C₂₀H₁₈O₂ [156°] which yields Ag₂A", by boiling polyporic acid with conc. KOHAq. White crystalline powder (from hot water), sol. alcohol.—Na₁A''4aq.—BaA''.—MnA''6aq.—Ag₂A'': crystalline pp.

POPPY OIL. The oil expressed from the

POPPY OIL. The oil expressed from the seeds of Papaver somniferum contains glycerides of linoleic, stearic, and palmitte acids (Oudemans, J. 1858. 304; 1863. 333; cf. Mulder, J.

1865. 323)

POPULIN is the Benzoyl derivative of Salicin. PORPHYRINE C₂₁H₂₂N₃O₂. [97°]. Occurs, together with alstonine and alstonidine, in the bark of Alstonia constricta (Hesse, A. 205, 366). White amorphous substance, sol. alcohol, ether, and chloroform. Conc. H₂SO₄ gives a purple solution. Its acid solutions exhibit blue fluorescence.—B'₂H₂PtCl₆4aq.

POTASH. Potassium hydroxide, q. v., p.

POTASSIUM. K. At. w. 39.04. Mol. w. probably 39 04 (Ramsay, C. J. 55, 521; v. infra). M.P. 58° (Quincke, P. 135, 642); 62 5° (Bunsen, A. 125, 368). B.P. 719°-731° (Carnelley a. Williams, C. J. 35, 563); 667° (Perman, C. J. 55, 326). S.G. 865 at 15° (Gay-Lussac a. Thénard); ·875 at 13° (Baumhauer, B. 6, 655). V.D. not known with certainty; v. infra, molecular weight. S.H. from -78° to +10°·166 (Regnault, A. Ch. [3] 26, 286). C.E. (linear, 0°-50°) ·00008415 (Hagen, Verhandl. d. physikal. Ges. zu Berlin, Volume at $t^{\circ} = \text{vol.}$ at 0° 1882. No. 13). $(1 + \cdot 00023935t + \cdot 00000020925t^2)$ for interval 10° to 95° (Hagen, l.c.). E.C. at 0° (Hg at 0° = 1) 11.23; at 100° (liquid) 5.586 (Matthiessen, P. 100, 177). Refraction-equiv 7.51 (Kanonnikoff, J. R. 1884 (1), 119). The emission-spectrum is characterised by a line K, in the extreme red, and a line Ks in the violet; for measurements of all the chief lines v. B. A. 1884. 436; for the absorption-spectrum v. Roscoe a. Schuster, C. N. 29, 268. H.C. $[K^2,O] = 66,050$ (Woods, P. M. [4] 2, 268).

Occurrence. — Compounds of K are widely distributed in large quantities. Felspar, mica, &c., contain silicates of this metal; carnallite, sylvine, &c., contain KCl; alum-stone and other

minerals contain K₂SO₄; saltpetre is chiefly KNO₂. Sea-water contains c. 5 to 7 g. KCl per litre. Potassium salts, e.g. acid tartrate and oxalate, are found in plants. Animals contain salts of K, chiefly phosphate and chloride.

Historical.—Potash was decomposed by Davy (T. 1808. [1] 5) in 1807, by passing the electric current from a Volta-pile of 200 plates, through a piece of potash placed in a Pt basin Gay-Lussac a. Thénard, in 1808, reduced potash and obtained the metal, by heating it to whiteness with iron filings (A. Ch. 65, 325). In the same year Curaudan (A. Ch. 66, 97) showed that charcoal could be used instead of iron. Brunner (S. 38, 517) made this process applicable on the large scale, and B.'s method was modified by several workers, especially by Donny a. Mareska (A. Ch. [3] 35, 147).

Formation.-1. By electrolysing KOH. The most effective method is to place a little conc. KOHAq, with pieces of solid KOH in it, in a Pt basin, to connect this with the negative pole of a powerful battery, to pour a little Hg into the basin, and place therein the wire from the positive pole. The K which is thus formed amalgamates with the Hg; the amalgam, after drying, is heated in a small retort, along with a little mineral oil, which drives out the air, and the K remains when the Hg has been distilled off.—2. By electrolysing a molten mixture of KCl and CaCl₂ in the ratio 2KCl:CaCl₂. Temperature is arranged so that a solid crust remains on the surface of the molten mixture. Electrodes of gas-coke are used. After electrolysis the mixture is allowed to cool for twenty minutes, and the contents of the crucible are scraped out under petroleum (Matthiessen). Linnemann electrolyses molten KCN (J. pr. 73, 415).—3. By heating KOH to redness with iron filings or charcoal. - 4. By heating an intimate mixture of K₂CO₂ and C, or a mixture of K₂CO₃ or KOH, C, and very finely-divided iron (Castner, C. N. 54, 218).-5. By heating sulphide of K with iron filings (Dolbear, C. N. 26, 33).—6. By heating potash with sodium under petroleum to c. 170° (Williams, C. N. 3, 21).—7. By the action of Na on dry molten K acetate (Wanklyn, C. N. 3, 66).

Preparation.—An intimate mixture of K₂CO₃ and C is formed by heating cream of tartar (K-H tartrate) in a closed crucible. The mixture is then strongly heated in an iron bottle, connected with a flat receiver about 30 centim. long, 12 centims. wide, and 6 centims. from the inner surface of one side to that of the other; this receiver is made of two pieces of sheet-iron (about 4 mm. thick) screwed together, and has an opening at the end farthest from the retort, to allow the escape of CO. The reaction may be represented thus: $K_2CO_3 + 2C = 2K + 3CO$. The K distils over and condenses in the receiver; when full the receiver is removed, plunged under mineral oil and there opened, and the K is removed by a chisel. K combines with CO at a red heat to form a black, very explosive substance; the more rapidly the distilled K is cooled the less of this explosive compound is formed. The distilled metal is purified by wrapping it, when under oil, in a piece of linen, heating the oil to c. 65°, and pressing the molten K through the linen; the metal is then redistilled, from an iron bottle, into a copper tube

partly filled with mineral naphtha, and having an opening through which an iron rod may be passed to clear the tube which carries the K from the bottle into the receiver (Donny a. Mareska, A. Ch. [3] 35, 147). Castner (C. N. 54, 218) uses as reducer a mixture of C and a metallic carbide, or a mixture of very finely-divided metal and C (conveniently prepared by heating a mixture of Fe_2O_3 and tar in a covered vessel), and heats in an iron crucible with an exit-tube passing through the lid. The reaction, using KOH, Fe, and C, may be formulated as $3KOH + Fe + 2C = 3K + Fe + CO + CO_2 + 3H$.

Properties .- A silver-white, lustrous metal. As soft as wax at ordinary temperatures, brittle at 0°, melts at c. 60°. Boils when heated in a stream of H to c. 700°, and forms a brightgreen vapour. A sublimate, in a very thin film on glass, shows a rich purple colour in transmitted light (Dudley, C. N. 66, 163; Newth, N. 47, 55). Crystallises in quadratic octahedra, which have a greenish-blue sheen. These crystals are obtained by melting K in a glass tube, narrowed at one place, and filled with coal-gas, and allowing the semi-solidified metal to flow slowly through the constricted part of the tube (Long, C. J. 13, 122). Sol. liquid NH3, forming a blue liquid, from which the metal is obtained by allowing the NH, to evaporate (Seeley, C. N. 23, 169; v. Potassammonium, p. 299). Combines very rapidly with O; on exposing a freshly-cut surface to air, oxidation occurs at once. Decomposes cold water rapidly. K must be kept under a liquid free from O, such as mineral naphtha or rock-oil.

The atomic weight of K was determined by Berzelius, Penny, Marignac, Pelouze, Millon, Faget, and Maumené, from 1813 to 1846, by reducing KClO₂, KClO₄, and KIO₃ to KCl and KI, and by converting KCl, KBr, and KI to AgCl, AgBr, and AgI. The researches of Stas, in 1860 and 1865, have determined the at. w. with great accuracy. Stas heated KClO₃, thus getting KCl, decomposed KClO₃ by HClAq, determined the ratio of KCl and KBr to Ag required for complete ppn., converted KCl into KNO₃ and KBr into AgBr, and determined the ratio of KBr to AgNO₃ required for ppn. (Stas, Rech. 69, 70, 91, 118; Nouv. R. 244, 803).

The molecular weight of K is not known with certainty. Dewar a. Dittmar (C. N. 27, 121) and Dewar a. Scott (Pr. 29, 206, 490) determined the V.D. of K at c. 1000°; the earlier results pointed to a mol. w. of c. 90, and the later numbers to a value c. 45. V. Meyer (B. 13, 391) showed that the results were untrustworthy, as vapour of K attacks glass vessels. It has also been shown that vessels of Cu, Ag, Pt, or Fe cannot be used (Rieth, B. 4, 807; Meyer, l.c.). Ramsay (C. J. 55, 521) attempted to find mol. w. of K by dissolving in Hg, and finding the decrease of the vapour-pressure of Hg thereby produced. Assuming the mol. w. of liquid Hg to be 200, and also assuming that equal vols. of dilute solutions contain equal numbers of molecules, the results gave mol. w. of K as 29.1 and 30.2; it seems probable that the mol. of K is monatomic.

The only compound of K whose V.D. has been determined is KI; in this molecule the atom of K is monovalent.

K is a very strongly positive element. In all

its compounds it reacts as a metal, and does not enter into the composition of any negative radicles. K forms very few basic salts, and no oxyhaloid compounds; its halide compounds are very stable towards heat. K is closely related to Cs, Li, Na, and Rb (v. Alkalis, Metals of the, vol. i. p. 114).

Reactions.—1. Heated in dry air or oxygen, K₂O and K₂O, are formed (v. Oxides, p. 304).— 2. At about 300° K absorbs hydrogen readily, probably forming K.H (v. Hydride, p. 301). 3. Combines with chlorine, bromine, and iodine when heated (v. Chlorides, bromides, iodides, pp. 299, 300, 302).-4. Forms compounds with sulphur, selenion, and tellurium when heated (v. Sulphides, selenides, tellurides, pp. 305, 306).-5. Combines with phosphorus (v. Phosphide, p. 305).—6. Alloys with several metals (v. Alloys, infra). -7. Reacts rapidly with water, forming KOHAq and H.—8. Heated with hydrogen sulphide forms KHS (v. Hydrosulphide, p. 302).-9. Dissolves in liquid ammonia (Seeley, C. N. 23, 169; v. infra, Potassammonium). Heated in dry ammonia gas, KNH, is formed (v. Amide, infra). - 10. Combines with carbon monoxide to form KCO (v. Carbonyl compound, p. 300). At red heat decomposes CO with separation of C .-11. Decomposes carbon dioxide, and nitrous and nitric oxide, when strongly heated with these compounds.—12. Reacts with moist carbon dioxids to form K₂CO₂ and HCO₂K (Kolbe a. Schmitt, A. 119, 251).—13. Decomposes silica and boric acid, when heated with these compounds, probably forming silicide and boride (q. v., pp. 299, 305).

Potassium, alloys of. Alloys of K with most metals are known; K alloys easily with the more fusible metals. The alloys generally oxidise in air, and decompose cold water. K forms amalgams with Hg; according to Joannis (C. R. 113, 795), Hg₁₈K is formed by letting potassammonium (q. v.) in liquid NH₂ drop on to Hg. For polarisation of K amalgam in KClAq against Zn amalgam v. Le Blanc, Z. P. C. 5, 467. For supposed alloy with NH₄, v. infra, Potassammonium.

Potassium, aluminate of; v. vol. i. p. 141.

Potassium amide, KNH₂. Gay-Lussac a. Thénard (G. A. 29, 135; 32, 23) obtained dark olive-green crystals by heating K in dry NH₂. Baumert a. Landolt (A. 111, 1) confirmed G. a. T.'s formula KNH₂. The compound is formed by passing dry NH₂ into a flask of c. 50 c.c. capacity, fitted with a cork carrying an inlet and exit tube, quickly throwing in pieces of K, continuing the passage of NH₃, and heating very gently till the metal just ceases to dissolve in the greenish-blue liquid that is formed; on cooling, a yellowish-brown solid is obtained. KNH₂ is a non-conductor of electricity. Melts at a little over 100°, at a higher temperature NH₃ is given off, along with H and N, and K,N remains (v. Potassium nitrhide, p. 304). Burns when heated in O, or heated to redness in air, giving N and KOH; must, therefore, be kept under rock oil. Reacts energetically with water, giving NH₄ and KOHAq.

Potassium-ammonium or Potassammonium. 110°. Gerlach (Fr. 8, 285) By evolving dry NH₂, from AgCl.xNH₂, in a lowing from Kremer's data:

Faraday tube containing K in the other end. Weyl (P. 121, 697) obtained a substance which he regarded as an alloy, or compound, of K and NH₄. Seeley (C. N. 23, 169) regarded the blue liquid formed by dissolving K in liquid NH₂ as a solution, inasmuch as K was obtained again when the NH₃ was allowed to evaporate. Joannis (C. R. 109, 900, 965; 110, 238) examined the vapour-pressures of a solution of K in liquid NH₃, and also the thermal phenomena attending the formation of the solution. He found the vapour-pressure to decrease till a certain value was reached, when it became constant, provided temperature was not changed; on removal of more NH₃ a copper-red solid separated, and thereafter NH, was given off at constant pressure, till only K remained. This last part of the process was regarded by J. as a dissociation of NH_sK; analysis showed this ratio (NH_s:K) to be always attained when the final part of the change commenced. The heat of formation of NH₃K is given by J. as 6,300 when NH₃ is gas and K solid, and as 1,900 when NH, is liquid and K solid. Joannis (C. R. 113, 795) found that NH, K in liquid NH, is decomposed by Sb, Pb, and Hg, but not by Al, Cu, Ag, or Zn; by dropping the liquid on to Hg, and washing the product with liquid NH₂, he obtained a crystalline amalgam Hg₁₈K.

By measuring the depression of the vapourpressure of liquid NH, by solution of K therein, J. concludes that the mol. formula of potassammonium is $N_2H_6K_2$ (C. R. 115, 820).

Potassium, antimonide of. Probably SbK₃. A greyish-white, lustrous solid; decomposes water, giving KOH, Sb, and H. Formed by heating the elements together, or by fusing Sb₂O₃ with K₂CO₃ and C.

Potassium, antimonates of; v. vol. i. p. 286. Potassium, arsenates of; v. vol. i. p. 309.

Potassium, arsenites of; v. vol. i. p. 306.

Potassium, auricyanide of; v. vol. ii. p. 332.

Potassium, borates of; v. vol. ii. p. 332. Potassium, borates of; v. vol. i. p. 529.

Potassium, boride of. The brown solid formed when boric acid is reduced by heating K may contain a compound of B and K.

Potassium, borofluoride of; v. vol. i. p. 526.

Potassium, boronitride of. By very strongly heating a mixture of 7 parts B₂O₂ with 20 parts KCN, Balmain (J. pr. 27, 422) obtained a white, infusible solid; insol. water or KOHAq; decomposed by boiling aqua regia, leaving BN (vol. i. p. 527); heated in steam gave off NH₂, and left KOH and boric acid.

Potassium, bromide of, KBr. Formula probably molecular, from analogy of KI. Melts at 699° (Carnelley, C. J. 33, 279). S.G. 2-695 to 2-72 at 4° (Schröder, P. 106, 226); 2-712 at 12-7° (Clarke's Tables of Spec. Gravities, 2nd ed., 31); 2-199 fused (Quincke, P. 138, 141); v. also Spring (B. 16, 2724). S.H. (16° to 98°)·11322 (Regnault, A. Ch. [8] 1, 129). Vol. at 40° = vol. at 0° (1+00012602) (Fizeau, C. R. 64, 314). S. 53-48 at 0°, 64-6 at 20°, 74-62 at 40°, 84-74 at 60°, 93-46 at 80°, 102-04 at 100° (Kremers, P. 25, 119); Coppet (A. Ch. [5] 30, 411) gives S. 54-43+-5122t where t varies from —13-4° to 110°. Gerlach (Fr. 8, 285) calculated the following from Kremer's data:

	S.G. 1	KBrAq	
5 p.c.	1.037	30 p.c.	1.256
10 ,	1.075	35 ,,	1.309
15 ,,	1.116	40 ,	1.366
20 ,,	1.159	45 ,,	1.430
25 ,,	1.207		
	TD1 05 910 .	TT7 TD A 7 0.	0 0 2 A CO A

H.F. [K,Br] = 95,310; [K,Br,Aq] = 90,230 (Th.

8, 235). [KBr,O'] = 84,060 (Th., l.c.).

Formation. -- 1. K and Br combine with production of much heat (v. Balard, Merz, a. Weith, B. 6, 1518).—2. By treating BrAq with iron filings and ppg. with K₂CO₃. -3. By neutralising HBrAq by KOHAq or K₂CO₃Aq, and evaporating.-4. By decomposing CaBr₂Aq by K₂SO₄, filtering after 12 hours, adding K₂CO₃ as long as turbidity is produced, filtering, and evaporating (Klein, A. 128, 237).

Preparation.-A moderately conc. solution of KOH, free from K2CO3, is prepared from pure K2CO3 and pure CaO (v. Potassium hydroxide, p. 302); to this solution Br is added, little by tenth of the quantity of Br used, is added, and the liquid is evaporated to dryness (6KOHAq + 3Br₂ = 5KBrAq + KBrO₃Aq + 3H₂O); the dry residue is well powdered, and heated to dull redness in a closed crucible for some time $(2KBrO_3 + 3C = 2KBr + 3CO_2)$; the residue is extracted with warm water, the solution is filtered, and evaporated to the crystallising point.

Properties and Reactions.—KBr crystallises in very lustrous, white cubes, sometimes elongated to prisms or flattened to plates; it has a strongly saline taste. Easily sol. water, with considerable lowering of temperature; [KBr,Aq] =-5,080 (Th. 3, 235). Chlorine reacts with KBrAq to give KClAq and Br; Br is not set free by H₂SO₄ containing N₂O₃, nor by KNO₂Aq + dil. H₂SO₄Aq. KBr heated with potassium chromate and sulphuric acid gives off Br: KCl under similar conditions gives CrO, Cl2. Conc. sulphuric acid produces HBr, Br, and SO2; the amount of HBr (the primary product of the reaction) thus decomposed varies very nearly in proportion to the quantity of H.SO. used; if the H2SO4 is in such large excess that the water formed in the reaction causes no sensible dilution, only SO2 and Br are produced; by using dil. H₂SO₄Aq (c. 30 p.c.) a mere trace of Br is obtained, the products being almost wholly KHSO, Aq and HBrAq (v. Addyman, C. J. 61, Hypochlorous acid solution produces KClAq and KBrOsAq, with evolution of Br and Cl. Potassium permanganate solution has no action even when boiled; but Br is set free if a little H₂SO₄ is added, even without warming, the decomposition being soon completed (Hempel, A. 107, 160). Fusion with potassium chlorate produces KBrO.

Combinations.—With many bromides of less positive metals to form double salts; some of these are best regarded as K salts of acids containing Br and a metal less positive than K, e.g. AuBr. KBr is best looked on as KAuBr. (v. the various metallic bromides). Also with iodine bromide, to form KBr.IBr (Wells a. Wheeler, Am. S. [3] 43, 475). Also with some metallic chlorides; thus SbCl₂ forms SbCl₂ 3KBr identical with SbBr, 3KCl, and therefore to be regarded as SbK, Cl.Br. (Atkinson, C. J. 43, 290). Feit (J. pr. [2] 39, 373) describes MgBr₂.KBr.6aq, analogous to carnallite (MgCl2.KCl.6aq). Schiff (A. 228, 72) describes a compound with arsenious

oxide KBr. As O.

Potassium tribromide KBrs. According to Berthelot (A. Ch [5] 21, 370), orange crystals of this composition are formed by adding Br to KBrAq; no analyses are given. The substance

decomposes rapidly. B. gives [KBr,Br²] = 2940.

Potassium, carbonyl compound of. (Carbonic oxide potassium.) The black solid formed in the preparation of K was thought by Berzelius (P. 4, 31) to be a carbide; Liebig (A. 11, 182) found that the same compound was formed by passing CO over K heated just to melting; Brodie (C. J. 12, 269) confirmed Liebig's result, and determined that one molecule CO is absorbed for each atom K; hence the empirical formula is probably KCO. This substance is extremely explosive; many serious accidents have occurred with it. If the CO used is quite dry, and the substance is washed rapidly in alcohol as soon as it is prepared, the residue is not nearly so explosive (Nietzki a. Benckiser, B. 18, 1833, where details of the method of preparation are given); probably the alcohol dissolves out unchanged K. Potassium carbonyl is a greyish solid; on standing in air it becomes yellowish and very explosive; it dissolves in water, with evolution of much gas (according to E. Davy [A. 23, 144] this gas contains C₂H₂), and generally with combustion or explosion; heated to redness it is resolved into K and CO. By exposure to moist air, or by other processes of oxidation, a series of organic compounds is obtained (v. CROCONIC ACID, vol. ii. p. 275; HEXA-OXY-BENZENE, vol. iii. p. 678; Tetra-oxy-quinone, vol. iii. p. 771; RHODIZONIC ACID, this vol.).

Potassium, chloride of, KCl. Formula probably molecular, from analogy of KI. Melts at 734° (Carnelley, C. J. 33, 279). S.G. 1 9775 at 4° (Playfair a. Joule, C. S. Mem. 2, 401); 1 9453 at 15° (Stolba, J. pr. 97, 503); 1.612 at M.P. (Braun, C. J. [2] 13, 31); 1.87 fused (Quincke, (Brain, C. v. [2] 10, 21, 10 fraction (California), 12, 135, 642); v. also Spring (B. 16, 2724). S.H. 14° to 99° 17295 (Regnault, A. Ch. [3] 1, 129; v. also Kopp, T. 155 [1] 71). Vol. at 40° = vol. at 0° (1 + 00011408) (Fizeau, C. R. 64, 314). S. 32 at 10°, 33·4 at 15°, 34·7 at 20°, 37·4 at 30°, 40·1 at 40°, 42·8 at 50°, 45·5 at 60°, 48·3 at 70°, 51 at 80°, 53·8 at 90°, 56·6 at 100° (Mulder, Scheikund. Verh. 1864. 39); Coppet (A. Ch. [5] 30, 411) gives S. 28·51 + ·2837t°, where to varies from -11° to 109°. S. at 0° in alcohol is given in following table (Gerardin, A. Ch. [4] 5, 139); where S.G. of alcohol is at 0° , and $S+t^{\circ}$ is solubility at to (cf. Schiff, A. 126, 167):

Alcohol S.G.	8.	to	Alcohol S.G.	s.	to
.9904	$23 \cdot 2$.27	9573	7.1	·162
.9848	19.4	.255	•9390	4.2	.125
.9793	15.7	.233	8767	1.89	.061
.9726	11.9	.205			

Gerlach (Fr. 8, 281) gives the following data:

			B.G.	KCLAq			
1	per	cent.	1.0065	20	per	cent.	1.1361
5	٠,,	**	1·0065 1·0325	24	-,,	12	1.1657
10	"	,,	1.0658	24.9	**	99	1.1723
15	**	**	1.1004	ll .			

H.F. [K,Cl] = 105,610; [K,Cl,Aq] 101,170 (Th. 3,235). 30 parts KCl dissolved in 100 parts water at 13.2° lower the temperature to .6°, i.e. through 12.6° (Rüdorff, B. 2, 68). Solution of 1 g. KCl in 100 g. water freezes at -446° ; saturated solution freezes at -10.9° (Rüdorff, P. 114, 63; 122, 337). Saturated solution boils at 108°.

Occurrence. - In small quantities in sea-water, and in some mineral springs. As chloride in sylvine; as double chloride, especially as carnallite KCl.MgCl₂.6aq. In crude pearl ash, in plant ash, and in kelp.

Formation.—1. By the direct union of the elements; K burns in Cl at the ordinary temperature.—2. By heating K in HCl gas.—3. By passing Cl over KOH or KI heated to redness .-4. By the action of HClAq on KOH or K2CO3. 5. By decomposing various metallic chlorides by fusion with K.

Preparation.—1. Commercial KCl is prepared chiefly from carnallite. The mineral is dissolved in warm water, heated by steam to c. 120°, and allowed to cool; at 60°-70°, MgSO, aq, CaSO,, and NaCl separate, and on further cooling c. 70 p.c. of the KCl is obtained; the crystals of KCl are washed with a little cold water, to remove NaCl and MgCl₂, and a product containing c. 95 p.c. KCl is thus obtained (for details v. Dic-TIONARY OF APPLIED CHEMISTRY) .- 2. Pure KCl is prepared by neutralising pure HClAq by pure KOH or K₂CO₃, evaporating to the crystallising point, and recrystallising from water.

Properties.—White salt, crystallising in cubes.
Saline taste. Unchanged in air. Decrepitates when heated; melts at c. 740°, and volatilises at a higher temperature. Fairly sol. water or aqueous alcohol; insol. absolute alcohol, or conc.

HClAq; sol. in 20 p.c. KC₂H₂O₂Aq.

Reactions.—1. With acids, KCl generally gives K salt of the acid used and HCl.—2. Said to combine with sulphuric anhydride, and chromic anhydride, probably forming SO2.Cl.OK and CrO2.Cl.OK respectively (H. Rose, P. 38, 117).—8. Fused with potassium, in H, a blue substance is formed, supposed by H. Rose to be a subchloride (P. 120, 15). What is probably the same substance is formed during electrolysis of molten KCl (Bunsen a. Kirchoff, P. 113, 344). 4. For action of water and oxygen, in presence and absence of acids, v. Schulze, J. pr. [2] 21,

Combinations. — 1. With sulphuric and chromic anhydrides, v. Reactions, No. 2 .- 2. With most metallic chlorides; some of the compounds are best regarded as K salts of metal-containing acids, e.g. KAuCl, (v. the different metallic chlorides). — 8. With iodine trichloride, to form KCl.ICl₃; prepared by mixing KClAq and ICl₃Aq, by passing Cl into warm KIAq containing HCl, or by dissolving 1 part KIO, in 8 parts HClAq of S.G. 1.176, at 40°-50°. Forms lustrous yellow prisms, smelling of Cl; on heating gives Cl, KCl, and ICl; with water forms KIO₃; ether dissolves out ICl₃ (Filhol, J. Ph. 25, 435, 506).— 4. With iodine monochloride, to form KCl.ICl (Wells a. Wheeler, Am. S. [3] 43, 475).

Potassium, cyanide of : v. vol. ii. p. 346.

Potassium, chromicyanide of; chromisulphoeyanide of; and chromocyanide of; v. vol. ii. p.

Potassium, cobalti- and cobalto-cyanide of: v. vol. ii. p. 330.

Potassium, ferrate and ferrite of; v. vol. ii.

Potassium, ferri- and ferro-cyanide of; v. vol. ii. pp. 339, 336.

Potassium, fluorides of. Two fluorides are known, KF, and KHF2 or KF.HF.

POTASSIUM FLUORIDE KF. Formula probably molecular, from analogy of KI. Prepared by neutralising HFAq by K₂CO₂ in a dish of Ag or Pt, evaporating to dryness, and heating till HF ceases to be given off. According to Berzelius (P. 2, 218), KF is obtained in crystals by slowly evaporating an aqueous solution at 35°-40° in a very shallow dish. Deliquescent; fusible; solution has alkaline reaction to litmus, and etches glass. Guntz (A. Ch. [6] 3, 5) says KFAq is neutral, but the salt is decomposed in solution and becomes alkaline. Evaporation of a very conc. solution gives KF.2H.O, according to H. Rose. S.G. 2⁴54 (Bödeker); 2[.]096 at 21[.]5° (Clarke, Am. S. [3] 13, 291). Combines with HF to form KF.HF (v. infra); with BF₃ to form. KBF. (v. Potassium borofluoride, vol. i. p. 526); with SiF₄ to form K₂SiF₅ (v. p. 305); with B₂O₃ to form 2KF.B₂O₃, obtained by dissolving B₂O₄ in molten KF (Schiff, A. 228, 72); with TeF, to form KF.TeF₄, by evaporating ${\rm TeO_3}$ in ${\rm HFAq}$, with addition of K₂CO₃ (Högborn, Bl. [2] 35, 60); and with many metallic fluorides.

POTASSIUM-HYDROGEN FLUORIDE KHF, or KF.HF. Prepared by dividing a quantity of HFAq into two equal parts, neutralising one, adding the other, and evaporating; also by evaporating KF in acetic acid. White cubes or four-sided tables; e. sol. water, almost insol. dil. HFAq. When heated gives HF, leaving KF. By electrolysis of HF containing KHF2, F is obtained (v. Fluorine, vol. ii. p. 561). Moissan (C. R. 106, 547) says that two other compounds of KF and HF are obtained by dissolving dry KF in liquid HF, and cooling; to these compounds he gives the formulæ KF.2HF and KF.3HF. Guntz (A. Ch. [6] 3, 5) gives [HF,KF] = 21,100.

Potassium, haloid compounds of. The compounds KF, KCl, KBr, and KI are the chief halides of K; as KI has been gasified and the molecular weight corresponds with the simplest formula, it is probable that KX expresses the molecular composition of these halides. KI. also exists, but is decomposed by heat; and there are indications of the existence of KBr. KF combines with HF to form KHF2, and, according to Moissan, also KF.2HF and KF.3HF.

Potassium, hydride of, K.H. The absorption of H by heated K was observed by Gay-Lussac a. Thénard (A. Ch. 74, 203), and was confirmed by Jacquelain. Troost a. Hautefeuille A. Ch. [5] 2, 273) found that the absorption of H by K begins at c. 200° and becomes rapid at 350°-400°; if the action continues for some time 126 vols. H are absorbed by 1 vol. K. The product is brittle, and much resembles Ag amalgam; it is fusible without change in H or in vacuo; takes fire in contact with air; heated* in vacuo, dissociation begins at 200° (for vapourpressures of H given off, v. Dissociation, vol. ii, p. 398). The formula K₂H requires 124.6 vols-H to 1 vol. K.

Potassium, hydrosulphide of, KHS (Potassium sulphydrate). Gay-Lussac a. Thénard (A. Ch. 115, 165) obtained this compound by heating K in dry H₂S gas; it is more readily formed by heating K₂CO₃ to redness in H₂S (H₂O and CO₂ are given off) (Berzelius, P. 6, 438). By saturating KOHAq with H2S, and evaporating the conc. solution over CaO or CaCl2 in vacuo, Schöne (P. 131, 380) obtained lustrous, rhombohedral crystals of 2KHS.H,O; and by dehydrating this in a stream of dry H,S, Sabatier (A. Ch. [5] 22, 5) obtained KHS as a yellow, amorphous solid. Prepared in the dry way, KHS appears reddish-black when molten, and white when cold; it crystallises in prisms; is very deliquescent, and reacts strongly alkaline; easily sol. in alcohol. Thomsen (Th. 3, 235) gives [K,S,H,Aq] = 65,140. KHSAq gives off H_2S at 70°, according to Drechsel (J. pr. [2] 4, 20). When a current of an inert gas is passed through KHSAq, H,S is given off, and K,SAq remains (Gernez, C. R. 64, 606). Electrolysis produces H and KOH at the negative pole, and H_2S at the positive (Bunge, B. 3, 911). KHSAq dissolves S with evolution of H2S; it ppts. MnS, or PbS, from a neutral solution of a Mn or Pb salt, at the same time giving off H₂S. These reactions distinguish KHSAq from K₂SAq; the latter does not give off H₂S while dissolving S or ppg. MnS or PbS. KHSAq dissolves several sulphides of the less positive metals, e.g. Sb, As, Sn, forming K thiosalts.

Potassium, hydroxide of, KOH (Caustic potash. Potassium, or potassic, hydrate). S.G. 2·1 (Dalton); 2·044 (Filhol, A. Ch. [3] 21, 415). H.F. [K,O,H] = 103,170; [K,O,H,Aq] = 116,460

(Th. 8, 235).

Formation.—1. By the reaction between K and H₂O, or K₂O and H₂O, followed by evaporation.—2. By boiling K₂CO₃Aq with CaO.—3. By adding powdered K₂SO₄ to warm conc. BaOAq, evaporating, filtering, and again evaporating (Schubert, J. pr. 26, 117).—4. By heating KNO₅ with 2-3 parts thin copper turnings, to redness, in an iron, or copper, crucible, extracting with water when cold, filtering from CuO and Cu₂O, and evaporating (Wöhler, A. 87, 373).—5. By heating I part KNO₅ with I part Fe₂O₅, in a closed crucible of Cu, H being passed in; treating with water, and drawing off the clear liquid, and evaporating it (Schulze, Z. 1861. 109). Evaporation of KOHAq should be performed in vessels of polished iron, or, better, of silver.

Preparation.—1. A solution of pure K2CO, in 10-12 parts water is boiled, in a dish of silver or polished iron, with milk of lime, added little by little, till a portion of the clear supernatant liquid gives no effervescence with an acid; rather more than half as much CaO is required as the weight of K₂CO₃ taken; large excess of CaO should be .avoided; water should be added as the boiling proceeds, because if the quantity of water is less than c. 10 times the quantity of K,CO, used the KOHAq begins to decompose the CaCO, formed, and the change of K.CO. to KOH stops (Liebig, A. 1, 124). When the change is completed, the vessel is closed, and, after a few hours, the clear liquid is drawn off by a syphon; the liquid is then rapidly evaporated in an iron vessel, allowed so stand for an hour or two, the vessel being closed, syphoned off from any pp. that has formed,

evaporated in a silver dish till the oily liquid thus produced begins to volatilise in white clouds, and cooled in an exsiccator. Impure KOH may be freed from all impurities, except KCl and traces of K_2CO_3 and $KC_2H_3O_2$, by dissolving in absolute alcohol, allowing to settle, draining off, and evaporating, at first on a water-bath, in a silver dish; the resinous matter which is produced is removed from the warm evaporated semi-solid mass, by a silver spatula, and the KOH is then poured out on to a plate of polished iron, or, better, of silver.

Graeger (J. pr. 96, 188) recommends to heat the $K_2\text{CO}_3\text{Aq}$ used with Ag_2CO_3 , to filter from AgCl and excess of Ag_2CO_3 , to boil, in a silver dish, with pure CaO made by strongly heating pure CaCO_3 , to filter the solution of KOH through pounded marble which has been washed with water till free from very fine particles, and to evaporate in a silver dish.—2. Water and benzene are placed in a silver dish, and small pieces of K, cut from the inside of a lump, are thrown in one by one; the K remains near the surface of separation of the two liquids, and reacts fairly slowly with the water. The benzene is removed by warming, and the aqueous solution of KOH is then evaporated to dryness.

Properties.—After fusion, KOH is a white, hard, brittle solid, often showing a fibrous texture. Melts below red heat, and volatilises at full redness in white pungent vapour. Very deliquescent; sol. water with production of much heat, [KOH,Aq] = 13,290 (Th. 3, 235). Easily sol. alcohol. Skey (C. N. 36, 48) says KOH is very sl. sol. ether. Absorbs CO₂ rapidly from the air, forming KHCO₃. KOH has a slight, but nauseous, odour, a strongly acrid taste, and acts as a powerful cautery towards both animal and vegetable matter. KOHAq should be kept in glass vessels free from Pb, as it corrodes leadglass; it attacks vessels of glass or porcelain when heated in them. To keep KOHAq free from carbonate, it is advisable to store it in bottles fitted with corks carrying a syphon-tube, and a rather wide tube filled with a mixture c. equal parts CaO and Na₂SO₄, well rubbed together, dried over a flame, and passed through a sieve to remove fine powder; the air which enters through this mixture is quite free from

KOHAq is strongly alkaline; the affinity of KOH in solution is large, about equal to that of NaOH and LiOH, and about 50 times greater than NH,Aq (Ostwald). KOH saponifies ethereal salts, and ppts. most of the heavier metals as oxides or hydroxides from solutions of their

The table on next page, given by Lunge, shows the composition of KOHAq of different S.G.

Expansion occurs when conc. KOHAq is diluted; Frankenheim (J.~1847-8.~69) gives $V=1+000415t+00000577t^{\circ}$ as representing the volume of KOHAq S.G. 1-2788 at 0°, between 13° and 100°.

Reactions. -1. According to Deville (C. R. 45, 857), KOH is decomposed to K, O, and H by heating to white heat.—2. Strongly heated with non-volatile acidic anhydrides, water and potassium salts are formed. 3. Heated with potassium, K,O and H are formed; heated with sodium under a liquid free from O, an alloy of K and Na

8.G. K OHAq	Baumé	Twaddell	100 pts. by weight contain		l cub. metre contains Kilos	
	m m		K ₂ O	кон	K,O	кон
1.007	1	1.4	0.7	0.9	7	9
1.014	2	2.8	1.4	1.7	14	17
1.022	3	4.4	2.2	2.6	22	26
1.029	4	5.8	2.9	3.5	30	36
1.037	5	7.4	3.8	4.5	39	46
1.045	6	9.0	4.7	5.6	49	58
1.052	7	10.4	5.4	6.4	57	67
1.060	8	12.0	6.2	7.4	66	78
1.067	9	13.4	6.9	8.2	74	88
1.075	10	15.0	7.7	9.2	83	99
1.083	11	16.6	8.5	10.1	29	109
1.091	12	18.2	9.2	10.9	100	119
1.100	13	20.0	10.1	12.0	111	132
1.108	14	21.6	10.8	12.9	119	143
1.116	15	23.2	11.6	13.8	129	153
1.125	16	25.0	12.4	14.8	140	167
1.134	17	26.8	13.2	15.7	150	178
1.142	18	28.4	13.9	16.5	159	188
1.152	19	30.4	14.8	17.6	170	203
1.162	20	32.4	15.6	18.6	181	216
1.171	21	34.2	16.4	19.5	192	228
1.180	22	36.0	17.2	20.5	203	242
1.190	23	38.0	18.0	21.4	214	255
1.200	24	40.0	18.8	22.4	226	269
1.210	25	42.0	19.6	23.3	237	282
1.220	26	44.0	20.3	24.2	248	295
1.231	27	46.2	21.1	25.1	260	309
1.241	28	48.2	21.9	26.1	272	324
1.252	29	50.4	22.7	27.0	284	338
1.263	30	52.6	23.5	28.0	297	353
1.274	81	54.8	24.2	28.9	308	368
1.285	32	57.0	25.0	29.8	321	385
1.297	83 84	59.4	25.8	30.7	335	398 416
1.308	35	61·6 64·0	26.7	41.8	349	432
1·320 1·332	36	66.4	27.5	32·7 33·7	363	449
1.345	37	69.0	28·3 29·3	34.9	377 394	469
1.357	38	71.4	30.2	35.9	410	487
1.370	39	74.0	31.0	36.9	425	506
1.383	40	76.6	31.8	37.8	440	522
1.397	41	79.4	32.7	88.9	457	543
1.410	42	82.0	33.5	39.9	472	563
1.424	43	84.8	34.4	40.9	490	582
1.438	44	87.6	35.4	42.1	509	605
1.453	45	90.6	36.5	43.4	530	631
1.468	46	93.6	37.5	44.6	549	655
1.483	47	96.6	38.5	45.8	571	679
1.498	48	99.6	39.6	47.1	593	706
1.514	49	102.8	40.6	48.3	615	731
1.530	50	106.0	41.5	49.4	635	756
1.546	51	109.2	42.5	50.6	655	779
1.563	52	112.6	43.6	51.9	681	811
1.580	53	116.0	44.7	53.2	706	840
1.597	54	119.4	45.8	54.5	731	870
1.615	55	123.0	47.0	55.9	759	905
1.634	56	126.8	48.3	57.5	789	940
!	!	!		1		

Is said to be formed (Williams, Rep. Chim. pure, 8,177).—4. Iron decomposes KOH at white heat, forming Fe₂O₂, H, and K.—5. Molten KOH acts generally as an oxidiser: e.g. Fe, As, Sb, Pt, &c. is decomposed form ferrate, arsenate, antimonate, and platinate of K; Cr₂O₂ forms K₂CrO₄, &c. Salts are generally decomposed by molten KOH, giving 121, 225).

K salts, and setting free the bases.—6. KOHAq neutralises acids, forming salts.—7. KOHAq decomposes most metallic salts in solution, ppg. oxides or hydroxides of the metals.—8. According to Schöne (A. 193, 241), addition of hydrogen peroxide to KOHAq produces K₂O₄ (v. Potassium tetroxide, p. 305).

Combinations.—1. With carbon dioxide, to form KHCO₁.—2. With water, to form hydrates. Pickering (C. J. 63, 850) obtained KOH.H₂O freezing at 143°, KOH.2H₂O freezing at 35·5°, and KOH.4H₂O freezing —32·7° (cf. Walter, P. 39, 192; Schöne, P. 131, 147). P. (l.c.) gives full data for freezing points of KOHAq. Göttilg (B. 20, 1094) described two hydrates, 2KOH.9H₂O and 2KOH.5H₂O, obtained from an alcoholic solution of KOH containing some water; but in a later paper (B. 20, 1907) G. says that these hydrates contained alcohol besides water. 3. With methyl alcohol, to form 3KOH.5MeOH (Göttig, B. 20, 1832).—4. With ethylic alcohol, to form KOH.2EtOH (Engel, C. R. 103, 155); decomposed by heating to EtOK, EtOH, and

Potassium, iodides of. Two iodides are known, KI and KI₃.

Potassium iodide, KI. Mol. w. 165.57. Melts at 634° (Carnelley, C. J. 33, 279). S.G. 3.059 (Playfair a. Joule, C. S. Mem. 2, 401); 3.077 to 3.081 (Schröder, P. 106, 226); fused 2.497 (Quincke, P. 138, 141); v. also Spring (B. 16, 2724). V.D. 91.5 (Dewar a. Scott, Pr. 29, 206); 84.6 at above 1300°, in N (Mensching a. Meyer, Z. P. C. 1, 157). S.H. (20° to 99°) 08191 (Regnault, A. Ch. [3] 1, 129). Vol. at 40° =vol. at 0° (1+00012796) (Fizeau, C. R. 64, 314). S. 132·1 at 5°, 136·1 at 10°, 140·2 at 15° 144.2 at 20°, 152.3 at 30°, 160 at 40°, 168 at 50°, 176 at 60°, 184 at 70°, 192 at 80°, 201 at 90° 209 at 100°, 218 at 110° (Mulder, Scheikund. Verhandel, Rotterdam, 1864.162). Coppet (A. Ch. [5] 30, 411) gives S. at $t^{\circ} = 126.23 + .8088t$, where t varies from -5.9° to 120°. S. in aqueous alcohol is given in following table (Gerardin, A. Ch. [4] 5, 139); the values hold for 0° to

S.G. alcohol	8.	[S.G. alcohol	8.
·9904	130.5	•9528	76 ·9
•9851	119.4	•9390	66.4
•9726	100.1	•9088	48.2
•9665	89.9	•8464	11.4
		•8399	6.2

H.F. $[K,\Gamma] = 80,130$; [K,I,Aq] = 75,020 (Th. 3, 235).

Formation.—1. By the direct union of K and I.—2. By neutralising HIAq by K₂CO₃, and evaporating.—3. I is added to water and iron filings till the iron is almost wholly dissolved, the solution is filtered, and K₂CO₃ added so long as FeCO₃ ppts.; the liquid is filtered (if alkaline it is neutralised by HIAq) and evaporated, any Fe₂O₃ which separates being filtered off (Baup, J. Ph. 9, 37, 122).—4. By adding I to KOHAq, evaporating, heating with C, dissolving, filtering, and evaporating (v. Preparation).—5. BaI₂Aq is formed by the action of I and water on BaS, and is decomposed by K₂SO₄; BaSO₄ is filtered off, and the liquid is evaporated. Liebig (A. 121, 222) used CaI₂ in place of BaI₂ (cf. Pettenkofer, A. 121, 225).

Preparation.—Moderately, but not too, conc. KOHAq, quite free from K2CO3, is prepared from K.CO.Aq and CaO (v. Potassium Hydroxide, Preparation, p. 302); I is added, little by little, to the slightly warm liquid till a slight yellow colour is produced; very finely powdered charcoal is now added, equal to c. $\frac{1}{10}$ of the weight of I used, the liquid is evaporated quite to dryness, the residue is powdered, and heated, in a closed crucible, to dull redness for some time $(6KOHAq + 6I = 5KIAq + KIO_9Aq + 3H_2O; KIO_3 + 3C = KI + 3CO)$. The contents of the crucible, when cold, are extracted with water, the liquid is filtered, neutralised by HIAq if alkaline, and crystallised. Morse a. Burton (Am. 10, 321) secommend to remove traces of KIO2 by boiling for some time with Zn amalgam and water, filtering, and crystallising; neither Zn nor Hg is found in the filtrate. The Zn amalgam is made by agitating zinc-dust with Hg in presence of tartaric acid solution, and washing with water.

Properties.—White cubes; non-deliquescent. Has a sharp taste. Solution in water is attended with fall of temperature; 140 pts. in 100 pts. water at 10.8° produce a fall to -11.7° (Rüdorff,

P. 136, 276).

Reactions.-1. Unchanged in dry air, but decomposed in ordinary air when exposed to sunlight (v. Downes a. Blunt, Pr. 29, 319; Loew, Fr. 1870. 251). Air containing ozone also decomposes KI (Houzeau, J. 1858. 60).—2. Heated in air to 230°, some KIO₃ is formed.—3. Heated in steam, I is evolved (Petterson, Fr. 1870. 362) and HIAq formed (Schindler, Mag. Pharm. 31, 33).-4. Decomposed, with separation of I, and formation of KOH, by heating with lead dioxide, manganese dioxide, arsenic oxide, antimonic oxide, chromium trioxide, hydrogen peroxide, &c. (v. Weltzien, A. 138, 134).—5. Chlorine decomposes hot KI to KCl and I. KIAq with ClAq gives KClAq, and I; with excess of Cl, ICl, is formed and remains in solution, or combines with the KCl and separates as KCl.ICl, according to the quantity of water present. Bromine acts similarly to Cl.—6. Nitrous acid solution sets I free; KNO, and dil. HClAq act in the same way (v. Price, C. J. 4, 155).—7. Conc. sulphuric acid, or nitric acid, decomposes KI when heated with it, giving sulphate or nitrate of K, and I; with H₂SO₄, SO₂ and H₂S are also formed. Addition of conc. H.SO, to KIAq, or evaporation with HNO,Aq, sets I free. If KIAq contains KIO, a few drops of a dilute acid suffice to give free I (5HIAq + HIO,Aq = 3I₂ + 3H₂O). Aqua regia sets I free from KIAq (for the delicacy of this reaction v. Harting, J. pr. 22, 46).—8. Heated with sulphuric anhydride, K₂SO₄, SO₂, and I are formed (H. Rose, 38, 121).—9. Manganese dioxide and sulphuric acid produce I, MnSO,, and K2SO, when heated with KI.—10. KI fused with potassium-hydrogen sulphate gives K2SO. and I.-11. Fusion with potassium chlorate produces KIO₃-12. Heating with barium nitrate forms KIO, and BaO, (Henry, J. Ph. 18, 845).-13. By heating with ammonium chloride, I, AmI, and KCl are formed.—14. Conc. potassium ferricyanide solution gives I K.FeOy, Aq; on dilution the action is reversed (Mohr, A. 105, 57).—15. Heated with potassium borate, KIO, and a polyborate (? probably $K_2B_2O_{10}$) are formed (Schiff, A. 228, 72).

Combinations.—1. With most iodides of less positive metals (v. various metallic iodides). Several of the compounds produced are best regarded as K salts of metal-containing acids (v. Remsen, Am. 11, No. 5).—2. With arsenious oxide to form KI.As.O. (Schiff, A. 228, 72).—3. With potassium ferricyanide, to form a very unstable compound, KI.K., FeCy. (Preuss, A. 29, 823; Mohr, A. 105, 57; Blomstrand, J. pr. [2] 3, 207; Kern, C. N. 33, 184).

Potassium tri-iodide KI₃. Dark-blue, lustrous needles; almost black by reflected light; melts at 45°; S.G. 3·498 at c. 15°; by saturating KIAq with I, and evaporating over H₂SO₄ (Johnson, C. J. 31, 249). Very deliquescent; in a little water some I is deposited; sol. alcohol, in a very small quantity water, and in saturated solution of I in KIAq. Agitation with CS₂ of cone. KIAq saturated with I abstracts the dissolved I (Baudrimont, C. R. 51, 827). Jörgensen (J. pr, [2] 2, 347) found, however, that CS₂ did not remove the I from an alcoholic solution of KI containing excess of I in ratio KI:2I, and that alcoholic KI completely decolourised solutions of I in CS₂.

Potassium di-iodide ?KI₂. A solution of 3 pts. I in 4 pts. KI in water is said by Guyard (Bl. [2] 31, 297) to contain an unstable compound of this composition; with Pb(NO₃)₂Aq or Pb(C₂H₃O₂)₂Aq, this solution gives an almost black prossid by G to be PbI

black pp., said by G. to be Pbl..

Potassium, iodobromide of. The compound
KBr.IBr may be called iodobromide of potassium;
v. Potassium bromide, Combinations, p. 300.

Potassium, iodochlorides of. The compounds KCl.ICl₂ and KCl.ICl may be called iodochlorides of potassium; v. Potassium chloride, *Combinations* Nos. 3 and 4, p. 301.

Potassium, iridicyanide of; v. vol. ii. p. 332.

Potassium mangani- and mangano-cyanide of; v. vol. ii. p. 342.

Potassium, nitride of, K₃N. Formed by heating to redness KNH₂ (v. Potassium amide, p. 299) in absence of O. A dark-grey, almost black, solid. Sublimes partially at white heat. Takes fire in air, with evolution of N; reacts with water to give KOHAq and NH₂. K₃N is a conductor of electricity. Combines with S and P (Gay-Lussac a. Thénard, A. Ch. 65, 325).

Potassium, nitroprusside of; v. vol. ii. p. 341.

Potassium, osmocyanide of; v. vol. ii. p. 343.

Potassium, oxides of. Two oxides have been certainly isolated, K_2O and K_2O_4 ; there are indications of the existence of others.

Potassium oxide K₂O (Potassium monoxide). A mixture of K₂O and K₂O₄ is formed by burning K in air; when K is burnt in a stream of dry O only K₂O₄ is produced, but on strongly heating this product O is given off, and K₄O remains (Davy, T. 1808 [1] 5). K₂O is also formed by heating KOH and K in the ratio KOH: K; according to Beketoff (Bl. [2] 37, 491), K and KOH do not react. B. says that K₂O may be prepared by heating a mixture of K₂O₄ and K in a silver dish. To prepare pure K₂O Kühnemann (C. C. 1864. 491) recommends to lead air, freed from CO₂, over K heated just to fusion; the K spreads out into a thin layer

which burns to K_2O . K_2O is a greyish, non-lustrous solid; S.G. 2.656 (Karsten, S. 65, 394); melts at full red-heat, and volatilises at a very high temperature. Thomsen (Th. 3, 235) gives $[K_1^*O,Aq] = 165,460$. K_2O reacts with water to form KOHAq, with production of much heat. Von Schaffgotsch (A. 43, 17) says that $K_2O.Fe_2O_3$ is obtained, but not pure, by heating Fe_2O_3 with K_2CO_3 .

Potassium tetroxide K2O4 (Potassium peroxide). The formation of this compound by heating K in dry air or O was observed by Gay-Lussac a. Thénard (A. Ch. 65, 325). The preparation and properties of the compound were examined by Vernon Harcourt (C. J. 14, 267). Prepared by heating K till it melts, in a hard glass flask filled with N, then sending in a slow stream of dry air, and continuing to heat in the air-stream until the metal is completely changed to a yellowish amorphous solid, and heating this for some time in dry O. It is also formed by throwing small pieces of K into molten KNO₃ (Bolton, C. N. 52, 289). A mixture of K_2O_4 with KOH.H₂O is said to be formed by mixing equivalents of KOH and H2O2 in Aq, and evaporating in vacuo (Schöne, A. 193, 241). K,O, is an amorphous powder, having the colour of PbCrO₄. It begins to soften at 280°, and melts at red heat to a black liquid, which crystallises in leaflets on cooling. At white heat gives off O and leaves K₂O. Deliquesces in moist air, giving off O. In water forms KOHAq and H2O2Aq, and evolves O. Heated in H produces KOH and H₂O. K₂O₄ oxidises P, S, C, &c., and many metals to their highest oxides. Reacts with CO, when heated, giving K₂CO₃ and O. No action with N₂O; with NO forms KNO₂ and KNO, and gives off N oxides.

Oxides of Potassium other than K₂O and K₂O₄. Harcourt (l.c.) asserted the existence of K₂O₂, obtained by oxidation of K. Lupton (C. J. 1876. [2] 565) thought that the oxides K₆O₃, K₄O₄, and K₆O₄ are formed by the action of air on K at c. 65°; these oxides may be regarded, according to L., as 3K₂O.K₂O₂, K₂O.K₂O₂, and 2K₂O.K₂O₂ respectively. The substance described as a suboxide, K₄O, is said by L. to be a mixture of K and K₂O. Schöne (A. 193, 241) obtained K₂H₄O₆, probably K₂O₂·2H₂O₃, by evaporating, in vacuo, at -10°, a mixture of 1 equiv. KOH and 2 equivs. H₂O₂ in Aq.

Potassium, phosphide of. K and P combine when heated under rock-oil, or in N, forming a dark-yellow solid (Magnus, P. 17, 517; Vigier, A. 122, 331). A compound, or compounds, of K and P seems also to be formed when K is heated in PH₃, and also when K is heated in P in a stream of H (H. Rose, P. 12, 547). The compound prepared by Rose was a crystalline, reddish, lustrous solid; decomposed by water, giving KH₂PO₂Aq and PH₃, and burning, when heated in air, to K phosphate.

Potassium, platinooyanide of, and allied salts; v. vol. ii. pp. 344-5.

Potassium, salts of. Compounds formed by replacing H of acids by K. The chief salts of K, besides the halides and sulphides, are carbonates, chlorate and perchlorate, chromate and derivatives, iodate and periodates, nitrate and nitrite, phosphates and derivatives, silicates, Von IV

sulphates and derivatives, sulphite, thiosulphate (v. Carbonates, Phosphates, &c.).

Potassium, selenides of. Berzelius (Lehr. buch, 5th ed. 2, 222) investigated the conditions of combination of K and Se, without isolating and analysing any definite compounds. Se and K combine when melted together; a steel-grey solid sublimes, which dissolves in water with formation of a red liquid that decomposes in air with separation of Se, and gives off H2Se with acids. Compounds of Se and K are also formed by reducing K₂SeO₄ or K₂SeO₅ by C or H at red heat (Wöhler a. Dean, A. 97, 5; Rathke, A. 152, 211). Mixtures of selenate and selenides of K are formed by fusing KOH or K2CO, with Se. Boiling KOHAq dissolves Se, forming a brown liquid, from which a brown powder separates on evaporation, and which gives Se when heated with acids, or when diluted and allowed to stand in air.

Fabre (C. R. 102, 613) obtained $K_2Se.xH_2O$, x=9, 14, and 19, by passing a rapid current of H_2Se into KOHAq of different concentrations; these hydrates of potassium monosclenide are all very easily decomposed. Fabre (C. R. 102, 703) gives the following thermal data: $[K^2,Se] = 98,340$ from solid elements; $[2KOHAq,H_2SeAq] = 7,520$; $[2KOHAq,H_2Se] = 16,920$; $[K^2Se,Aq] = 8,540$.

Potassium, selenocyanide of, v. vol. ii. p. 348. Potassium, selenostannate of, K.SnSe, 3aq. Yellow, unstable crystals; by saturating K.SeAq with SnSe₂ and evaporating in vacuo (Ditte, C. R. 95, 641).

Potassium, selenothiostannate of,

K.SnSe.S 3aq. Yellow, octahedral crystals; by boiling K.SAq with Sn and Se, and evaporating in vacuo. Sol. water, with red colour. Both solid and solution in water very readily undergo decomposition, with separation of Se (Ditte, l.c.).

Potassium, silicide of. By reducing SiO₂ by K, Berzelius obtained a brown solid, which reacted with water to give H, KOHAq, and SiO₂ (*Lehrbuch*, 5th ed. 2, 84).

Potassium, silicofluoride of, K2SiF6. Prepared by adding H.SiF.Aq to a fairly conc. solution of a K salt, washing, and drying. Small tesseral white crystals. Preis (Listy Chem. 13, 150; abstract in C. J. 58, 694) found hexagonal, and also regular, crystals of this salt, along with CaSO, in a basin wherein phosphoric acid K2SiF6 had been prepared from phosphorite. is isodimorphous with the ammonium salt. 2.665 at 17.5° (Stolba, J. pr. 97, 503). Easily sol. hot water, very sl. sol. cold water. Melts at low red heat, giving off SiF, and leaving KF (v. Bothe, Hofman's Ber. uber die Entwick. der Chem. Industrie, 1, 318). Decomposed by conc. H₂SO₄, giving off SiF₄ and HF. Decomposed by boiling (not cold) solutions of alkali hydroxides or carbonates, with separation of gelatinous silicic acid and formation of KF.

Potassium, siliconitride of. The white, infusible solid, obtained by heating SiO₂ with KCN, is probably a compound of K, Si, and N (Balmain, P. M. October 1842).

Potassium, sulphides of. K and S combine when heated together, the K burning brilliantly. Five compounds of K and S are known: K₂S₂, K₂S₃, K₂S₄, and K₂S₅. They are all solwater, the polysulphides also in alcohol. K₂SAq is colourless; solutions of the others are coloured

yellowish brown, the colour being deeper the greater the proportion of S. Solutions are decomposed by acids; K₂SAq with evolution of H₂S, the other solutions also with ppn. of S. Solutions decompose in air; K₂SAq to K₂S₂O₃Aq and KOHAq, K₂S₂Aq to K₂S₂O₃Aq, and the others to K₂S₂O₃Aq with separation of S. Solutions of the polysulphides react with K₂SO₄Aq to form K₂SAq and K₂S₂O₃Aq; these solutions are reduced to K₂SAq by shaking with Hg. With EtI or EtBr they form Et₂S₂, which dissolves the S set free; these solutions dialyse unchanged (Spring a. Demarteau, Bl. [3] 1, 311).

POTASSIUM MONOSULPHIDE K28. Formed by reducing dry K2SO4; by heating in H (Berzelius, P. 6, 438); by heating with C (Berthier, A. Ch. 22, 233; Bauer, *J. pr.* 75, 246; Wittstock, *P.* 55, 536). Not obtained free from polysulphides by these methods; if vessels of glass or porcelain are used the product contains SiO₂, as K₂S acts on silicates. The product of the foregoing reactions is a brown, crystalline, deliquescent solid; sol. in water, with production of much heat and formation of KSHAq and KOHAq; Thomsen (Th. 3, 235) gives $[K^2, S, Aq] = 113,800$. Sabatier (A. Ch. [5] 22, 5) says that K_2S has never been obtained pure. An aqueous solution of K2S is prepared by saturating KOHAq with H₂S, and adding an equal quantity of the same KOHAq. By evaporating this solution in vacuo at a low temperature, the pentahydrate K2S.5H2O is obtained in orthorhombic crystals, which lose $3H_2O$ by heating to 150° , out of air (Schöne, P. 131, 880; cf. Sabatier, l.c.). K₂SAq is colourless out of contact with air; it is very alkaline and caustic; in air it becomes yellow, H₂S is given off, and K2CO2Aq and K2S2O3Aq are formed. K.SAq dissolves S, forming polysulphides; it

dissolves Fe or Fe₂O₃, forming a green liquid.

Potassium disulphide K₂S₂. Formed by allowing an alcoholic solution of K₂S to stand in air till the surface is covered with a film of K₂S₂O₃, then pouring off, and evaporating in vacuo (Berzelius, l.c.). Also by heating 2 equivalents KHSO₄ with 7 or more equivalents C (Geiger). Berzelius obtained K₂S₂ by heating 4 equivalents K₄CO₃ with rather less than 7 equivalents S. A yellowish red, deliquescent solid; K₂S₂Aq is yellow; in air it oxidises to K₂S₂O₃Aq without separation of S; acids give off H₂S and

cause ppn. of S.

Potassium trisulphide K₂S₂. Formed by passing CS₂ over K₂CO₃ heated to redness (Berzelius, *l.c.*; cf. Schöne, *l.c.*). Also by heating 100 parts K₂CO₃ with 58-22 parts S to dull redness, till CO₂ ceases to be given off; the product contains some K₂SO₄. Also, with some S, by passing H₂S over strongly-heated K₂SO₄. By heating K₂S₄ to over 800°. A yellowish-brown solid, black when molten; not decomposed at 900°, gives off S at white heat (B.). Easily sol. water and alcohol; K₂S₄Aq in air gives K₂S₂O₄Aq with separation of S; decomposed by acids, S separating and H₂S being evolved.

Potassium tetrasulphide K₂S₄. Formed by passing CS₂ over strongly-heated K₂SO₄ so long as CO₂ is produced. Also by melting K₂CO₂ with 1 to 2 parts S, heating till excess of S is removed, and then reducing K₂SO₄ formed by heating in H₄S. A red-brown, crystalline solid (B., Lc.; S., Lc.). Decomposed at 600°-900° to K₂S₂ and S.

The dihydrate K₂S₄.2H₂O is obtained by boiling K₂SAq with the proper quantity of S, and evaporating in vacuo (S., l.c.). Sabatier (A. Ch. [5] 22, 53) obtained the octahydrate, K₂S₄.8H₂O, by adding 90 p.c. alcohol to K₂S₄Aq, separating the brown oily liquid, and letting it crystallise.

brown oily liquid, and letting it crystallise.

Potassium pentasulphide K,S., Formed by heating any of the lower sulphides with S until the excess is removed; Schöne (l.c.) says temperature should not exceed 600° (v. B., l.c.). A solution of K,S., containing also K,S.O., is formed by heating K,CO.,A with S (Fordos a. Gélis, C. R. 23, 211); or by digesting a solution of a lower sulphide with S (B., l.c.). A red, fusible solid; decomposed by heating in steam, giving K,SO., and H,S (Drechsel, J. pr. [2] 4, 20). Very sol. water, also in alcohol; K,S.Aq slowly decomposes in air, giving K,SO.,Aq, K,CO.,Aq, and S; acids evolve H,S and separate S.

Liver of sulphur. This name is given to the brown solid obtained by heating S with K₂CO₃ in a closed vessel; it contains various sulphides of K, along with K₂SO₄, and generally

some K₂CO₃.

Potassium, sulphydrate of; v. Potassium hydrosulphide, p. 302.

Potassium, sulphocyanide of; v. vol. ii. pp. 351-2.

Potassium, telluride of. Probably K₂Te. By heating together K and Te in H; also by heating a mixture of 100 pts. H₂TeO₂, 20 pts. KOH, and 10 pts. C. A brittle, crystalline, coppercoloured solid. Easily sol. water; exposed to air Te separates from the solution. HClAq evolves H₂Te (Davy; v. also Berzelius, Lehrbuch (5th ed.) 2, 247).

Potassium, tellurofluoride of, KF.TeF4; v.

Potassium fluoride, p. 301.

Potassium, thio-arsenates and thio-arsenites of; v. vol. i. pp. 317, 316.

Potassium, thiocarbonate of; v. vol. i. p. 703.

M. M. P. M.

PRASEODYMIUM. The constituent of didymium which yields green-coloured salts; v. DIDYMIUM, vol. ii. p. 383.

DIDYMIUM, vol. ii. p. 383.

PREHNITENE v. c-DURENE.

PREHNITIC ACID C₁₀H₂O₈ i.e. C₆H₂(CO₂H)₄[1:2:3:4]. Mol. w. 254. [237°-250°]. Formed by heating either of the tetrahydrides of pyromellitic acid with H₂SO₄ (Baeyer, A. 166, 254), and by the oxidation of c-durene and of c-tetra-ethyl-benzene (Töhl, B. 21, 907; Galle, B. 16, 1746; Jacobsen, B. 17, 2518). Crystallises from water in large prisms (containing 2aq) resembling the mineral prehnite. Converted by fusion into an anhydride C₁₀H₄O, [239°]. May be reduced to a viscid hydride.—KH₂Al'aq.—BaH₂A₂''3aq.—BaH₂Al' aq: needles.—Pb₂Al': small needles, insol. water.

Methyl ether Me,A'v. [104°-108°].

PREHNITOL v. c-DURENOL. PREHNOMALIC ACID C10H,O, i.e.

C₀H₃(OH)(CO₂H)₄. Oxy-prehnitic acid dihydride. [210°]. A product of the action of H₂SO₄ on the tetrahydrides of mellitic acid (Baeyer, A. 166, 325; B. 4, 275). Needles, readily converted by heat into an anhydride $C_{20}H_{14}O_{17}$ [210°]. By heating with H₂SO₄, or by treatment with Br, it is converted into prehnitic acid.—Ag,A¹⁷.

PRIMULA CAMPHOR O₁₁H₁₂O₅. [49°]. (above 200°). Contained in the root of *Primula*

veris (Mutschler, A. 185, 222). Six-sided plates, v. sl. sol. water, v. sol. alcohol and ether. Smells like anise. FeCl, colours its aqueous solution violet. Yields salicylic acid on oxidation.

PRIMULINE BASE C28H18N4S3 i.e. $C_{\bullet}H_{\bullet}Me < N > C.O_{\bullet}H_{\bullet} < N > C.C_{\bullet}H_{\bullet} < N > C.C_{\bullet}H_{\bullet} < N > C.C_{\bullet}H_{\bullet}NH_{\bullet}$

Prepared by heating p-toluidine (2 mols.) with sulphur (4 or 5 atoms), H_2S being evolved (Green, C. J. 55, 234). Yellowish powder, nearly insolall solvents. Not decomposed at 400°. Its salts are decomposed by water. The Na salt of its sulphonic acid (primuline), is taken up by unmordanted cotton. KOH at 260° yields amidotolyl-mercaptan and p-amido-benzoic acid (Gattermann, B. 22, 425, 1066).

Dehydrothiotoluidine C14H12SN2 i.s.

 $C_eH_3Me <_S^N > C.C_eH_4.NH_2.$ [191°]. (434° at 766 mm.) (Pfitzinger a. Gattermann, B. 22, 1066). An intermediate body in the preparation of primuline. Formed, together with primuline base, by heating p-toluidine with S (Dahl & Co., G. P. 35790; Green, C. J. 55, 230). Got also by the action of HI and P upon primuline (Jacobsen, B. 22, 330). Yellowish iridescent needles (from isoamyl alcohol), v. sol. HOAc, sl. sol. hot alcohol, v. sl. sol. benzene, almost insol. boiling water. Its alcoholic solutions have violet-blue fluorescence. Ppd. by adding water to its solution in HClAq.

Reactions.—1. Yields p-toluidine on distillation with zinc-dust. - 2. Yields primuline on heating with S.—3. Converted by diazotisation in

boiling alcoholic solution into $C_6H_3Me < N > CPh$

[123°].—4. When diazotised and boiled with water it gives $C_{14}H_{11}(OH)N_2S$ [256°], which gives $C_{14}H_{11}(OAc)N_2S$ [132°].—5. Fuming H_2SO_4 (containing 70 p.c. SO₈) forms, below 50°, a sulphonic acid C14H11(SO3H)SN2, which crystallises in yellow needles (containing aq), or orange plates (containing 2aq), sl. sol. hot water, and forms NH, A'aq, sl. sol. hot water, CuA', 2aq, a reddishbrown pp., and AgA', a white pp.—6. MeI yields $C_{1_i}H_{1_0}(NMe_2)NS$ [197°] (434°), and $C_{1_i}H_{1_0}(NMe_3)NS$, a bright-yellow powder, v. sol. water. MeOH and HCl at 150°200° give water. MeOH and HCl at 150°-200° give C₁₄H₁₀(NMe,Cl)NS, a yellow powder which forms $(C_{14}H_{10}NS.NMe_{3}Cl)_{2}PtCl_{4}$ [234°].

Acetyl derivative C14H11AcN2S. [227°]. White plates or prisms, sl. sol. HOAc.

PROPACONIC ACID v. Lactone of Oxy-BUTYLMALEÏC ACID.

PROPANE C.H. i.e. CH3.CH2.CH3. Mol. w. 44. (-17°). S. (alcohol) 6. H.F.p. 35,110. H.F.v. 33,370 (Thomsen, Th. 4, 52). Occurs in the gases given off from the petroleum springs of the United States (Ronalds, C. J. 18,54; Lefevre, Z. [2] 5, 185). Formed by the action of HI on propyl iodide, allyl iodide, isopropyl iodide, acetone, and glycerin (Berthelot, Bl. 7, 60; 9, 13, 184). Prepared by heating n-propyl iodide (9.6 pts.) with AlCl, (2.5 pts.) at 140° (Köhnlein, B. 16, 560), or by heating isopropyl iodide with zinc and HClAq (Schorlemmer, A. 150, 209). Gas. On compression with water at a low temperature it yields a hydrate, dissociating at 8.5° (Villard, C. R. 111, 302).

References .- DI-BROMO-, BROMO-IODO-, BROMO-

NITRO-, CHLORO-, CHLORO-IODO-, DI-CHLORO-NITRO-, DI-10DO, and NITRO- PROPANE

PROPANE CARBOXYLIC ACID v. BUTTRIC ACID.

Propane dicarboxylic acid v. GLUTARIC. Pyrotantaric, and Ethyl-malonic acids.

Propane tricarboxylic acid C.H.O. i.e. CH, CH(CO₂H).CH(CO₂H)₂ (Bischoff, B. 13, 2164; 14, 614; 15, 1107; 17, 2783; 22, 8180; A. 214, 53). [146°]. Formed by saponifying the ether with dilute alcoholic potash. Crystalline mass, v. e. sol. water, alcohol, and ether. Splits up when heated alone, or with dilute HClAq, into CO2 and pyrotartaric acid. Bromine forms bromopyrotartaric and bromocrotonic acids.—Ba₃A'''₂: bulky pp.

ds.—Ba,A''', bulky pp.

Methyl-di-ethyl ether MeEt,A'''. (268°). S.G. 15 1078. Formed from methyl a-chloro-

propionate and sodium malonic ether.

Ethyl ether Et₃A'''. (270.3° cor.). S.G. 20 1.0698. $\mu_D = 1.4288$ at 20°. Formed from sodium malonic ether and α-bromo-propionic ether. Oil, miscible with alcohol and ether. NaOEt yields C,H,Na(CO,Na)(CO,Et),.

Propane tricarboxylic acid CH_3 . $C(CO_2H)_2$. CH_2 . CO_2H .

Methyl ether Me.A.". Oil (Barthe). Ethyl ether Et.A.". (273.5° cor.). S.G. \ref{Grade} 1.077. $\mu_D=1.4311$ at 20°. Formed from sodium methyl-malonic ether and chloro-acetic ether, and also from sodium ethane tricarboxylic ether and MeI or MeCl (Bischoff a. Von Kuhlberg, B. 23, 635). Yields on saponification pyrotartaric and a little succinic acid.

Methyl ether of the mono-nitrile CH₂.C(CN)(CO₂Me).CH₂.CO₂Me. Formed from methyl cyanosuccinate, MeOH, and at 70° (Barthe, Bl. [3] 1, 303; C. R. 108, 297; 112, 1013). Oil. The corresponding CH, C(CN)(CO,Et).CH, CO,Et is also an oil (185° at 35 mm.).

Propane tricarboxylic acid

CO₂H.CH₂.CH₂.CH(CO₂H)₂. Ethyl ether Et₃A". (161° at 13 mm.). S.G. 20 1.0808. Formed from sodium malonic ether and β -bromo-propionic ether (Emery, B. 24, 282). The free acid splits up into CO, and glutaric acid on boiling with HClAq.

Isomeride v. TRICARBALLYLIC ACID. Propane tetracarboxylic acid

 $CO_2H.CH_2.CH(CO_2H).CH(CO_2H)_2.$ Ethylether Et, A' (204° at 18 mm.). S.G. 20 1.1184. Formed from sodium malonic ether and chloro-succinic ether (Emery, B. 23, 3759). Oil, v. sol. alcohol and ether. The free acid

splits up into CO2 and tricarballylic acid. Propane tetracarboxylic acid (CO₂H)₂CH.CH₂CH(CO₂H)₂. Dicarboxy-glutaric acid. [170°]. An acid formed by the action of

sodium amalgam on dicarboxyglutaconic ether C,H,Et,O, in alkaline solution (Conrad a. Guthzeit, A. 222, 257). One of the products of the action of chloro-di-methyl oxide on sodium malonic ether (Kleber, A. 246, 106). Got also by boiling its ether with dilute alcoholic potash Guthzeit a. Dressel, B. 21, 2234; 22, 1423; A. 256, 174). Crystalline powder (from ether), v. e. sol. water. At 180° it splits up into CO₂ and glutaric acid [98°].-Ba,A'v 2aq.-Ca,A'v 2aq.-Pb2A" 2aq.

Ethyl ether Et.Air. (285% at 80 mm.).

S.G. 20 1.116. Formed from sodium malonic ether by treatment with CH_2Cl_2 or CH_2I_2 , and got also by reducing dicarboxy-glutaconic ether with zinc-dust and HOAc (W. H. Perkin, jun., C. J. 59, 993). Oil, with very bitter taste. NaOEt forms a di-sodium derivative, which reacts with EtI forming oily C17H28O8 (c. 248° in vacuo).

Propyl-tri-ethyl ether Et, PrA'v. (195°-202° at 15 mm.). Formed from propyl triethyl dicarboxy-glutaconate, zinc-dust, and HOAc.

Colourless oil.

Propane tetracarboxylic acid

CO₂H.CH₂·C(CO₂H)₂·CH₂·CO₂H. [151°]. Got by saponifying its ether with alcoholic potash. Prisms (from water), v. sol. water, alcohol, and ether. At 170° it splits up into CO2 and tricarballylic acid.—K₃HA¹, 2₂aq (at 100°). Zn₂A'v Saq (dried at 100°).—Pb₂A'v aq.—Ag₄A'v: insoluble pp.

Ethyl ether Et.A. (295° uncor.). S.G. \$ 1.102. Formed from malonic ether, alcoholic NaOEt, and chloro-acetic ether, followed by a second dose of NaOEt and chloro-acetic ether (Bischoff, B. 13, 2163; A. 214, 61). Oil.

Ethyl ether of the mono-nitrile CO₂Et.CH₂.C(CN)(CO₂Et).CH₂.CO₂Et. [41°]. (200°-215° at 10 mm.). Formed from sodium cyano-succinic ether and chloro-acetic ether (Haller a. Barthe, C. R. 106, 1413). crystals, sol. alcohol and ether.

Methyl ether of the mono-nitrile $CO_2Me.CH_2.C(CN)(CO_2Me).CH_2.CO_2Me.$ Prepared in like manner (Barthe, C. R. 111, 344). Prisms, insol. water and alkalis.

Propane pentacarboxylic acid $CH(CO_2H)_2$. $C(CO_2H)_2$. CH_2 . CO_2H . [151°]. Got by saponifying its ether. Spherical aggregates (from ether).— K_5A^* 4aq.— Ba_5A^* 24aq: crystalline

Ethyl ether Et,A. (275°-280° at 188 mm.). S.G. $\frac{15}{15}$ 1·121 (Bischoff, B. 15, 1107; 21, 2118). Formed from sodium-malonic ether and CCl(CO₂Et)₂.CH₂.CO₂Et. Formed also from sodium ethane tricarboxylic ether and chloroacetic ether. Oil.

Reference.—OXYPROPANE TRICARBOXYLIC ACID.

PROPANE PHOSPHONIC ACID

C₂H₇.PO(OH)₂. [60°-70°]. Prepared by oxidation of propyl-phosphine with fuming HNOs (Hofmann, B. 6, 304). Wax-like mass, sol. water.

PROPANE SULPHONIC ACID

CH₂.CH₂.CH₂.SO₂H. Formed by oxidation of propyl mercaptan with nitric acid (S.G. 1.3) (Spring a. Winssinger, Bl. [2] 48, 110). ICl, at 160° forms chloro-propane sulphonic acid, CCl, and C,Cl,.

Propane sulphonic acid (CH_s)₂.CH.SO₃H. [below 100°]. Formed by oxidising isopropyl mercaptan with HNO, Obtained also from acetone by successive treatment with P2S5 and HNO_s (Claus, B. 5, 660; 8, 533; Spring, Bl. [2] 40, 66). Crystalline. Its salts are v. e. sol.

Propane disulphonic acid

CH₂(SO₃H).CH₂.CH₂(SO₃H). Formed by boiling trimethylene bromide with a saturated solution of ammonium sulphite (Monari, B. 18, 1345). Deliquescent needles, v. sol. water and alcohol.-Na,A" 41aq. BaA" 2aq: very soluble needles.

Propane disulphonic acid

 $CH_3.CH(SO_3H).CH_2(SO_3H).$ Formed in like manner from propylene bromide (M.).—Na₂A"aq.

-BaA": sparingly soluble crystals.

Propane disulphonic acid C3H6(SO3H)2. Got from butyramide and fuming H.SO. (Buckton a. Hofmann, A. 100, 153), and also by heating butyric acid with ClSO₈H at 140° (Baumstark, д. 140, 83).—BaA".—PbA": crystalline.

Propane trisulphonic acid CH₂(SO₃H).CH(SO₃H).CH₂(SO₃H). Got by boiling s-tri-chloro-propane with aqueous K2SO, (Schäuffelin, 4. 148, 117).—Ba₃A'''₂: crystalline powder, sl. sol. waver.

References .- CHLORO- and OXY-PROPANE SUL-

PHONIC ACID

DIPROPARGYL C.H., i.e. CH:C.CH₂.CH₂.C:CH. Hexunene. (85°). 18 ·81. H.F.p. -96,040. H.F.v. -97,200 (Thomsen, Th.). Formed by distilling diallyl tetrabromide with solid KOH, and boiling the resulting dibromo-diallyl with alcoholic potash (Henry, B. 6, 956; 7, 21; 14, 399; 17, 1132). Limpid, highly refractive oil, v. sol. ether. Burns with smoky flame. Combines with explosive violence with bromine, forming C₆H₆Br₄, a thick liquid S.G. 19 2.464, from which, by further action of Br, crystalline CaHaBrs [141°] can be obtained. Iodine in KI forms dipropargyl tetraiodide C₆H₈I₄ [113° crystallising from CS_2 in prisms. Dipropargyl gives with ammoniacal cuprous chloride a yellow pp. $Cu_2C_0H_4$ 2aq, which explodes at about 100°. Aqueous AgNO₂ ppts. $Ag_2C_0H_4$ 2aq, which explodes below 100°. Dipropargyl is very easily polymerised, forming a resin.

ymerised, forming a resum.

PROPARGYL ALCOHOL C₃H₄O i.e.

76. (115°). V.D. 1.88 CH:C.CH₂OH. Mol. w. 56. (115°). V.D. 1·88 (calc. 1·93). S.G. $^{24}_{0}$ ·9715. $\mu_{\beta} = 1\cdot437$. R_{\infty} = 24·01 (Br\(\text{ihl}\)). H.F.p. -3,500. H.F.v. -4,370 (Thomsen, Th.). Formed by boiling bromo-allyl alcohol with KOHAq (Henry, B. 5, 274, 449, 569; 6, 728; 7, 20, 761). Formed also from propargylamine oxalate, NaNO2, and HCl (Paal a. Hempel, B. 24, 3039). Colourless liquid of peculiar odour, sol. water.

Reactions.—1. Ammoniacal Cu₂Cl₂ ppts. canary-yellow Cu₂(C₂H₃O)₂ which explodes when heated, and is decomposed by dilute acids with reproduction of propargyl alcohol.-2. Ammoniacal AgNO, ppts. white AgC, H,O.-3. Anhydrous BaO forms Ba(C,H,O),C,H,O, crystallising in small plates.—4. On heating with solid KOH it yields formic acid and acetylene.-5. HBr

forms bromo-allyl alcohol.—6. HNO2 yields CO2 and oxalic acid.

Acetyl derivative C_3H_3 .OAc. (125°). S.G. 29 1·0052. $\mu_{\beta} = 1\cdot427$. $R_{\infty} = 89\cdot71$ (Bruhl).

Got from the alcohol and AcCl.

Methyl ether C3H3OMe. $(62^{\circ}).$ 1275 ·83. H.F.p. -10,910. H.F.v. -12,860 (Thomsen, Th.). Formed by boiling s-tri-bromo-propane with KOH dissolved in MeOH (Liebermann, A. 135, 287) and by boiling the dibromide of methyl allyl oxide with alcoholic potash (Henry). Ammoniacal AgNO, ppts. lemon-yellow C.H.Ag.OMe, whence I in KI forms oily $C_{2}H_{2}I.OMe[12^{\circ}].$

(81°-85°). S.G. Ethyl ether C,H,OEt. 20 ·8326. $\mu_{\beta} = 1.4096$. $R_{\infty} = 39.5$ (Brühl). This body is formed by the action of alcoholic potash on CH_Br.CHBr.CH_Br, on CH_.CBr_.CH_Br, on dichloropropylene, or on ethyl bromoallyl oxide (Liebermann, A. 135, 278; 158, 230; Henry, B. 5, 274; Baeyer, A. 138, 196). Mobile liquid, with penetrating odour, v. sl. sol. water, miscible with alcohol. Combines · with Br, forming C3H3Br2.OEt. Boiling very dilute (1 p.c.) H₂SO₄ splits it up into alcohol and propargyl alcohol. Sodium forms C₃H₂NaOEt. Alcoholic silver nitrate forms crystalline (C₃H₂Ag.OEt)₂AgNO₃ converted by ammonia into amorphous C₃H₂AgOEt, whence I in KI forms crystalline C3H2IOEt and oily C3H2I3OEt. An ammoniacal solution of AgCl added to an alcoholic solution of C3H,OEt ppts. white Ammoniacal Cu2Cl2 gives (C₂H₂AgOEt),AgCl. yellow amorphous C,H,CuOEt.

Isoamyl ether C.H.OC.H... (140°-145°). PROPARGYLAMINE C, H, NH,. Formed by the action of NaOEt on bromo-allyl-amine and on di-bromo-propyl-amine (Paal a. Hermann, B. 22, 3080). The free base has not been isolated. -B'H₂C₂O₄. [143°]. Large tables (from water), sl. sol. alcohol. Gives a white pp. with ammoniacal AgNO₃.—B'C₆H₂(NO₂)₃OH. [189°]. Large red plates or tables.—B'HCl.—B'HBr. [130°-171°].—B'HI. [205°]. White plates, v.

e. sol. water.

PROPARGYL-ISOAMYL-AMINE C₈H₁₅N i.e. C₃H₃.NHC₅H₁₁. Formed from di-bromo-isoamylamine and NaOEt (Paal a. Hermann, B. 22, 3084). Liquid.—B'H₂C₂O₄ aq. [204]. White needles (from water).—B'HBr. [186°]. Pearly

PROPARGYL BROMIDE CH:C.CH, Br. (89°). S.G. 11 1.59. Formed from propargyl alcohol and PBr₃ (Henry, B. 6, 728). Liquid. Propargyl tribromide v. Tri-bromo-propyl-

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PROPARGYL-ISOBUTYL-AMINE C, H,3N i.e. CH:C.CH₂.NH.CH₂Pr. (135°). Formed from di-bromo-propyl-isobutyl-amine and NaOEt (Paal a. Heupel, B. 24, 3045). Liquid, miscible [148°].-B'2H2PtCl6. water.—B'HCl. [210°]. Needles and plates, [172°].—B'H₂C₂O₄. m. sol. cold water.

PROPARGYL CHLORIDE C3H3C1. S.G. 5 1.045. Formed from propargyl alcohol and PCl, (Henry, B. 8, 398). Mobile oil.

PROPARGYLIC ACID is Propiolic ACID.

PROPARGYL IODIDE CH:C.CH₂I. (c. 115°). S.G. 2 2.018. Formed from propargyl bromide and NaI in alcohol (Henry, B. 17, 1132). Yellowish liquid. Iodine forms CHI:CI.CH2I. [41°].

PROPARGYL-PROPYL-AMINE C.H., N i.e. C3H3.NH.C3H7. The salt B'HBr [180°] is got from propyl bromide and propargylamine (Paal

a. Hermann, B. 22, 3084)

PROPARGYL SULPHOCYANIDE C.H.SCN. Formed from propargyl bromide and potassium sulphocyanide (Henry, B. 6, 729). Oil, smelling

like mustard. Decomposed by heat.

PROPARGYL-DI-THIO-CARBAMIC ACID CH:C.CH₂.NH.CS₂H. [115°]. Formed by boiling propargylamine with CS_2 and alcohol (Paal a. Heupel, B. 24, 3041). Needles (from dilute alcohol) or plates (from benzene-ligroin).

PROPENYL-o-AMIDO-PHENOL OXYLIC ETHER C.H. < NH > CMe.CH. CO.Et. [108°]. Formed by heating a mixture of o-amidophenol and acetoacetic ether (Hantzsch, B. 16, 1948). Flat prisms, readily resolved into the parent substances by dilute acids, alkalis, or even boiling water.

PROPENYL-O-AMIDO-PHENYL MERCAP. C₉H₉NS i.e. C₆H₄<8 CEt. Formed by heating o-amido-phenyl mercaptan with propionyl chloride at 150° (Hofmann, B. 13, 21). Heavy oil.—B'₂H₂PtCl_s: large prisms.

PROPENYLAMINE CH3.CH:CH.NH2. Formed by the action of baryta on β -bromopropylamine (Hirsch, B. 23, 968). Does not decolourise bromine water. -B'sH,I,2BiI,: sixsided plates.

PROPENYL-BENZENE SULPHONIC ACID CH2: CMe. C6H4. SO3H. The salts of this acid are formed by heating the dry salts of the acid CMe₂(OH).C₆H₄.SO₄H (R. Meyer, A. 219, 302).

Amide C, H, SO, NH2. [152°]. o-PROPENYL-BENZOIC ACID

CH₂:CMe.C_aH₄.CO₂H. [61°]. Formed from dimethyl-phthalide and KCy at 260° (Wislicenus, A. 248, 64). Needles (from water).

p-Propenyl-benzoic acid CH₂.CMe.C₆H₄.CO₂H. [161°]. Formed by heating (CH₃)₂.C(OH).C₆H₄.CO₂H with dilute HCl on a water bath (R. Meyer a. J. Rosicki, B. 11, 1791; A. 219, 270). White satiny plates (from 1 pt. alcohol and 1 pt. water) or slender needles (from water). Sl. sol. hot water, v. e. sol. alcohol Sodium amalgam reduces it to and ether. cuminic acid.

Salts. - NH, A'. - BaA', aq. - CuA', 7aq. -AgA': amorphous pp.

Methylether. MeA' [53°]. (254° cor.). V.D. 6.09. Insol. water.

Isopropenyl-benzoic acid C3H3.C6H4.CO2H [255°-260°]. By boiling oxy-propyl-benzoic acid (CH_s)₂C(OH).C₆H₄.CO₂H with fuming HCl (R. Meyer, B. 12, 1076; A. 219, 281). Minute needles grouped in stars (from alcohol). water. Does not decolourise bromine, and is not reduced by sodium amalgam, but is converted by HI into cuminic acid. Hence it would appear to be a polymeride of propenyl-benzoic acid, not

containing a C:C group.

Salts.—NH,A'aq. Sl. sol. water.—BaA'2 aq. -CaA', 1 aq. —CuA', —AgA': pp., insol. water. Methylether MeA'. [835]. Decomposed

by heat.

References .- Amido-, Nitro-, and Oxy- Pro-PENYL-BENZOIC ACID.

PROPENYL BROMIDE v. BROMO-PROPYLENE. PROPENYL CARBINOL v. BUTENYL ALCOHOL. PROPENYL-TRICARBOXYLIC ACID v. Pro-PANETRICARBOXYLIC ACID.

PROPENYL CHLORIDE v. CHLORO-PROPYL-

PROPENYL-PHENOL. Methyl derivative v.

PROPENYL-ISOPROPYL BENZENE CioHie i.e. Pr.C.H., CH: CHMe. (230°). S.G. 12 890. Got by boiling Pr.C.H., CH., CBr.Me.CO.H with Na.CO.Aq (Perkin, C. J. 1877, ii. 660). Dibromide C.H., Br. [59°].

PROPENYL-SALICYLIC ACID v. Oxy-PRO

PENYL-BENZOIC ACID.

PROPERTONE v. PROTEÏDS.
PROPHETIN C₂₂H₃₂O₄. A bitter resin got from the fruit of Cucumis prophetarum, split up

by boiling dilute HClAq into glucose and propheteïn C₂₀H₂₀O₄ (Walz, J. 1859, 566). PROPINENE v. ALLYLENE.

PROPINENE DIPHTHALIDE C, H, O, i.e. -Ç:CMe.CH:Ç Ö.CO C6H4. Formed by C.H. < CO.O heating phthalic anhydride (3 pts.) with pyrotartaric acid (3 pts.) and sodium acetate (1 pt.) at 240°-290° (Roser, B. 17, 2776). Fine yellow needles, not melted at 280°.

PROPIO-GUANAMINE C,H,N, CH₂Me.C:N.C(NH) NH. Got by heating guanidine propionate at 230° for an hour (Haaf, J. pr. [2] 43, 78). Granules (from NaOHAq) or dimetric prisms (from water).

PROPIO-HOMO-FERULIC ACID v. DI-OXY-

PHENYL-CROTONIC ACID.

PROPIOLIC ACID C₂H₂O₂ i.e. CH¹C.CO₂H. Propargylic acid. [6°]. (c. 144°). Formed by boiling the acid K salt of acetylene dicarboxylic acid with water (Bandrowski, B. 13, 2340). Liquid, solidifying at about 4°. Smells like acetic acid. Turns brown in air. Sol. water, alcohol, and ether. Reduces HgCl₂, AgNO₃, and

Reactions.—1. Ammoniacal AgNO, gives a white crystalline explosive pp.-2. Ammoniacal Ou₂Cl₂ gives a green amorphous explosive pp., turning brown.—3. Sodium-amalgam reduces it to propionic acid.-4. Combines with HCl, forming β-chloro-acrylic acid. HBr and HI act in like manner.-5. Br forms di-bromo-acrylic acid [85°].-6. I forms CHI:CI.CO,H [104°] (Bruck, B. 24, 4120).

Salt.-KA' aq. Transparent prisms, v. sol.

water, deflagrating at 105°.

Ethyl ether EtA'. (119°). V.D. 3.47. Liquid, smelling like horse-radish. Oxidised by alkaline K_sFeCy_e to diacetylene dicarboxylic acid (Baeyer, B. 18, 677, 2269).

Reference. - Bromo- and Iodo- Propiolic

ACTD.

A. 265, 167).

PROPIONAMIDE v. Amide of PROPIONIC ACID PROPIONAMIDINE C3H8N2 i.e.

CEt(NH).NH2. The hydrochloride is formed by leading HCl into a solution of propionitrile in isobutyl alcohol, and heating the product with alcoholic NH_s (Pinner a. Klein, B. 11, 1484; 17, 176; 18, 2845). Very unstable oil, sl. sol. ether. On heating with NaOAc and Ac,0 it yields a base $C_8H_{18}N_8$ [204°].—B'HCl. [129°]. Needles, insol. ether.—B'₂H₂PtCl₈. [200°].—B'HNO₈. [116°]. Crystals (from alcohol), v. sol. water (C. Lossen,

Reference.—OXY-PROPIONAMIDINE. PROPIONE v. DI-ETHYL RETONE.

PROPIONIC ACID C3H6O2i.e. CH3.CH2.CO2H. Metacetonic acid. Mol. w. 74. $[-23^{\circ}]$ (Pettersson, J. pr. [2] 24, 295). (140·7° cor.) (Zander). S.G. $\frac{20}{3}$ '9946 (B.); $\frac{15}{15}$ '9983; $\frac{25}{25}$ '9896 (Perkin). M.M. 3·462 at 20°. $\mu_{\theta} = 1.391$. $R_{\infty} = 28.01$ (Brühl). S.V. 85·3. C.E. (0°-10°) '00106. H.F.p. 109,450. H.F.v. 107,710 (Thomsen, Th.). S.H. ·440 at 0° (R. Schiff, A. 234, 300). Critical temperature 340° (Pawlewski, B. 16, 2634). Occurs in the fruit of Gingko biloba (Béchamp, A. 130, 364), in suint (Buisine, Bl. [2] 48, 639), and in wood vinegar (Anderson, C. N. 14, 257; Barré, Z. [2] 5, 445).

Formation.—1. By exidation of metacetone with chromic acid (Gottlieb, A. 52, 121).-2. By potash-fusion from sugar, starch, mannite, and angelic acid (G.).—3. By reduction of acrylic acid (Linnemann, A. 125, 317), of lactic acid (Lautemann, A. 113, 217), and of pyruvic acid (Wislicenus, A. 126, 229).—4. By heating potassium oxalate with dry NaOEt (Van't Hoff, B. 6, 1107).-5. By heating NaOEt with CO at 180° (Fröhlich, A. 202, 290; cf. Hagemann, B. 4, 877).—6. From ZnEt₂, sodium, and CO₂ (Wanklyn, A. 107, 125).—7. Together with di-ethyl ketone, by heating ZnEt₂ with CO₂ at 160° (Schmitt, J. pr. [2] 42, 568).—8. By the action of sunlight on a solution of succinic acid containing a uranium salt (Seekamp, A. 133, 253). 9. By heating barium succinate with NaOMe for 3 hours at 300° (Mai, B. 22, 2133).—10. Among the products of destructive distillation of colophony (Renard, C. R. 103, 157).

Preparation .- 1. By oxidation of n-propyl alcohol (Pierre a. Puchot, A. Ch. [4] 28, 75).-2. By saponification of the nitrile (Linnemann, A. 148, 251; Beckurts a. Otto, B. 10, 262).-3. By heating lactic acid with HIAq (Freund, J. pr. [2] 5, 446).

Properties.-Liquid, miscible with water, alcohol, ether, benzene, and ligroïn. Separated from its aqueous solution by CaCl, but not by NaCl and Na, SO. Heated in a sealed tube with ICl, it gives C, Cl, CO, and HCl (Krafit, B. 9, 1085). On boiling the dry acid with Br and P for some time the product is a-bromo-propionic acid (Volhard, A. 242, 141).

Salts.—NH,A'.—(NH,)HA'₂ (Sestini, Z. 1871, 34).—NaA'. S. 99 at 15°; 187 at 100° (R.). S. (alcohol) 2·4 at 20°, 3 at 100°.—NaA'aq.— KA'aq: deliquescent plates (Renard, C. R. 104, 913). S. (of KA') 178 at 16°; 309 at 100°. KHA'₂.—LiA' aq.—CaA'₂ aq. S. 54 at 17° (L.); 41 at 0°; 39 at 80° (K.). Needles, insol. alcohol.— Ca₂HA'₅ 5aq.—BaA'₂ aq. S. 60 at 17° (Linnemann, A. 160, 220); 48 at 0°; 68 at 80° (von Krasnicki, M. 8, 605).—BaA'₂ 6aq.—BaHA'₃ 3aq (Mixter, Am. 8, 343).—SrHA'₃ 3½aq: long thin orystals. [67°-75°].—SrA'_26aq. Crystals.—Ba,Mg,A'₁₈ 12aq (Fitz, B. 13, 1312; 14, 1085).—Mg,Pb,A'₁₈ 12aq.—Ca,Pb,A'₁₈ 12aq: monometric crystals.—BaCa₂A'₆: monometric octahedra. SrCa, A'₆.—MgA'₂ aq.—Ba₆A'₁₀Ac₂ (Fitz, B. 11, 1897). — PbA'₂. [100°]. Crystalline mass.—Pb, A'₆O₄. S. c. 11 at 14°.—Pb₂A'₂O. S. 8·8 at 20°; 6·3 at 85°.—CuA'₂aq; green prisms.—Cu, A'₂O aq. - * CdA'2. V. e. sol. water. — ZnA'2 aq: needles. - $^{\text{L}}$ CdA'₂. V. 6. sol. water. - $^{\text{L}}$ ZnA'₂ aq: needles. S. (of ZnA'₂) 32 at 15°. - $^{\text{L}}$ HgA'₂. [110°]. S. 19·2 at 15°. - $^{\text{L}}$ Hg₂A'₂. [225°]. S. 1·4 in the cold. - NiA'₂ 2aq. - $^{\text{L}}$ CoA'₂ 3aq. - $^{\text{L}}$ FeA'₃: pale-green crystals. - $^{\text{L}}$ Fe₂A'₃. - $^{\text{L}}$ A'₄O: white pp. - $^{\text{L}}$ Crystals. S. 9 at 18.

Methyl ether MeA'. (80°). S.G. 2 .9373 (Elsässer, A. 218, 313); $\frac{9}{0}$ ·9403 (Gartenmeister). C.E. (0°-10°) ·001296. $\mu_{\rm D} = 1$ ·3812. H.F.p. 107,830. H.f.v. 105,010 (Thomsen, Th.). S.V. 104·6.

Ethyl ether EtA'. Mol. w. 102. (99° cor.). S.G. 2 9124 (E.); ½5 958; ½5 8688. M.M. 5-452 at 15-7° (Perkin). C.E. (0°-10°) 001215. S.V. 127.7 (R. Schiff, A. 220, 110). Formed from AgA' and EtI and also by distilling the acid with alcohol and H2SO4. Oil.

Propyl ether PrA'. (122° cor.). S.G. 2 9019 (Elsässer. A. 218, 821); \frac{15}{5} \cdot 868; \frac{25}{25} \cdot 8781 (Perkin, C. J. 45, 497); \frac{2}{6} \cdot 9023 (Gartenmeister). M.M. 6429 at 21.8°. C.E. (0°-10°) \cdot 001052. S.V. 150.

Isopropyl ether PrA'. (110° cor.). S.G. 9831 (Pribram a. Handl, M. 2, 687); \(\frac{15}{15}\) \(\frac{8}{15}\) \(\frac{15}{15}\) \(\frac{15}{15}\) (110° cor.). S.G.

25 ·862C. M.M. 6·595 at 14°.

(145°). S.G. 2 n Butyl ether C,H,A'. ·8953. S.V. 173·2. C.E. (0°-10°) ·00106 (Gartenmeister, A. 233, 265).

Isobutyl ether PrCH,A'. (137°). S.G. P :8876. C.E. (0°-10°) :001015 (E.). S.V. 174

(R. Schiff, A. 220, 332)

Isoamyl ether C₃H₁₁A'. Mol. w. 144. (160°). S.G. ² ·8877 (Elsässer, A. 218, 330). C.E. (0°-10°) 00099. S.V. 197 (Schiff); 195

Heptyl ether C,H,A'. (208°). S.G. 8 ·8847. C.E. (0°-10°) ·00096. S.V. 247.1 (Gar-

tenmeister).

Octylether C₈H₁₅A'. (226°). S.G. § 8833. C.E. (0°-10°) 00093. S.V. 270.5 (Gartenmeister). Ethylene ether $C_2H_4A'_2$. (2 S.G. $\frac{15}{18}$ 1.0544 (Perkin, C. J. 45, 505).

Phenyl ether PhA'. (200° i.V.) at 744 mm

Chloride C.H., COCl. (80°). S.G. 20 1.0646 (Brühl, A. 203, 14; cf. Sestini, Bl. [2] 11, 470). $\mu_{\rm p} = 1.4107$. $R_{\infty} = 34.12$. ZnMe₂, followed by water, forms methyl ethyl ketone and oily C₂H₁₄O (168°). S.G. ² ·877 (Pawloff, A. 188, 138).

Bromide C₃H₃OBr. (104° i.V.). S.G. ⁹⁻⁸
1·52 (Lobry de Bruyn, R. T. C. 3, 389).

Iodide C₃H₄OI. (128°).

Anhydride (C₃H₂O)₂O. Mol. w. 130. (162°)
at 708 mm. (L.); (165°) (Limpricht a. Uslar, A.
94, 322); (167°) (Kahlbaum, B. 16, 2481); (169°)
(Perkin, C. J. 28, 10). Formed by the action of Pb(NO)₂, (also of nitrates of other heavy of Pb(NO_s)₂ (also of nitrates of other heavy metals) upon propionyl chloride; PbCl2 is formed, and N₂O₄ and oxygen evolved; the yield is 77 p.c. of theoretical (Lachowicz, B. 18, 2991). Reacts with hydroxylamine hydrochloride, forming CH3.CH2.C(OH):NOH [85°] (Miolati, B. 25, 700).

Amide C,H,NO, i.e. C,H,CO.NH, Mol. w. 73. [79°]. (213°). H.C.p. 436,000. H.C.v. 435,600. H.F. 88,400 (Berthelot, C. R. 111, 145; A. Ch. [6] 22, 20; Bl. [3] 4, 229). Formed from propionic ether and NH₃ (Dumas, C. R. 25, 657; Sestini, Cimento, 4, 21; L. Meyer, B. 22, 24). Prepared by heating ammonium propionate at 230° under pressure; the yield being 55 p.c. of the theoretical amount (Hofmann, B. 15, 981). Got also by heating propionic acid (46 g.) with ammonium sulphocyanide (18 g.) for 4 days, and distilling the product (J. Schulze, J. pr. [2] 27, 517).—B'2HCl. Needles, v. e. sol. water, v. sl. sol. ether. - Hg(NH.C₃H₇O)₂. Large dimetric plates, sl. sol. cold water.

C,H,O.NHBr. [80° Bromo-amide Formed from propionamide, bromine, and KOHAq (Hofmann, B. 15, 753). Colourless needles, sol. alcohol. Converted by hot KOHAq into ethylamine. — C₂H₂O.NNaBr₂. [c. 100°]. Needles.

[204°] being formed at the same time (Pinner, B. 22, 1604).

Imide (C₂H₃O)₂NH. [154⁴] (Otto a. Tröger, B. 23, 759). (210°-220°). Formation.—1. By heating the amide in a current of HCl.-2. By heating the amide with the chloride and AlCl ..-3. By the action of AlCl, at 100° on a mixture of propionitrile with AcCl or propionyl chloride .-4. By heating propionic acid with propionitrile for some hours at 200°. Properties.—Rectangular tables or silky needles, sl. sol. cold water. May be sublimed.

Anilide C.H.ONHPh. [105°]. S. 42 at 24°. Formed by heating the amide with aniline (Kelbe, B. 16, 1200). Plates, sol. ether. With oxalic ether, benzene, and NaOEt it forms

CO.CO NPh [192°] (Wislicenus a. Sattler,

B. 24, 1256).

Methyl-anilide C₃H₅O.NPhMe. [58.5°]. Formed from the anilide and MeI (Norton a. Allen, B. 18, 1998).

Ethyl - p - toluideC₃H₅O.NHC₆H₄Me. (c. 270°). From the chloride and ethyl-toluidine

(Norton a. Livermore, B. 20, 2271).

Nitrile C.H.N, i.e. Et.CN. Propionitrile. Ethyl cyanide. (97°) . S.G. $\frac{4}{2}$ $^\circ$ 1998; $\frac{25}{25}$ $^\circ$ 7815 (Perkin, O. J. 55, 701); $\frac{9}{2}$ $^\circ$ 8010 (Thorpe). C.E. $(0^\circ$ - $10^\circ)$ $^\circ$ 00125; $(0^\circ$ - $100^\circ)$ $^\circ$ 00148. M.M. 8 $^\circ$ 331. μ_D 1 $^\circ$ 3659 at 24° (Gladstone, O. J. 59, 290). H.F.p. -9,670 (Thomsen, Th.); +8,700 (from diamond) (Berthelot, C. R. 108, 1219). H.F.v. -10,830. H.C. 446,700 (Berthelot); 471,450 (Thomsen, Th. 4, 129). S.V. 78.4. Formation.—1. By distilling dry KCy with KEtSO, or Ba(EtSO,)2 (Pelouze, A. 10, 249).—2. By boiling KCy with EtI and alcohol (Williamson, P. M. [4] 6, 205; Buckton a. Hofmann, C. J. 9, 250; Rossi, A. 159, 79).—3. From the amide and P₂O₈ (Dumas, A. 64, 334).—4. From $ZnEt_2$ and cyanogen or CyCl (Gal. Z. 1868, 252; Frankland a. Graham, C. J. 37, 740).—5. By the action of Ac₂O on the oxim of propionic aldehyde in the cold (Dollfus, B. 25, 1915). Preparation.—The nitrile cannot be freed from alcohol by distillation, since the compound C₂H₂N3HOEt boils constantly at 79°, and is not decomposed by water or CaCl2. It is best, therefore, to prepare it from the amide. Properties.—Colourless liquid, v. sol. water, but separates on addition of NaCl or CaCl₂. Miscible with alcohol and ether. Combines with metallic chlorides, yielding (C₂H₄N)SbCl₃, C₃H₄NAuCl₃, C₄H₄NPtCl₄, (C₅H₄N)₂SnCl₄, and (C₄H₅N)₂TiCl₄ (Henke, **4.** 106, 280). Forms also (C,H,N),FeCy, 6aq, (C,H,N),FeCy,Et,Cl, 6aq (Buff, A. 91, 253), and (EtCy),PtCy, 2aq (Than, A. 107, 315). Reactions.—1. Potash forms pro-pionic acid. Dilute H,SO, acts in like manner. 2. Dry chlorine forms CH2. CCl2. CN (Otto, A. 116, 195).—3. Bromine forms C₂H₃NBr₂ [64°], converted by water into (C₂H₄BrO)₂NH (Engler, A. 142, 65).—4. Dry HCl slowly forms C₂H₂NCl [121°] (Gautier, A. 142, 289), while HBr gives C₃H₂NBr₂ [50°-55°] (Engler, A. 149, 807). H1 below 0° forms EtCl₂NH₂ crystallising in plates (Biltz, B. 25, 2542).—5. Sodium converts it on heating into the polymeric cyanethine (q. v.). Sodium added to its solution in dry ether gives Acetyl-amide C₂H₂O.NHAc. [82°]. (230°— off H and ethane, and forms a product which is 240°). Formed from propionamidine hydrochloride, NaUAc, and Ac₂O, a compound C₂H₁₂N₂ (Et.C(NH).CMeNa.CN?). This powder is converted by water into C_sH₁₀N₂ [48°] (258° uncor.), and by HClAq into C_sH₅NO or C₂H₅.CO.CHMeCN (193.5°). The white powder treated with MeI yields C,H₁₂N₂ (43°) (262°), whence HClAq forms liquid C,H₁₁NO (175°), S.G. 2°945, mol. w. 125 (obs.). EtI and HClAq acting on the white powder yield C₈H₁₈NO (195°), S.G. 2°943 (E. von Meyer, J. pr. [2] 38, 336; Hanriot a. Bourer B. C. 1911, R. 1911, 191 veault, C. R. 108, 1171; Bl. [3] 1, 551). The compounds C,H,1NO and C,H,1NO are converted by heating with HClAq in sealed tubes into ethyl isopropyl ketone and ethyl butyl ketone respectively.-6. A mixture of propionitrile and benzonitrile in ethereal solution is converted by Na, followed by water, into CPh(NH).CHMe.CN [97°], whence warm dilute HClAq forms oily C.H. CO.CHMe.CN (E. von Meyer, J. pr. [2] 39, 189).—7. Zinc and HClAq yield propylamine together with smaller quantities of di- and tripropylamine (Pisanello, G. 16, 446).—Ac₂O at 200° forms C₇H₁₄N₂O₈ [68°] (220°) (Gautier, Z. 1869, 127).

References.—Amido-, Bromo-, Bromo-nitro-, Chloro-, Chloro-bromo-, Iodo-, Nitro-, and Oxy-amido-propionic acid and propionitrile.

PROPIONIC ALDEHYDE C_3H_0O i.e. $CH_2.CH_2.CH_O.$ Mol. w. 58. (48°). S.G. $\frac{15}{25}$.7966 (Perkin, C.J. 45, 476); $\frac{20}{4}$.8065; $\frac{25}{25}$.7966 (Perkin, C.J. 45, 476); $\frac{20}{4}$.8066 (Brühl, A. 203, 13). S. 16 at 20°. MM. 3·332 at 13·6°. μ_{β} 1·368. $R_{\infty}=25\cdot42$. S.V. 75·4. H.F.p. 55,240. H.F.v. 53,790 (Thomsen). Formed by oxidation of propyl alcohol (Michaelson, A. 134, 69; Chancel, A. 151, 301; Lieben a. Zeisel, M. 4, 14) and by distilling calcium formate with calcium propionate (Rossi, C.R. 70, 129). Pungent liquid, sol. water. Readily oxidised by air. Reduces ammoniacal AgNO₃, forming a mirror. Readily dissolves in aqueous NaHSO₃. Decomposed by hot KOHAq. Coloured brownish-red by alkaline sodium nitroprusside, the colour being destroyed by HOAc (Von Bitto, A. 267, 376).

Reactions.—1. Reduced in aqueous solution by sodium-amalgam to n-propyl alcohol.—2. H₂S passed into its aqueous solution, acidified by HCl, forms oily (C₂H₆S)C₃H₆O (Alexejeff, B. 10, 1739).—3. Dry ammonia passed into propionic aldehyde dissolved in ligroïn cooled by a freezing mixture forms a white flocculent pp. of C₂H₆(OH)NH₂, which readily deliquesces, forming C₁₃H₁₉N₃ [74°], crystallising in triclinic tables, a:b:c = 1:1·185: 801, insol. water, sol. alcohol and ether (Waage, M. 4, 708).—4. PH₃ and HCl form (CH₃.CH₂.CHO),PH₄Cl [128°], while PH₃ and HBr give (C₃H₆O),PH₄Br [106°]. These compounds are decomposed by water (Messinger a. Engels, B. 21, 331). PH₄I at 0° forms, in like manner, (C₂H₆O),PH₄I [96°] (Girard, A. Ch. [6] 2, 24).—5. Mercaptan and ZnCl₂ give CH₃.CH₂.CH(SEt), (198°) (Fromm, A. 253, 150). 6. Propyl alcohol and HOAc at 100° form CHEt(OPr)₂ (166° cor.) (Schudel, M. 5, 247).

Oxim C₂H₅.CH:NOH. (131°). Formed from the aldehyde and aqueous hydroxylamine (Petraczek, B. 15, 2784). Liquid, sol. water. It is a syn-oxim, because it is converted by Ac₂O and ether in the cold into propionitrile (Dollfus, A. 25, 1915).

Phenyl-hydraside CHEt:N₂HPh (152° in vacuo) (Bischoff, B. 23, 1916).

References.—Bromo-, Chloro-, Iodo-, and Oxy- Propionic aldehyde.

PROPIONIC IMIDO-ETHYL ETHER

C₂H₅.C(NH).OEt. The hydrochloride, B'HCl, formed by passing gaseous HCl into a mixture of propionitrile and alcohol diluted with ether (\frac{1}{3}\text{ vol.}), crystallises in prisms, decomposing at about 92° into EtCl and propionamide (Pinner, B. 16. 1654).

PROPIONITRILE v. Nitrile of Propionic

PROPIONYL-ACETOPHENONE v. PHENYL-METHYLENE-ETHYL-DIKETONE.

PROPIONYL-BENZOIC ACID v. PHENYL ETHYL KETONE CARBOXYLIC ACID.

PROPIONYL BROMIDE v. Bromide of Propionic Acid.

PROPIONYL-BUTYRIC ACID

C2H5.CO.CMe2.CO2H.

Methylether MeA'. (188° uncor.). Formed by the action of MeOH and HCl on the nitrile C₂H₃.CO.CMe₂.CN (175°), which is got by the action of cold HCl upon C₂H₃.C(NH).CMe₂.CN [43°] which is made by the action of MeI on the product of the action of Na on propionitrile (Bouveault, C. R. 111, 531). Liquid.

PROPIONYL CHLORIDE v. Chloride of Pro-

PIONIC ACID.

PROPIONYL CYANIDE C4H5NO i.e.

CH₂CH₂CO.CN. (109°). Prepared by heating propionyl chloride (12 g.) with silver cyanide (8 g.) in closed tubes for 1 hour at 100°. The product is fractionally distilled (Claisen a. Moritz, C. J. 37, 692). HCl converts it into ethylglyoxylic acid (a, v.).

glyoxylic acid (q. v.).

Di-propionyl di-cyanide (Et.CO)₂C₂N₂. [59°].
(208°). V.D. 5·6 (calc. 5·7). Obtained in the preparation of the preceding (C. a. M.) and by the action of AgCy on propionyl bromide (Lobry de Bruyn, R. T. C. 3, 390). Prisms, sl. sol. water, sol. alcohol and ether.

PROPIONYL-CYANO-ACETIC ETHER v.

CYANO-ACETIC ACID.

PROPIONYL-MALONIC ETHER $C_{10}H_{14}O_3$ i.e. C_2H_3 .CO.CH($CO_2Et)_2$. (239°-242° uncor.). Formed by the action of propionyl chloride upon sodio-malonic ether (Lang, B. 20, 1326). Colourless liquid.

PROPIONYL-PHENOL v. Oxy-phenyl ethyl retone.

 α -PROPIONYL-PROPIONIC ACID $C_sH_{10}O_s$ i.e. CH_s - CH_2 -CO.CHMe. CO_2H .

Methyl ether C₂H₅·CO.CHMe.CO₂Me or C₂H₆·C(OH):CMe.CO₂Me. (187° cor.). S.G. ²²
1·07. Formed by the action of Na on methyl propionate (Pingel, A. 245, 84). Colourless liquid with aromatic odour. Converted by ammonia into C₇H₁₅NO₂ and a hygroscopic substance C₁₂H₁₇N₄O₄ (?) [82°]. Aniline forms oxymethyl-ethyl-quinoline [295°] (Bouveault, C. R. 111, 41). NaOHAq yields a mixture of di-ethyl ketone and methyl ethyl ketone. NaOEt followed by alkyl iodides yields the alkyl derivatives: C,H₁₁MeO₃ or CEt(OMe):CMe.CO₂Me [194°], C,H₁₁EtO₃ (208°), C,H₁₁PrO₃ (220°), and C,H₁₁(CH₂Pr)O₃ (231°), all being liquid.

Ethyl ether Eth'. (199°). S.G. ² ·995; 15 ·983. Formed by digesting propionic ether in Et O. with Na. (Oppending a Hollow Ethology.

Ethyl ether Eth'. (199°). S.G. 2 '995; 15 '983. Formed by digesting propionic ether in Et₂O with Na (Oppenheim a. Hellon, B. 10, 699; Israel, A. 231, 197). Gives no colour with FeCl₂. Ammonia forms C₈H₁₂NO₂ (195°), insol. water, and a hygroscopic substance [75°]. NaOHAq forms di-ethyl ketone. NaOEt and EtI yield propionic ether and CH₃.CHEt.CO₂H (Geuther, A. 239, 386). Na followed by EtI added to the solution in benzene forms C.H₃.C(OEt):CMe.CO₂Et (206°), S.G. ¹⁵ 966. Reduced by HI and P to CHMePr.CO₂H (193°). Nitrous acid yields CH₃.C(NOH).CO₂Et [94°] (Hantzsch a. Wohlbrück, B. 20, 1320).

A mide C₃H₅O.CHMe.CO.NH₂. [158°]. Formed by adding water to the product of the action of AlCl₃ on a mixture of propionyl chloride and propionitrile (Otto a. Tröger, B. 22, 1455).

Needles, sl. sol. cold water.

Nitrile C₂H₃.CO.CHMe.CN. (c. 188°). Formed by the action of cold conc. HClAq on the product of the action of Na on propionitrile (Meyer, J. pr. [2] 38, 339). Oil. Converted by ammonia into the imide [45°]. Hydroxylamine forms the oxim C.H₅.C(NOH).CHMe.CN [44°] (180° at 200 mm.) (Hanriot, C. R. 112, 796). Aniline yields amido-methyl-ethyl-quinoline (Hanriot a. Bouveault, Bl. [3] 1, 552).

Propionyl-propionic acid CH₃.CH₂.CO.CH₂.CH₂.CO₂H. [33°]. A product of the action of boiling water on di-bromo-hexoic acid (from hydrosorbic acid and Br) (Fittig, A. 268, 69). Crystalline, v. e. sol. water, alcohol,

and ether.—ČaA'₂1½qq: plates.—AgA'.

PROPIONYL - PROPIONIC ALDEHYDE

C.H., CO.CHMe.CHO. [c. 40°]. (165°). Formed
by adding a well-cooled mixture of formic ether
(11 g.) and di-ethyl ketone (13 g.) to NaOEt
(10·5 g.) suspended in dry ether (10 g.). After
12 hours the product is acidified and distilled
under 50 mm. pressure (Claisen a. Meyerowitz,
B. 22, 3273). Crystals, v. sol. water. Deliquesces
and turns brown in air. NH₃ forms deliquescent
crystals of (C_nH₁₀O₂)NH₃. Cu(OAc)₂ added to its
alcoholic solution forms Cu(C₀H₉O₂)₂ [168°].
Phenyl-hydrazine produces C₁₂H₁₄N₂ (283°).

PROPIOPHENONE v. PHENYL ETHYL KETONE.
PROPYL. The radicle C₄H₇. n-Propyl
CH₂.CH₂.CH₂ changes in some reactions to iso-

propyl (CH₂)₂CH and vice versa.

Pr changes to Pr: 1. When cumyl alcohol is reduced to n-cymene by boiling with zinc-dust (Kraut, A. 192, 225; Jacobsen, B. 12, 434).—2. When cumyl chloride is reduced to n-cymene by zinc and HCl (J. 1879, 369).—3. When nitro-cymylene chloride is reduced to n-cymidine by boiling with zinc and HCl (Widman, B. 15, 166; 19, 245).—4. When isopropyl-cinnamic acid is reduced by sodium-amalgam to n-cumyl-propionic acid.—5. When o-amido-isopropyl-cinnamic acid is reduced by sodium-amalgam to

(B. 3)-n-propyl-(Py. 3)-oxy-quinoline dihydride. Pr changes to Pr: 1. When n-cymene is oxidised to cuminic acid—(a) by passing through the organism, (b) by air and gaseous NaOH (B. 5, 749; 12, 512; 11, 369; 14, 1144).—2. When n-cymene sulphonic acid is oxidised to oxy-isopropyl-sulpho-benzoic acid by alkaline KMnO₄ (Meyer a. Boner, B. 14, 1136, 2391; A. 220, 6).—3 When n-cymene is oxidised to oxyisopropyl-benzoic acid by alkaline KMnO₄.—4. When thymol is oxidised by KOH fusion to the oxy-cuminic acid, which is obtained from m-nitro-cuminic acid by reduction and diazotisation (Barth, B. 11, 1571).—5. When carvacrol (which contains n-propyl, since by distillation with P₂S_a

it gives n-cymene) is oxidised by KOH fusion to the oxy-cuminic acid [93°], which is obtained from o-amido-cuminic acid by diazotisation (B. 6, 936; 9, 1061).—6. Since cuminol is always accompanied in plants by n-cymene, it is probably produced by oxidation of the latter.—7. When n-propyl bromide is boiled with AlBr₃ (Gustavson, C. J. 44, 565).

From the above may be deduced the general law: When a propyl group in a benzene derivative occurs in the para-position to Me, CHO, or CO₂H, it is predisposed to be NORMAL by Me, 150 by CHO or CO₂H (Widman, B. 19, 251, 273, 2769, 2781; cf. Fileti, G. 16, 300, 497).

Dipropyl v. Hexane.

DI-PRÔPYL-ACETAL v. ALDEHYDE.
PROPYL ACETATE v. Acetyl derivative of
PROPYL ALCOHOL.

PROPYL-ACETIC ACID v. VALERIC ACID.
Di-propyl-acetic acid v. Octoic acid.
PROPYL-ACETOACETIC ETHER v. vol. i.

p. 23. DI-PROPYL-ACETONE v. METHYL HEPTYL

PROPYL-ACETYLENE v. PENTINENE.
DI-PROPYL-ACRYLIC ACID v. ENNENOIG

n-PROPYL ALCOHOL C, H, O i.e. CH₃.CH₂.CH₂OH. Mol. w. 60. (97.4° i.V.). S.G. 9 *8177 (Zander, A. 224, 79); $\frac{15}{15}$ *8088; $\frac{25}{25}$ *8025 (Perkin); $\frac{20}{4}$ *8044 (Brühl, A. 203, 268). C.E. $(0^{\circ}-10^{\circ}) \cdot 00082$. M.M. 3.77 at 15.6° . $\mu_{\beta} 1.3901$. R_{∞} 28.0. R_{∞} 28.0. H.F.p. 82,000 (Berthelot); 65,690 (Thomsen). S.V. 81.3 (R. Schiff, A. 220, 101). S.H. '670 (Diaconoff, Bl. [2] 38, 172). Occurs in fusel oil, especially in that of wine (Chancel, C. R. 37, 410; 68, 659, 726; Pierre a. Puchot, C. R. 66, 302; 70, 406; Linnemann, A. 160, 195; Fittig, Z. [2] 4, 44; Ekman, Chem. Zeit. 12, 564). Thus 40 g. of it were found in 100 litres of old cognac (Ordonneau, C. R. 102, 217; Clandon a. Morin, C. R. 104, 1187; 105, 1019). A litre of potato-fusel oil was found to contain 30 c.c. of n-propyl alcohol, 150 c.c. of isopropyl alcohol, 115 c.c. of butyl alcohols, 335 c.c. of amyl alcohols, 125 c.c. of water, and 245 c.c. of other bodies (Rabuteau, C. R. 87, 501). Propyl alcohol also occurs, together with isopropyl alcohol, among the products of the lactic and butyric fermentation of sugar (Bouchardat, C. R. 78, 1145; Meyer a. Forster, B. 9, 535), and in the fermentation of glycerin (Fitz, B. 13, 36, 1311).

Formation.—1. By reducing propionic anhydride with sodium-amalgam (Linnemann, A. 148, 251; 160, 231; 161, 18).—2. By adding sodium-amalgam and dilute H₂SO₄ to a solution of propionic aldehyde (1 pt.) in water (18 pts.) (Rossi, C. R. 70, 129).—3. By heating allyl alcohol with KOH (Tollens, Z. [2] 7, 242).

Properties.—Liquid with alcoholic odour, miscible with water, but separated from the solution by CaCl. Does not yield isopropyl alcohol when heated in sealed tubes (Aronstein, R. T. C. 1, 134). When propyl alcohol (71.46 c.c.) is mixed with water (28.54 c.c.), contraction (1.85 c.c.) is observed (Ramsay a. Young C. J. Proc. 4 101).

Young, C. J. Proc. 4, 101).

Reactions.—1. Chromic acid oxidises it to propionic aldehyde and acid.—2. HBr and H₂SO₄ give propyl bromide, propylene bromide,

and C₂H₂Br₂ (Niemilovitch, M. 10, 820).—3. I and KOHAq form iodoform (Lieben, A. Suppl. 7, 280).—4. Fused ZnCl₂ forms propylene, isobutyl alcohol, CH₂:CMe₂, and CHMe:CHMe (Le Bel a. Greene, C. R. 89, 418).—5. Phosphoplatinous chloride followed by water forms crystalline P(OPr)₃PtCl₂ (Pomey, C. R. 104, 364).—6. Calcium chloride yields (C₂H₂O)₃CaCl₂ crystallising in needles (Göttig, B. 23, 181).

Metallic derivatives. KOPr (De Forcrand, C. R. 104, 68).—NaOPr2HOPr (Fröhlich, A. 202, 295). Ba₂(OPr)₂(OH)₂. Formed by heating propyl alcohol with BaO under pressure (Destrem, A. Ch. [5] 27, 32). Yields propylene, methane, and hydrogen when heated. The corresponding lime compound yields di-ethyl ketone when heated.—Al(OPr)₂. [c. 60°]. S.G. \$\frac{1}{2}\$ 1.026. Formed in small quantity from propyl alcohol, AlI₂, and Al (Gladstone a. Tribe, C. J. 39, 6). Isopropyl alcohol does not react in this way.

Acetyl derivative ProAc. Propyl acetate. (102°). S. 1.5 at 16° (Linnemann, A. 161, 30). S.G. 2° 9091 (E.); 2° 8856 (Brühl); 8° 9093 (Gartenmeister); 15° 8933; 25° 8840 (Perkin, C. J. 45, 421). C.E. (0°-10°) 001197 (Elsässer, A. 218, 320). S.V. 128°3. µs 1.3890. R_∞ 43°11. M.M. 5·487 at 15·2°. Liquid. Not changed into isopropyl acetate by heating in a sealed tube (Aronstein, R. T. C. 1, 134).

Benzoyl derivative BzOPr. (230° cor.). S.G. 12 1.032 (Linnemann).

Isopropyl alcohol (CH₃)₂CH.OH. (83° cor.). S.G. § '7996 (Zander); $\frac{5}{15}$ '7914; $\frac{25}{25}$ '7845 (Perkin); $\frac{2}{4}$ ° '7887 (Brühl, A. 203, 12). $\mu_{\mathcal{S}}$ = 1'3821. \hat{R}_{∞} 27'99. M.M. 4'019 at 17'9°. S.V. 81'7 (R. Schiff, A. 220, 331); 82'8 (Zander). C.E. (0°-10°) '00108. H.F.p. 71,000. H.F.v. 68,970 (Thomsen, Th.).

Formation.—1. By distilling PrO.SO₃H (made from propylene and H₂SO₄) with water (Berthelot, J. 1855, 611).—2. By reducing acetone or propylene oxide with sodium-amalgam (Friedel, A. 124, 327; Linnemann, A. 140, 178).—3. By heating a solution of n-propylamine nitrite (Linnemann, A. 161, 43; B. 10, 1111; Meyer, B. 9, 535).—4. By the action of sodium-amalgam on (CH₂Cl)₂CH.OH dissolved in wet ether (Buff, Z. [2] 4, 124).—5. From CH₂I.CH₂OH by successive treatment with ZnMe₂ and water (Butlerow a. Ossokin, A. 145, 257).

Preparation.—1. Acetone (1 vol.) is diluted with water (5 vols.) and sodium-amalgam is slowly added. The liquid is distillate (Linnemann, A. 136, 37).—2. By boiling isopropyl iodide (1 pt.) with water (10 pts.) and Pb(OH)₂ (Flavitzky, A. 175, 380).—3. By heating isopropyl iodide (1 pt.) with water (20 pts.) at 100° for 40

hours (Niederist, A. 186, 391).

Properties.—Liquid with peculiar odour, not solid at -20° . Inactive to light. Miscible with water, yielding on distillation the hydrate $(C_2H_0O)_2$ 2aq (80°) , S.G. 15 882. When K_2CO_2 is added to an aqueous solution of the alcohol a hydrate $(C_2H_0O)_2$ aq separates (Erlenmeyer, A. 126, 307). After drying over CuSO₄ the hydrate $(C_2H_0O)_2$ aq (81°) S.G. 15 800 is left. Dilute CrO₂ forms acetone. Bromine forms acetone and FrBr.

Acetyl derivative ProAc. Isopropyl

acetate. (91°). S.G. 2 ·917 (Friedel, 4. 124, 827; Pribram a. Handl, M. 2, 686).

Benzoyl derivative ProBz. (218°) (Silva, Bl. [2] 12, 225). Liquid. Split up by distillation into benzoic acid and propylene (Linnemann, A. 161, 51).

References.—Bromo-, Chloro-, Chloro-iodo-, and Iodo- Propyl alcohol.

PROPYL ALDEHYDE v. Propionic Alde-

PROPYL-DI-ALLYL-CARBINOL v. DEGINYL ALCOHOL.

PROPYLAMIDOACETIC ACID

NHPr.CH₂.CO₂H. Formed from CH₂Br.CO₂Et and NH₂Pr (Chancel, *Bl.* [3] 7, 409; *C. R.* 114, 756). Needles, v. sol. water and alcohol, insole ther.—(HA')₂H₂PtCl₈ aq.—CuA'₂2aq: crystals, v. sol. water and alcohol.

PROPYL-AMIDO-ISOPROPYL ALCOHOL C₆H₁₅NO i.s. CH₃.CH(OH).CH₂.NHC₃H₃. Formed by heating allyl-propyl-amine with H₂SO₄ (3½ pts.) at 185° (Liebermann a. Paal, B. 16, 532).—B'₂H₂PtCl₆ 2aq: nodules (from water).

Di-propyl-amido-isopropyl alcohol CH₂.CH(OH).CH₂.N(C₃H₇)₂. Formed in like manner from allyl-di-propylamine.—B'₂H₂PtCl₆.

n-PROPYLAMINE C_3H_9N i.e. $CH_3.CH_2.CH_2.NH_2.$ Mol. w. 59. (49°). S.G. $\frac{15}{25}$ ·7222; $\frac{25}{5}$ ·7144. H.F.p. 22,760. H.F.v. 20,420 (Thomsen, *Th.*). S.V.85·6. M.M. 4·564 (Perkin, *C. J.* 55, 693). $μ_D = 1·3873$ (Gladstone, *C. J.* 59, 290).

Formation.—1. By reduction of propionitrile. Using 36 g. of propionitrile, 500 g. alcohol, 200 g. water, zinc, and 500 g. of 20 p.c. HClAq, the yield is 9 g. (Mendius, A. 121, 129; Linnemann, A. 161, 44).—2. From n-propyl iodide by successive treatment with silver cyanate and KOHAq (Silva, Z. [2] 5, 638).—3. By heating n-propyl iodide with alcoholic NH₂, and distilling the product with potash. The fraction boiling below 78° is mixed with alcohol and oxalic ether, and the di-propyl-oxamide [161.5°] thus formed distilled with alcoholic potash (Vincent, C. R. 103, 208). n-Propyl iodide heated with conc. NH,Aq for 13 hours at 100°-150° forms propylamine hydriodide and NPr. I (Malbot, A. Ch. [6] 13, 480, 546).—4. When n-propyl chloride is heated with aqueous ammonia in equimolecular proportions at 140°-165°, one-fifth of the product is n-propylamine and is nearly all contained in the lower layer. The upper layer consists of free di- and tri- propyl-amines, water, and alcohol (Malbot, C. R. 104, 998).-5. By heating n-propyl nitrate with alcoholic NH, at 100 (Wallach, A. 214, 311).

Preparation.—1. A mixture of equimolecular quantities of bromine and butyramide is run into an excess of a 10 p.c. solution of KOH at 60° and the propylamine distilled off; the yield is 80–90 p.c. (Hofmann, B. 15, 769). It may be dried over CaO and afterwards over Na.—2. PrCl (1 mol.) heated with aqueous NH₂ (1 mol.), and enough alcohol to effect solution, in sealed tubes at 110° forms 45 p.c. propylamine, 35 p.c. dipropylamine, and 20 p.c. tri-propyl-amine (Chancel, Bl. [3] 7, 405). The product is neutralised with HCl, the alcohol distilled off, cooled, filtered from separated NH₂Cl, the syrup mixed with NaOH, distilled, and fractionated.

The bases may be further purified by means of | their oxalates.

Properties .- Mobile liquid with ammoniacal smell; mixes with water, becoming hot. When wet it boils at 78°-82°. Its aqueous solution ppts. salts of Cu and Fe, the pp. not dissolving in excess, but with AgNO, it gives a pp. sol. excess.

Reactions.—1. Chromic acid yields propionic acid (Chapman a. Thorpe, A. 142, 176).—2. Nitrous acid forms n- and iso- propyl alcohols and propylene (Meyer a. Forster, B. 9, 535).—3. Benzoic aldehyde forms C.H. CH:NPr (210°) (Zaunschirm, A. 245, 282).—4. Oxalic ether added to a mixture of equal volumes of propylamine and water forms CONHPr.CONHPr and CO₂H.CONHPr [110°], which yields CaA'₂ 2aq. S. 1.38 at 17°

Salts .-- B'HCl. [158°] (Topsoë, J. 1883, 621).—B'HHg₂Cl₈. — B'HHg₈Cl₁₁.—B'HAuCl₄. B',H,PtCl₃. Monoclinic crystals. — Oxalate. NH,Pr C₂H₂O₄ jaq. Plates. S. 14·8 at 18°; S. (alcohol) ·6 at 18°.—Picrate [135°]. Nitramine CH₃.CH₂.CH₂.NH.NO₂. (129° at 40 mm.). S.G. ¹⁵ 1·102. Formed by heating

PrN(NO2).CO2Et in a current of dry NH3, and decomposing the product with dilute H₂SO₄ (Thomas, R. T. C. 9, 75). Liquid, crystallising at -23°. Decomposes at 142°. Sl. sol. water; decomposes carbonates.-PrNK.NO2. Formed by dissolving the nitramine in alcoholic potash and ppg. with ether. Small pearly plates, forming an alkaline solution in water.—PrNAg.NO2. Slender needles, blackening in light. yields N₂O.

Isopropylamine (CH₃)₂CH.NH₂. (32°). S.G.

18 ·690.

Formation.—1. By the action of HClAq on isopropyl-carbamine (Siersch, A. 148, 263; Gautier, C. R. 67, 723).—2. By running a mixture of bromine (1 mol.) and isobutyramide (1 mol.) into an excess of a 10 p.c. solution of KOH at 60°; the yield being 65 p.c. (Hofmann, B. 15, 768).—8. By reducing acetoxim dissolved in alcohol with sodium-amalgam and HOAc at 40° (Goldschmidt, B. 20, 728).—4. By heating isopropyl iodide with conc. NH₃Aq (Malbot, C. R. 111, 650; cf. Jahn, M. 3, 166).

Preparation.—By reduction of a solution of acetone-phenyl-hydrazide (1 pt.) in 96 p.c. alcohol (10 pts.) by means of acetic acid (2½ pts.) and sufficient 67 p.c. sodium-amalgam, the temperature being kept below 25°; the yield is 70 p.c. of the theoretical (Tafel, B. 19, 1926).

Properties.—Mobile liquid, with ammoniacal smell, miscible with water. Converted by nitrous

salts. — B'HCl. [1 [140°]. Deliquescent

cubes.—B',H,PtCl_s: orange plates, m. sol. Aq.
Nitramine PrNH.NO_r [-4°]. (91 (91°) S.G. 15 1.098. Formed from PrN(NO2).CO2Et by successive treatment with dry NH, and dilute H₂SO₄ (Thomas, R. T. C. 9, 78). Needles. -PrNK.NO. Slender hygroscopic needles. PrNAg.NO. Thin plates. Converted by PrI into PrNPr.NO. (67° at 10 mm.).

Di-n-propylamine Pr.NH. (110°) (P.; C.); (98°) (V.). S.G. $\frac{12}{3}$ ·7430; $\frac{25}{35}$ ·7357 (Perkin, C. J. 55, 693). M.M. 7, 549. $\mu_T = 1$ ·4083 (Gladstone, C. J. 59, 290). A product of the action of alcoholic NH, on PrI at 100° (Vincent, C. R. 103, 208;

Bl. [2] 46, 287) or of aqueous NH, on PrCl at 190° (Malbot, C. R. 105, 575). Purified by means of its nitrosamine. Liquid with ammoniacal smell, sl. sol. water. The pps. that it forms with salts of Ag and Al are sol. excess; those with salts of Cu and Zn are insol. excess.

Salts.—B'H₂C₂O₄. Flat needles. S. 5 at 17°. S. (alcohol) '33 at 18° (Chancel, *Bl.* [3] 7, 406).—Picrate. [75°]. S. 2·2 at 19°. Melts under water at 45°.

Nitrosamine Pr.N.NO. (206°). S.G. 9 931. Liquid smelling like hay. Decomposed by HClAq, with formation of dipropylamine.

Nitramine Pr.N.NO. A liquid, got from Prnk.NO₂ and PrBr (Thomas, R. T. C. 9, 79).

Di-isopropylamine Pr₂NH. (84°). S.G. 22 ·722. A product of the action of alcoholic NH. on PrI at 100° (Van der Zande, R. T. C. 8, 202; cf. Siersch, Z. [2] 5, 145). Liquid with very B'HNO₂. [188°].—B'₂H₂PtCl₈: orange tables.

Nitrosamine Pr₂N.NO₂. [46°]. (195°).

Nitramine Pr₂N.NO₂. (56° at 10 mm.).

Formed from PrNAg.NO2 and PrI (Thomas,

R. T. C. 9, 82). Liquid.

Tri-n-propylamine Pr,N. (157°). S.G. $\frac{9}{0}$ '770; $\frac{15}{5}$ '760; $\frac{9}{25}$ '754 (Perkin). S.V. 222·1 (Zander, A. 214, 171). M.M. 11·664. $\mu_{\rm D}=1$ ·4171 (Gladstone, C. J. 59, 290). C.E. (0°-10°) ·00105. A product of the action of PrI on alcoholic NH, It is the chief product got by heating PrCl with conc. NH, Aq (Malbot, A. Ch. [6] 13, 480, 546). Purified by means of its picrate (Chancel, Bl. [8] 7, 407). Liquid with faintly ammoniacal odour, almost insol. water. Picrate. S. 3 at 18°; S. (94 p.c. alcohol) 4.4 at 19°.

Propylo-iodide Pr.NI. The chief product of the action of PrI on NH, Aq at 150°-190°. Prisms. Yields (Pr.NCl), PtCl, and deliquescent Pr. NOH, which is decomposed by heat into Pr.N, propylene, and water (Roemer, B. 6,

References .- Bromo-, Chloro-, Chloro-1000-, and Oxy- PROPYLAMINE.

PROPYL-ISOAMYL-GLYOXALINE

C₈H₁(C₈H₁₁)N₂. Oxalisoamyl-butyline. (251°). S.G. 12 '920. Formed from propyl-glyoxaline and C₅H₁₁I (Rieger, M. 9, 609).

PROPYL ISOAMYL OXIDE Pr.O.C,H,,. (125°-130°) (Chancel, A. 151, 305).

PROPYL-ANILINE CoH13N i.e. PhNHPr. (222° i.V.) (Pictet a. Crépieux, B. 21, 1111). S.G. 4 949. Formed by heating aniline with PrBr (Claus a. Roques, B. 16, 909; Wacker, A. 243, 290). Liquid. Salts.—*B'HCl. [150].—
*B'₂H₂C₂O₄. [152°]. Large tables, v. sol. water.
Formyl derivative PhNPr(CHO). (269°

i.V.). S.G. 16 1.044. Got from formanilide and

n-propyl bromide.

derivative PhNPrAc. Acetyl[48°]. (269° i.V.). Hexagonal plates (from ligroïn).

Nitrosamine. Yellow oil.

Isopropyl-aniline PhNHPr (213° i.V.)

Liquid. Yields an oily nitrosamine. B'2H2PtCl. Red prisms (P. a. C.).

Formy i derivative PhNPr.NO. (264°i.V.).

Formed from formanilide and PrBr.

Acetyl derivative PhNPrAc. [39°]. (265° i.V.). Transparent leaflets (from ligroIn).

Di-n-propyl-aniline NPhPr₂. (245° i.V.). S.G. ²₆ 9240. S.V. 243·1 (Zander, A. 214, 168). C.E. (0°-10°) .00081. Formed from aniline and PrI (Lippmann a. Fleissner, M. 3, 711).-B'2H2PtCl6. Decomposed by water.

Di-isopropyl-aniline NPhPr₂. (220° i.V.). S.G. ^a ·9338. S.V. 235·4 (Zander). C.E. (0°-10°) ·00087.

References .- NITRO- and NITROSO- PROPYL-ANILINE.

DI-PROPYL-ANTHRACENE DIHYDRIDE

C₂₀H₂₁ i.e. C₆H₁< CPr₂ C₆H₄. Formed from dipropyl-anthrone, HI, and P at 140°-170° (Hallgarten, B. 22, 1070). Plates, getting soft at 47°. Di-isopropyl-anthracene dihydride

 $C_0H_3Pr < CH_2 > C_0H_3Pr$. (above 360°). $[90^{\circ}].$ Formed by boiling C₆H₄Pr.CH₂Cl alone or with ZnCl₂ (Errera, G. 14, 280). Amorphous yellow powder, insol. alcohol, sol. ether. Its solutions are red with green fluorescence.

PROPYL-ANTHRANOL. Propyl derivative. $C_{20}H_{22}O$ i.e. $C_{e}H_{4} < \stackrel{CPr}{C(OPr)} > C_{e}H_{4}$. [72°]. Formed by boiling anthranol with KOHAq and PrI (Hallgarten, B. 22, 1070). ether and alcohol, insol. alkalis. Needles, sol. Chromic acid solution converts it in the cold into propyloxanthranol $C_{17}H_{16}O_2$ [164°], v. sol. benzene.

DI-PROPYL - ANTHRONE C, H, O C_oH₄<CO_{CPr₂}>C_oH₄. [124°]. Formed, together with the preceding body, by boiling anthranol (5 g.) with KOH (5 g.), water (25 c.c.), and PrI (18 g.) (Hallgarten, B. 22, 1069). Trimetric crystals, sol. hot alcohol. CrO, in HOAc oxidises it to anthraquinone.

TRI-PROPYL-ARSINE v. vol. i. p. 319. PROPYL-BENZENE v. CUMENE.

Di-n-propyl-benzene C_eH₄Pr₂[1:4]. (219°). Formed from p-di-bromo-benzene, n-propyl bromide, and Na (Körner, B. 11, 1863; A. 216, 223). Formed also, together with the m-isomeride, by the action of AlCl, and HCl on n-cumene (Heise a. Töhl, A. 270, 165). Liquid, volatile with steam. Bromine forms C₆H₂Br₂Pr₂ [48°]. n-iso-Di-propyl-benzene C₆H₄Pr₂Pr[1:4].

(212° cor.). S.G. 2 ·8713. Formed from C,H,Pr.CH,Cl and ZnEt, (Paterno a. Spica, B. 10, 1746) and from PrBr, p-bromo-cumene, and Na. Formed also, together with the m- isomeride, by the action of AlCl, at -2° on a mixture of n-cumene and PrBr (Heise, B. 24, 772).

m-Di-isopropyl-benzene $C_6H_4Pr_2[1:3]$. (204°). Formed, together with the p-isomeride, by the action of propyl or isopropyl chloride on benzene in presence of AlCl. (Silva, Bl. [2] 43, 320; Uhlhorn, B. 23, 3142) and by the action of AlCl. and HCl at 100° on cumene (H. a. T.). Yields isophthalic acid on oxidation.

p-Di-isopropyl-benzene C₆H₄Pr₂[1:4]. (c. 202°). Formed as above. Yields terephthalic acid on oxidation.

PROPYL-BENZENE SULPHONIC ACID v. CUMENE SULPHONIC ACID.

p-Di-n-propyl-benzene sulphonic acid C₆H₂Pr₂·SO₅H. [62°]. Formed from di-propylbenzene and fuming H₂SO₄ (Körner, B. 11, 1865; A. 216, 224; Remsen, Am. 5, 162). Deliquescent needles, with pearly lustre.—NaA'4aq.—KA'4aq. -BaA', 1aq.-BaA', 2aq.-CaA', 9aq.-PbA', aq. Amide C₆H₃Pr₂SO₂NH₂. [103°]. Hexagonal

crystals (from alcohol), sl. sol. hot water.

n-iso-Di-propyl-benzene sulphonic acid [1:4:x] C₈H₈PrPr.SO₃H. [60°]. Needles (Heise, B. 24, 771).—BaA'₂ xaq.—PbA'₂ aq: minute needles, m. sol. cold water.—PbA'₂ 8aq: needles. Amide. [96°]. Needles.

m-Di-isopropyl benzene sulphonic acid $C_0H_3Pr_2.SO_3H[1:3:x]$. Formed by shaking with H_2SO_4 ($1\frac{1}{2}$ pts.) the fraction (200°-210°) got by the action of AlCl, on a mixture of PrCl and benzene, and separated from the accompanying p-di-isopropyl-benzene sulphonic acid by means of its Ba salt (Uhlhorn, B. 23, 3142). -BaA'22aq: needles, sl. sol. water. $-\text{CuA}'_2$ 4 aq. $-\text{MgA}'_2$ 4 aq. $A \ mid \ c \ C_6 H_3 \text{Fr}_2 \cdot \text{SO}_2 \text{NH}_2$. [145°].

p-Di-isopropyl-benzene sulphonic acid $C_6H_3Pr_2(SO_3H)$ [1:4:2]. Formed as above.--BaA'₂: nodules, v. sol. water.—CuA'₂6½aq. $Amide\ C_{b}H_{3}Pr_{2}.SO_{2}NH_{2}.$ [102°].

m-Di-n-propyl-benzene disulphonic acid $C_6H_2Pr_2(SO_3H)_2$. Deliquescent tables (Heise).— $K_2A''x$ aq.—BaA'' $1\frac{1}{2}$ aq: tables, v. sol. water.— PbA" 13aq: tables, v. e. sol. warm water.

Amide. [195°]. Needles.

PROPYL-BENZOIC ACID v. CUMINIC ACID. PROPYL-BENZOYL-ACETIC ACID v. Benz-OYL-ACETIC ACID and PHENYL BUTYL KETONE CARBOXYLIC ACID.

PROPYL-BENZYL- v. CUMINYL-. n-PROPYL BORATE (PrO),B. (172°-175°). S.G. 16 867 (Cahours, C. R. 76, 1383).

Isopropyl borate (PrO)₃B. (140° cor.). Formed by heating B₂O₈ with isopropyl alcohol at 120° (Councier, J. pr. [2] 18, 389). Mobile oil, slowly saponified by water.

n-PROPYL BROMIDE PrBr. (71°). S.G. 9 1·3835 (Zander, A. 214, 159); $\frac{20}{1}$ 1·3520 (Brühl, A. 203, 13); $\frac{1}{15}$ 1·3611; $\frac{25}{25}$ 1·3474 (Perkin). C.E. (0°-10°) ·00123. μ_{β} = 1·4406. R_{∞} = 38·20. S.V. 97. M.M. 6·885 at 19·2°. H.F.p. 30,850. H.F.v. 29,110 (Thomsen, Th.). Formed from n. propyl alcohol and HBr (Linnemann, A. 161, 40; Pierre a. Puchot, J. Ph. [4] 13, 9). When heated in sealed tubes at 280° it partially changes to isopropyl bromide (Aronstein, R. T. C. 1, 134). When heated for a short time with AlBr, it is completely changed to isopropyl bromide (Kekulé a. Schrötter, B. 12, 2279; Gustavson, J. R. 15, 61). Bromine forms propylene bromide. — C₃H,Br(H₂S)₂23aq (De Forcrand, **A.** Ch. [5] 28,

Isopropyl bromide PrBr. $(60^{\circ}).$ 13397 (Z.); \$\frac{29}{2}\$ 1-3097 (B.); \$\frac{15}{15}\$ 1-3198; \$\frac{25}{25}\$ 1-3052 (Perkin). C.E. (0^2-10^2) \cdot 00127, \$\mu_\theta\$ = 1-4317. \$\mathbb{R}_\tilde{\omega}\$ 38:58 (Br\(\omega\$hl)). M.M. 7-003 at 17-1°. S.V. 99. Formation.—1. From isopropyl alcohol and

HBr at 150° (Linnemann, A. 136, 41).—2. From n-propyl bromide and AlBr_s. —3. From propylene bromide and HI (Linnemann, A. 161, 57).

Preparation.—By adding bromine (65 g.) gradually to PrI (100 g.) in the cold; the yield being 45 g. (R. Meyer, J. pr. [2] 34, 105).

Properties.—Liquid, not attacked by Cl.

ISOPROPYL-BUTENYL-BENZENE v. Bu-TENYL-CUMENE.

n-PROPYL-ISOBUTYL-AMINE C,H,,N i.e. PrNH.C.H., (124°). Formed by reducing isobutyl-propargyl-amine in alcoholic solution with

sodium (Paal a. Heupel, B. 24, 3048).—B'HCl. [135°].—B'H₂C₂O₄. [224°]. Needles.

ISOPROPYL-BUTYL-AMINE DICARB-

OXYLIC ACID C.H., NO. i.e.

CO2H.CMe2.NH.CMe2.CH2.CO2H. A product of the oxidation of triacetonamine with chromic acid mixture (Heintz, A. 198, 69). Crystalline grains, m. sol. hot water. Yields amido-isobutyric acid when heated. — H₂A''HCl. –
KHA'' 2aq. — Zn(HA'')₂ 6aq. — ZnA''aq. —
CuA'' aq.—AgHA''.—Ag₂A''HNO₃ aq: grains.
PROPYL-ISOBUTYL-GLYOXALINE

CH.NC₁₀H₁₈N₂ i.e. CH.NPr C.C.H. Oxal-propyl-iso-(241° at 738 mm.). S.G. 18 .915. Formed from isobutyl-glyoxaline and PrBr (Radziszewski a. Szul, B. 17, 1295). Liquid.-B'2H2PtCla: minute orange prisms.

Propyl-butyl-glyoxaline CH.N:CPr CH.N(C,H₉) (243°) at 728 mm. S.G. 19 938. Formed from propyl-glyoxaline and butyl iodide. Oxidised by $\mathbf{H}_2\mathbf{O}_2$ to butyl-oxamide [198°] (Rieger, M. 9, 609). $-\mathbf{B'HZnCl_{s}} - \mathbf{B'_{2}H_{2}CdCl_{4}} - \mathbf{B'_{2}H_{2}PtCl_{6}}$

Propyl-isobutyl-glyoxaline CH.N:CPr CH.N(C,H,0)>. (232°) at 736 mm. S.G. 13 940. Formed from propyl-glyoxaline and isobutyl iodide (Rieger).

PROPYL ISOBUTYL KETONE Pr.CO.C.H. (155°) at 750 mm. S.G. 9 831. Formed from isovaleryl chloride and ZnPr₂ (Wagner, J. R. 16, 668). Liquid.

Isopropyl isobutyl ketone Pr.CO.CH, Pr. (160°). S.G. 14 ·865. Formed by oxidising secoctyl alcohol with cold dilute K2Cr2O, and H2SO4 (Williams, C. J. 35, 130). Liquid, not solid at -17°. Does not form a crystalline compound with NaHSO.

PROPYL-ISOBUTYL-MALONIC ACID

 C_4H_9 . $CPr(CO_2H)_2$. [129°]. This appears to be the composition of the acid got by saponifying C₄H₂.C(C₂H₃)(CO₂Et)₂ (249°), which is got from malonic ether, isobutyl iodide, allyl iodide, and NaOEt (Ballo, B. 14, 335).

PROPYL BUTYL OXIDE Pr.O.C, Hg. (117°). S.G. § ·7773. C.E. (0°-10°) ·00124. S.V. 174·4 (Dobriner, A. 243, 7).

ISOPROPYL-ISOBUTYL-PHOSPHINE PHPr.CH₂Pr. (140°). Formed from isopropylphosphine and isobutyl iodide at 130° (Hofmann,

B. 6, 300)

ISOPROPYL-ISOBUTYL-QUINOLINE

-CH:Ç₽r $C_0H_4 < N = C.CH_2Pr$. (296°) at 710 mm. Formed by the action of isovaleric aldehyde and conc. HClAq upon aniline (Spady, B. 17, 1718; 18, 3373).—B'₂H₂PtCl₈. Yellow needles, sl. sol. 8373).—B'2H2PtCl8. Yellow needles, sl. sol. water.—B'C8H2(NO2)8OH. Large yellow plates. B'HCl aq. — B'HNO, aq. — B'H₂SO. B'H2Cr2O,: orange needles, sol. hot water.

Methylo-iodide B'MeI aq. Yellow needles, v. sol. alcohol.—B', Me, PtCl,: yellowish-

red triclinic prisms.

PROPYL-TRICARBALLYLIC ACID [152°]. CHPr(CO₂H).CH(CO₂H).CH₂.CO₂H. Formed by saponifying its ether, which is made by the action of fumaric ether on sodium propylmalonic ether (Auwers, B. 24, 311, 2898). Prisms (containing xaq), v. e. sol. ether.

Ethyl ether Et.A". (206° at 16 mm.).

8.G. ¥ 1.092.

Isopropyl-tricarballylic acid

CHPr(CO₂H).CH(CO₂H).CH₂.CO₂H. [162°]. Prepared in like manner from isopropyl-malonic ether. Plates or prisms, v. sl. sol. ligroïn.

Ethylether Et, A'". (206° at 20 mm.).

S.G. 19 1.085.

PROPYL CARBAMATE NH2.CO.OPr. [52°] (195°). Formed by heating urea with propyl alcohol (Cahours, C. R. 76, 1387). Formed also from Cl.CO₂Pr and NII₃ (Roemer, B. 6, 1102). Prisms, v. e. sol. water and alcohol.

Isopropyl carbamate NH2.CO.OPr. [87°]. Formed from Cl.CO₂Pr and dry NH, (Spica a. Varda, G. 17, 165). Hygroscopic needles.

ISOPROPYL-CARBAMINE PINC. S.G. 2.760. Formed from isopropyl iodide and AgCy (Gautier, C. R. 67, 723; A. 149, 155). HClAq forms isopropyl-formamide.

PROPYL-CARBAMIC ACID NHPr.CO.H.

Methyl ether MeA'. (180° at 755 mm.). S.G. 15 992. Liquid (Thomas, R. T. C. 9, 71). The nitramine NPr(NO₂).CO₂Me. S.G. 15 1.187 is made from NAgPr(NO₂) and Cl.CO₂Me.

Ethyl ether EtA'. (186°). Formed from propylamine and chloro-formic ether (Schreiner,

J. pr. [2] 21, 125).

Isopropyl-carbamic acid NHPr.CO.H. Methyl ether MeA'. (165.5°). S.G. 15 ·981. Liquid. The nitramine

NPr(NO₂).CO₂Me S.G. 15 1.159 is also liquid. PROPYL-CARBINOL v. BUTYL ALCOHOL.

Di-propyl-carbinol v. HEPTYL ALCOHOL. DI-PROPYL-CARBOBENZONIC ACIDS

C20H22O2. Two acids of this formula [139°] and [90°] are formed by heating phonyl benzyl ketone with n-propyl alcohol and KOH (Zagumenny, A. 184, 166). The acid [139°] forms a di-nitro- derivative [176°]

PROPYL CARBONATE CO(OPr)2. cor.). S.G. 17 .949 (Roese, A. 205, 230)

Propyl orthocarbonate C(OPr)₄. (224° cor.). S.G. 9 911. Formed from NaOPr and chloro-

picrin (Roese, A. 205, 253)

n-PROPYL CHLORIDE PrCl. Mol. w. 783. 36,020. C.E. (0°-10°) 00137. Formed by the action of chlorine on propane and on PrI. Prepared by saturating n-propyl alcohol with gaseous HCl, and heating at 125° with conc. HClAq (Malbot, Bl. [3] 2, 136; cf. Pierre a. Puchot, A. Ch. [4] 20, 234). Acts with extreme slowness on alcoholic NH, (Vincent, Bl. [2] 45, 504). Not changed to PrCl by heating in sealed tubes (Aronstein, R. T. C. 1, 134).—PrCl(H₂S)₂ 23aq (De Forcrand, A. Ch. [5] 28, 34).

Isopropyl chloride PrCl. (36°). S.G. § 8825 (Z.); ½ 8688; ½ 8575 (P.). M.M. 5 159 at 17 2°. S.V. 94. C.E. (0°-10°) 00142. Formed from isopropyl alcohol and HCl. Chlorine converts it into CH₃.CCl₂.CH₃ and CH₃.CHCl.CH₂Cl. Partially converted into NH2Pr by heating with conc. NH₃Aq at 140° (Malbot, C. R. 111, 650).

ISOPROPYL CINNAMIC ACID v. CUMYL-

ACRYLIC ACID.

References .- AMIDO-, CHLORO-, and NITRO-PROPYL-CINNAMIC ACIDS

PROPYL-m-CRESOL C.H.MePr.OH. (230°-235° at 734 mm.). Formed, together with its

propyl ether and di-propyl-m-cresol, by heating | m-cresol with propyl alcohol and MgCl2 (Mazzara, G. 12, 187, 332). Liquid, sl. sol. water, sol. alcohol and ether. Not solid at -15° . Not coloured by FeCl₃. Yields a crystalline nitrosoderivative [140°].

Methyl ether MeA'. (226°) at 740 mm. Propyl ether PrA'. (235°-240°).

Acetyl derivative AcA'. (240°) at 743

Isopropyl-m-cresol C₆H₃PrMe.OH. (238°). Formed in like manner by heating m-cresol with isopropyl alcohol and MgCl2 (Mazzara, G. 12, 505). Oil. Yields $C_6H_2(NO)$ PrMe.OH [167°].

Methyl ether MeA'. (215°-220°). Isopropyl ether PrA'. (230°-235°).

Isomeride v. THYMOL.

Propyl-o-cresol v. Carvacrol and Cymenol. Isopropyl-cresol CaHaPrMe.OH. (229° cor.). S.G. o 1.0012. Formed by potash-fusion from the m-isocymene derived from camphor (Spica, G. 12, 552). Not coloured by FeCl.

Ethyl ether EtA'. (228° cor.). Di-propyl-m-cresol C_eH₂Pr₂Me.OH. (250°). Formed as above. Liquid, yielding a liquid acetyl derivative (255°-260°).

 $C_6H_2Pr_2Me(OH)$. Di-isopropyl-m-cresol (251°). Formed at the same time as isopropylm-cresol (M.). Liquid smelling like phenol, sl. sol. water, insol. dilute potash.

Methyl ether MeA'. (243°).

Acetyl derivative AcA'. (255°-260°)

ISOPŘOPYL-CUMIDINE C.H.Pr.NHPr. (245°-250°). A product of the action of aniline and ZnCl₂ on isopropyl alcohol (Louis, B. 16, 111). Liquid. Forms a granular picrate.

ISOPROPYL CYANATE v. vol. ii. p. 315. PROPYL CYANIDE v. Nitrile of BUTYRIO

PROPYLENE C.H. Mol. w. 42. S. ·44 0°: ·22 at 20° (von Than, A. 123, 187). Mol. w. 42. S. ·44 at (alcohol) 13. S. (H,SO,) 200 (Berthelot). H.F.p.

3,220. H.F.v. 2,060 (Thomsen, Th.).

Formation.—1. By passing fusel oil, valeric acid, or light petroleum through a red-hot tube (J. W. Reynolds, A. 77, 118; Prunier, J. 1873 847).-2. By distilling calcium oxalate with KOAc (Dusart, A. 97, 127) .- 3. By the action of mercury (containing a trace of zinc) and fuming HClAq on allyl iodide (Berthelot a. De Luca, A. 92, 306).—4. Mixed with H ($\frac{1}{5}$ vol.), by reducing allyl iodide in alcoholic solution with granulated zinc and HClAq (Tollens a. Henninger, A. 156, 156).—5. From allyl iodide and HI (Butlerow, A. 145, 271).—6. A product of the action of ZnEt₂ on CCl₄ (Rieth a. Beilstein, A. 124, 242) on allyl iodide (Wurtz, Bl. 5, 51) and on CHBr, (Beilstein, Bl. [2] 2, 51).—7. By boiling n- or iso-propyl iodide with alcoholic potash (Freund, M. 8, 633; Erlenmeyer, A. 139, 228).—8. Together with other hydrocarbons, by treating propylene bromide with water and sodium-amalgam or with zinc and HClAq (Prunier, C. R. 76, 98).—9. By heating CCl₂Me₂ or CBr₂Me₂ with Na at 140° (Friedel a. Ladenburg, Z. 1868, 48; Reboul, A. Ch. [5] 14, 488).—10. Together with cresol, by heating thymol with P_2O_8 (Engelhardt a. Latschinoff, Z. 1869, 616).—11. In the manufacture of oil-gas (Armstrong, C. J. 49, 74).—12. Together with ethylene, by heating allyl alcohol with P.O. (Béhal, A. Ch. [6] 16, 860).— 13. By heating N(C₃H₇), with C₃H₇Cl at 190°

(Malbot, A. Ch. [6] 13, 546).

Preparation.—1. By dropping propyl alcohol on strongly-heated ZnCl₂ (Le Bel a. Greene, Am. 2, 23).-2. By passing gaseous HI into allyl iodide, PrI being also formed (Malbot, C. R. 107, 114; Bl. [2] 50, 449).—3. By pouring an alcoholic solution of allyl iodide on zinc (Gladstone a. Tribe, B. 6, 1550; Niederist, A. 196, 358).-4. By dropping propyl alcohol (4 pts.) on P₂O₃ (3 pts.) (Beilstein, B. 15, 1498).—5. By mixing isopropyl alcohol (200 g.) with ZnCl₂ (650 g.) and heating after twenty-four hours (Friedel a. Silva, J. 1873, 322).

Properties .- Colourless gas, liquefied by a pressure of eight atmospheres (Moltschanoffsky, $J.\ R.\ 21,31)$. Unites with Cl and Br. Its solution in H_2SO_4 yields isopropyl alcohol on distilling with water. Cuprous chloride in HClAq absorbs it somewhat. HClAq heated with propylene for seventy hours at 100° forms isopropyl chloride (Berthelot, A. 104, 184); HBr and HI act in like manner. Dilute CrO, forms acetone acetic acid and CO₂ (Berthelot, A. 150, 373). Alkaline KMnO, forms formic and acetic acids (Truchot, C. R. 63, 274). KCl added to a solution of propylene in PtCl2 and HClAq ppts. C₃H₄PtCl₂KCl aq (Birnbaum, A. 145, 72).

Di-propylene v. HEXYLENE.

References. - Bromo-, DI - BROMO - IODO-CHLORO-, CHLORO-10DO-, DI-CHLORO-NITRO-, and IODO- PROPYLENE.

PROPYLENE-ACETOACETIC ACID v. ACETO-ACETIC ACID and METHYL TRIMETHYLENYLMETHYL KETONE CARBOXYLIC ACID.

PROPYLENE-ALLYL-\u03c4-THIO-UREA

CH₂N(C₃H₃)>C:NH. Formed from propylene- ψ -thio-urea and allyl iodide (Hirsch, B. 23, 973). Oil.—B'C₆H₂N₃O₇. [126°]. Triangular prisms.

Isomeride CHMe.S C:NC₂H₃. [56°]. God

from β -bromo-propyl-amine and mustard oil. Prisms (from ligroin).—B'C₂H₄N₃O₇. [130°].—B'C₂H₅NCS. [52°]. White prisms.

PROPYLENE-DIAMINE C₃H₁₀N₂ i.e. S.G. ¹⁵ ·878. CH₂.CH(NH₂).CH₂.NH₂. (119°). S.G. 15 ·878. Formed by heating propylene bromide with alcoholic ammonia at 100°-150° (Hofmann, B. 6, 308; Strache, B. 21, 2358). Hygroscopic liquid, fuming in the air. Yields a hydrate $(C_3H_{10}N_2)_2$ aq, boiling at the same temperature as the base, and dehydrated by Na. Benzoic aldehyde forms oily C_sH₆(N:CHPh)₂, decomposed by HCl into the parent substances. Acetoacetic ether forms oily C3H6(NH.CMe:CH.CO2Et)2.

Salts.—B"H₂Cl. [220°]. Very hygroscopic needles.—B"H₂PtCl. Small four-sided tables. Acetyl derivative C,H,(NHAc)2. [139°].

Needles, v. e. sol. water and alcohol.

Benzoyl derivative C.H.Bz.N. [193°]. PROPYLENE-AMYL-y-THIO-UREA

heating allyl-amyl-thio-urea with conc. HClAq at 100° (Avenarius, B. 24, 264).

PROPYLENE BROMIDE v. DI-BROMO-PRO-

PROPYLENE TRICARBOXYLIC Ethyl ether CO.Et.CH.CH.CH.CO.Et),. (171°

at 15 mm.). Formed by the action of water or of NaOHAq upon CH C(CO,Et):C(OEt) O [94°], which is got by distilling di-carboxy-glutaconic ether in vacuo (Guthzeit a. Dressel, B. 22, 1425). It is identical with iso-aconitic ether.

Isomeride v. ACONITIC ACID.

Propylene tetra-carboxylic acid

(CO₂H)₂CH.C(CO₂H):CH.CO₂H. [192°]. Got by saponifying its ether. Colourless crystals (containing 2aq). Gives off CO2 at 180°.

Salts. Na, HAiv Saq. - Ba, Aiv 11 aq. -

Ca2Aiv 31aq.

Ethyl ether Et,Aiv. (220°-230° at 40 mm.). Formed from sodium-malonic ether and bromo-maleïc ether (Schacherl, A. 229, 91).

Isomerides .- DICARBOXY-GLUTACONIC ACID and TRIMETHYLENE TETRACARBOXYLIC ACID.

PROPYLENE CHLORHYDRIN v. CHLORO-PROPYL ALCOHOL.

PROPYLENE CHLORIDE v. DICHLORO-PROPANE.

PROPYLENE CHLOROBROMIDE v. CHLORO-BROMO-PROPANE.

PROPYLENE GLYCOL C₈H₈O₂ i.e. C₃H₆(OH)₂. Di-oxy-propane. Mol. w. 76. (189°). S.G. 2 1.0527 (Z.). C.E. (0°-10°) .00069. S. (ether) 11. S.V. 85.2 (Zander, A. 214, 177).

Formation.-1. By saponifying its acetylderivative (Wurtz, A. Ch. [3] 55, 438).--2. By reduction of CH,Cl.CH(OH).CH,OH by sodiumamalgam (Lourenço, C. R. 52, 1043).-3. By heating propylene bromide with PbO and much water at 150° (Eltekoff, J. R. 10, 210).-4. By boiling propylene bromide with aqueous K2CO3 for some days (Hartmann, J. pr. [2] 16, 383). 5. By boiling propylene bromide (1 pt.) with water (36 pts.), the yield being 43 p.c. (Niederist, A. 196, 359).—6. By the action of the copper-zinc couple and some HClAq on the product of the action of AcBr on glycerin (Hanriot, C. R. 86, 1139).-7. By treating allyl alcohol with dilute H2SO4 (20 p.c.) or HCl (10 p.c.) (Solonina, Bl. [2] 46, 816).

Preparation.—Glycerin (1,300 g.) is distilled with NaOH (550 g.) and the aqueous layer concentrated and distilled; the yield is 140 g. (Belohoubek, B. 12, 1872; Morley a. Green, C. J.

47, 132).

Properties. - Liquid, with sweet taste;

miscible with water and alcohol.

Reactions .- 1. Water containing a trace of HCl forms propionic aldehyde at 215° (Linnemann, A. 192, 61). It also yields propionic aldehyde on heating with ZnCl2 (Flavitzky, B. 11, 1256).-2. Conc. HIAq forms isopropyl iodide (Wurtz, A. Suppl. 1, 381) .- 3. PCl, forms dichloro-propane.-4. KOH at 250° forms oxalic acid and H.-5. Chromic acid mixture forms acetic acid (Flavitzky, B. 11, 1256).-6. Oxidised by platinum black and air to lactic acid.— 7. Glyceryl trimetric forms, at 100°, oily CH, CH(O.NO).CH, O.NO (109°), S.G. 2 1.44, which explodes in sunlight (Bertoni, C. C. 1887,

Acetyl derivative C, H₁₂O, i.e. C.H. (OAc). Propylens acetate. (186°). S.G. 1109. S. 11. Formed by heating C.H. Br. with AgOAe and HOAc (Wurtz, 4. 105, 202). Neutral liquid.

Mono-bensoyl derivative CH₂.CH(OBz).CH₂OH. Formed from the amine CH₃.CH(OBz).CH₂NH₂ and HNO₂ (Gabriel a.

Heymann, B. 23, 2501). Oil.

Di-benzoyl derivative CH₃.CH(OBz).CH₂OBz. (240° at 13 mm.). Liquid (Friedel a. Silva, C. R. 73, 1379).

 $Di\text{-}nitrate C_sH_6(O.NO_2)_2$. S.G. $\frac{\pi}{2}$ 1.335. Formed by dropping propylene oxide into strongly-cooled fuming HNO₃ (Henry, A. Ch. [4] 27, 261). Oil.

Chlorhydrin v. Chloro-propyl alcohol.

Bromhydrin v. Bromo-PROPYL ALCOHOL.

Iodhydrin C₃H₆I(OH). (105° at 60 mm.). Formed from propylene oxide and HI (Markownikoff, Z. 1870, 423). Liquid.

Active propylene glycol $C_3H_6(OH)_2$. $\alpha_D =$ 4° 35' to -1° 15' in a 22 mm. tube. Formed, together with propionic and lactic acids, by allowing Bacterium termo to breed in ordinary inactive propylene glycol (Le Bel, C. R. 92, 532). Yields active propylene oxide (35°) $a_D = +1^{\circ} 10'$ in a 22 mm. tube.

n-Propylene glycol v. Tri - METHYLENE GLYCOT.

PROPYLENE IODIDE v. DI-IODO-PROPANE. PROPYLENE MERCAPTAN $C_3H_6(SH)_2$. (152° in vacuo). Formed by reducing propylene sulphocyanide with Zn and HClAq (Hagelberg, B. 23, 1087)

PROPYLENE OXIDE C.H.O i.e.

CHMe O. Mol. w. 58. (35°). S.G. 2 ·859. V.D. 2.0. Formed by warming propylene chlorohydrin with KOHAq (Oser, Bl. 1860, 237; Morley, B. 15, 179). Dried over KOH. Neutral liquid, smelling like acetone. Miscible with water, alcohol, and ether. Reduced by sodium-amalgam to isopropyl alcohol. Yields acetic acid when oxidised by moist Ag₂O (Linnemann, M. 6, 369).

PROPYLENE-OXIDE CARBOXYLIC ACID v.

METHYL-GLYCIDIC ACID.

PROPYLENE PHENYL ETHYL KETATE CHMe.O CH₂—O CPhEt. (235° cor.). S.G. 22 .988. Formed by dropping C_sH_sCl.OBz into a mixture of ZnEt, and toluene, and then adding water (Morley a. Green, C. J. 47, 134; B. 17, 3015). Oil, with pleasant odour; sol. ether. Not attacked by alcoholic potash, by AcCl, by Na, by hydroxylamine, by phenyl-hydrazine, or by HClAq at 155°. HI at 200° yields PrI and phenyl ethyl ketone. By dissolving in H2SO. and pouring into water, it is split up into propylene glycol and C.H., CO.C.H., Nitric acid oxidises it to acetic and benzoic acids. Bromine forms a mono-bromo- derivative (230°)

PROPYLENE - PROPYL - ψ - THIO - UREA CHMe.S C.NHPr. [237°]. Formed by heating CH₂—N Class at 100° allyl-propyl-thio-urea with conc. HClAq at 100° (Avenarius, B. 24, 264). Oil.—Picrate [123°].

PROPYLENE SULPHIDE C.H.S. Formed from C₃H₆Br₂ and alcoholic Na₂S (Husemann, A. 126, 296). Amorphous powder.

PROPYLENE SULPHOCYANIDE C.H.N.S. i.e. CH₂CH(SCy).CH₂(SCy). Formed by heating C₂H₂Br₂ with potassium sulphocyanide and alcohol at 100° (Hagelberg, B. 23, 1086). Insol. water, sol. alcohol and ether.

PROPYLENE DISULPHONIC ACID v. Pro-PANE DISULPHONIC ACID.

PROPYLENE SULPHONIC ACID

CH₂:CH.CH₂.SO₃H. Formed by boiling allyl iodide with conc. K, SO, Aq (Von Rad, A. 161, 218).—KA'. Crystallir (KA'), K₂I₂. White solid. Crystalline solid. - (KA'), K, I,.

PROPYLENE TRI-THIO-CARBONATE

 C_3H_6 : CS_3 . S.G. 20 1·31. Formed from $C_3H_6Br_2$ and Na_2CS_1 (Husemann, A. 126, 269). Thick

liquid, of disagreeable smell.

PROPYLENE-4-THIO-UREA C.H.N.S i.e. CHMe.S C:NH. Imido-methyl-thiazole tetrahudride. Formed by heating allyl-thio-urea with HClAq (S.G. 1.17) under pressure at 100° (Gabriel, B. 22, 2985). Formed also from B-bromo-propyl-amine hydrobromide and potassium sulphocyanide (Hirsch, B. 23, 965). Liquid. — B'₂H₂PtCl₂. [212°]. — B'HAuCl₁. — B'C₆H₃N₃O₇. [200°]. — Methylo-iodide [172°]. Colourless prisms.

 $_{\rm CH_2,NH}^{\rm CHMe.O}$ > C:NH. PROPYLENE-\u03c4-UREA

Formed by heating allyl-urea with HClAq at 100° (Gabriel, B. 22, 2990), and by heating β -bromo-propyl-amine hydrobromide with potassium cyanate at 100° (Hirsch, B. 23, 966).-B'C.H.3N.3O. [186°]. Needles.—B'2H.PtCl. PROPYL ETHER v. DI-PROPYL OXIDE.

PROPYL-ETHYLENE v. AMYLENE.

PROPYL-EUGENOL v. EUGENOL.

PROPYL FLUORIDE PrF. (2°). V.D. 2·161 at 20° (calc. 2·175). S. 1 at 15°. Formed by gently warming AgF with PrI (Meslans, C. R. 108, 352). Gas, with ethereal odour and hot, sweet taste. Has no action on glass when dry. V. sol. alcohol, ether, and benzene.

Isopropyl fluoride PrF. (-5°). S. 1.5 at 15°. S. (alcohol) 29 at 16°. V.D. 2.171. Formed from AgF and PrI. Gas, not affected by KOHAq at 100°. Br at 30° forms a liquid (143°), while chlorine forms a liquid (105°).

ISOPROPYL-FORMAMIDE v. Formamide in article on FORMIC ACID.

DI-PROPYL-FURFURANE C4H2Pr2O. (120°). One of the products of the distillation of sodium citrate with lime (Bischoff, B. 23, 1918).

DI-PROPYL-GLUTARIC ACID C11H20O4 i.e. (CO₂H.CH.Pr)₂CH₂. [89°]. Formed by heating ((CO₂H)₂.CPr)₂CH₂ above 170° (Dressel, A. 256, 190). Needles, v. sl. sol. water, v. sol. ether.

PROPYL-GLYCOL v. PROPYLENE GLYCOL.

PROPYL-GLYOXAL. Mono-oxim. C.H., CO.CH:NOH. [51°]. Formed from methyl propyl ketone, amyl nitrite, and HCl or NaOEt (Claisen a. Manasse, B. 22, 528). Pearly plates. Di-oxim Pr.C(NOH).CH(NOH). [168°].

Di-phenyl-di-hydrazide

PrC(N₂HPh). (H(N₂HPh). [163°]. Needles. PROPYL - GLYOXALINE C₂H₁₀N₂ (219°–223°). S.G. ¹² · 267. Formed by heating glyoxaline with PrBr (Wallach, B. 15, 650; 16, 534; A. 214, 321). Liquid, miscible with water. HgCl. added to its aqueous solution gives a pp. sol. HClAq.—B'2H2PtCl. Crystals, sol. hot water.

Propyl-glyoxaline CH.NH CPr. Formed from glyoxal, n-butyric aldehyde and NH, (Rieger, M. 9, 603). Oil. — B',H,C,O,. [192°].—B'₂H₂C₂O₄ 2aq: [161°]; plates or needles. —B'₂H₂PtCl₆: yellow prisms. Isopropyl-glyoxaline. Glyoxal-isobutyline.

[129°]. (c. 250°). Formed from glyoxal and isobutyric aldehyde-ammonia (Radziszewski, B. 16, 747). Needles, sol. alcohol, benzene, and hot water.—B'HCl. [105°] (Rieger).—B'HBr. [105°] (Rieger).—B'HBr. [222°].—B'H₂C₂O₄. [195°].

Di-propyl-glyoxaline C6H,PrN2. (227°). S.G. 19 939. Formed by heating propyl-gly-

oxaline with PrI (Rieger, M. 9, 607)

n-PROPYL-GLYOXYLIC ACID Pr.CO.CO2H. (180°-185°) at 760 mm.; (115°) at 84 mm. Got by the action of HCl on the nitrile (133°-137°) which is made from AgCy and butyryl chloride (E. Moritz, C. J. 39, 16). Liquid.

Amide C,H,O.CONH2. [106°]. Formed

from the nitrile and conc. HClAq.

Oxim PrC(NOH).CO₂H. [144°]. Formed by the action of alcoholic soda and NaNO2 on propyl-acetoacetic ether (Fürth, B. 16, 2180). Small needles, v. sol. alcohol, sl. sol. water.

Isopropyl-glyoxylic acid *Pr.CO.CO.H. mixture (93° at 45 mm.) of this acid with isobutyric acid is got by the action of HCl on diisobutyryl dicyanide (Moritz, C. J. 39, 14).

Amide * Pr.CO.CONH... [126°].

PROPYL HEPTYL KETONE C,H,.CO.C,H, [12°]. (222°). S.G. 20 828. Formed by distilling

calcium butyrate (Limpricht, A. 108, 185).

PROPYL HEPTYL OXIDE ProC, H_{1s}. (188°).

S.G. § 7987. S.V. 245 6. C.E. (0°-10°) 00099

(Dobřiner, A. 243, 7).

DI - ISO - PROPÝL - HEXINYL DIKETONE Pr.CO.C₃H₅:C₃H₅.CO.Pr. Di-isobutyrone. 266°). Formed by the action of sodium on isobutyric ether (Bruggemann, A. 246, 151).

PROPYL - HEXYL - CARBINOL v. DECYL

ALCOHOL

PROPYL-HEXYL-GLYOXALINE C.H.15PrN2-Oxalpropylænanthyline. (286°). S.G. 17 ·919. Formed from hexyl-glyoxaline and PrBr (Karez, M. 8, 222). Oil.

PROPYL HEXYL KETONE Pr.CO.C. H13. $[-9.5^{\circ}]$. (207°]. S.G. $\frac{21}{0}.824$. Formed by oxidation of the corresponding alcohol (Wagner, J. pr. [2] 44, 271). Needles. Oxidised by K2Cr2O, and H2SO, to heptoic and propionic

Isopropyl hexyl ketone Pr.CO.C.H. (200°-210°). S.G. 17 ·841. Formed by distilling calcium isobutyrate with calcium heptoate (Fuchs, J. R. 7, 334). Yields acetone, HOAc, and heptoic acid on oxidation.

PROPYL HYPOPHOSPHATE Pr.P2O6. S.G. 15 1·134. Formed from Ag,P,O, and PrI (twice the calculated quantity) at 120° (Sänger, A. 232, 12). Oil. Decomposed by hot water.

Propyl-hypophosphate of barium

PrBaHP, O. 6aq. Needles. PROPYLIDENE-ACETIC ACID v. PENTENOIC

ISOPROPYLIDENE-AMIDO-PHENOL

CMe₂:N.C₆H₄.OH. [158°]. Formed from p-amidophenol, acetone, and HOAc (Haegele, B. 25, Colourless plates.

ISO-PROPYLIDENE-ANILINE C.H. N:CMe2. (228°). V.D. 66.4 (calc. 66.5). From acetone and aniline in presence of dehydrating agents (Engler a. Heine, B. 6, 638; Riehm, A. 238, 10). Liquid; rapidly turning brown in air. Its salts. are v. sol. water and alcohol.—B'2H2SO4: decomposes at 235°

PROPYLIDENE CHLORIDE v. DI-CHLORO-

PROPANE.

PROPYLIDENE - HYDRAZIDO - BENZENE p-SULPHONIC ACID CMe2:N.NH.C8H4.SO3H. Got from p-hydrazido-benzene sulphonic acid (phenyl-hydrazine p-sulphonic acid) and acetone (Pfülf, A. 239, 216). Plates, m. sol. hot water.

ISO-PROPYLIDENE HYDRAZINE CMe2:N.NH2. (125°). Formed from acetone and hydrazine hydrate (Curtius, J. pr. [2] 44, 543). Mobile liquid decomposing when kept, with

evolution of N and NH

ISO-PROPYLIDENE-PHTHALIDE C11H10O2 i.e. $C_6H_4 < CO \longrightarrow 0$. [96°]. Formed by heating phthalic anhydride with NaOAc and isobutyric acid or pyrotartaric acid (Gabriel a. Michael, B. 11, 1683; Roser, B. 17, 2776). Needles, sl. sol. hot water. Converted by boiling alkalis into Pr.CO.C.H4.CO2H.

DI-ISO-PROPYLIDENE DISULPHONE v.

TETRA-METHYL-DI-METHYLENE DISULPHONE.

ISOPROPYLIDENE - DI - THIO - DIGLY-LLIC ACID CMe₂(S.CH₂,CO₂H)₂. [127°]. Formed from thioglycollic acid, acetone, and ZnCl₂ (Bongartz, B. 21, 482). Crystals (from chloroform)

ISOPRÓPYL INDOLE C,1H,3N i.e.

 $C_sH_4 < CPr > CH$. (288°). Formed by heating the phenyl-hydrazide of isovaleric aldehyde with ZnCl₂ at 180° (Trenkler, A. 248, 106). Oil, solidifies by cold. Yields a dihydride (c. 260°) and a picrate [99°] crystallising in red needles. Di-isopropyl-indole $C_{14}H_{19}N$ i.e.

CPr:CH.C.CPr CH.CH.C.NH CH. [65°]. (295°-300°). Formed by the action of dilute H₂SO₄ on isopropyl-pyrrole in the cold (Dennstedt, B. 21, 3430). Needles (from dilute alcohol). Colours acidified pinewood red. Yields a picrate [115°] and an acetyl derivative [186°]. Benzoic aldehyde

and ZnCl, form $C_{35}H_{42}N_{2}$ [c. 164°]. n-PROPYL 10DIDE $C_{3}H_{1}$ I. (102·5°) (Brown, Pr. 26, 238). S.G. $\frac{9}{6}$ 1·7829 (Dobriner, A. 243, 24); $\frac{19}{6}$ 1·7673; $\frac{16}{5}$ 1·7585 (Perkin); $\frac{49}{5}$ 1·7427 (Brühl, A. 203, 15). C.E. (0°-10°) ·00105. μ_{β} 1·5157. R_{∞} 47·05. M.M. 11·08 at 18°. S.V. 106·9. Formed from propyl alcohol (60 g.), iodine (127 g.), and red P (10 g.), the yield being 90 p.c. of the theoretical amount (Chancel, Bl. [2] 39, 648;

cf. Linnemann, A. 160, 240). Oil.

Cr. Linnemann, A. 100, 240). Oli.

Isopropyl iodide Pri. (89°). S.G. 3 1·7033 (Brühl); ‡ξ 1·7163; ‡ξ 1·7005 (Perkin). M.M. 11·182 at 26°. μ_s 1·5108. R_∞ 47·48. S.V. 108·3. Formed by the action of HI on isopropyl alcohol, propylene glycol, glycerin, allyl iodide, or propylene (Linnemann, A. 161, 26; Wurtz, A. Suppl. 1, 381; Erlenmeyer, A. 126, 305; 189, 228; Maxwell Simpson, A. 129, 127; Berthelot, A. 104, 184; Butlerow, A. 145, 275). Prepared by distilling glycerin (200 g.) with I (300 g.), water (160 g.), and clear phosphorus (55 g. added slowly) in a current of CO₂ (Markownikoff, A. 138, 364; R. Meyer, J. pr. [2] 34, 98). Obtained also by saturating allyl iodide with HI and heating in sealed tubes at 100° (Malbot, C. R. 107, 114; Bl. [2] 50, 449; A. Ch. [6] 19, 352). Oil, converted by Br into PrBr. Vol. IV.

The copper-zinc couple decomposes it at 50°, giving off gases (Gladstone a. Tribe, C. J. 26,

PROPYL-ITACONIC ACID C₈H₁₂O₄ i.e.

CO₂H.CH₂.C(CO₂H):CHPr. [159°]. Formed by heating propyl-paraconic ether with alcoholic NaOEt, and saponifying the product (Fittig, A. 256, 106; cf. Schmidt, A. 255, 83). Tufts of prisms, insol. chloroform, v. sol. ether. Reduced by sodium-amalgam to butyl-succinic acid [81°]. Bromine, followed by hot water, gives rise to OHPr<0.00 CH [124°].—BaA".

DI-PROPYL-KETINE DI-METHYL-DI PROPYL-PYRAZINE.

DI-PROPYL KETONE C,H,4O i.e. COPr2. Butyrone. Mol. w. 114. (145°). S.G. $\frac{15}{25}$ 8217; 23 8145. H.C. 1,053,064 (Louguinine, Bl. [2] 41, 389). Prepared by distilling calcium butyrate alone or mixed with CaCO₃ (Chancel, A.52, 295; Kurtz, A. 161, 205; Schmidt, B. 5, 597). Formed also from ZnPr2 and butyryl chloride, by the oxidation of di-propyl-carbinol (Schtscherbakoff, J. R. 13, 346), and by heating butyryl chloride (1 mol.) with FeCl₃ (1 mol.) at 50° (Hamonet, Bl. [2] 50, 355). Got also by heating butyric anhydride with sodium butyrate at 180° (Perkin, C. J. 49, 325). It is also one of the products formed by the action of sodium on butyric ether (Brüggemann, A. 246, 140). Oil sol. alcohol. Does not combine with NaHSO. or NH3. Chromic acid yields propionic and butyric acids. Treatment with Zn and EtI followed by water gives ethyl-di-propyl-carbinol; while Zn, MeI, and water give methyl-di-ethyl-carbinol (A. Saytzeff, J. pr. [2] 31, 320). Sodiumamalgam forms sec-heptyl alcohol and a pina-cone C₁₄H₂₀O₂ [68°] (c. 260°). PCl₃ yields C₇H₁₄Cl₂ (181°) and C₇H₁₅Cl (141°) (Tavildaroff, B. 9, 1442). Gives on chlorination, when cooled by ice and salt, the compound Pr.CO.CHCl.Et (c. 175°) converted by NH₃ into tetra-propylpyrazine (Vladesco, Bl. [3] 6, 835). P₂O₅ forms (C,H₁₂)_x (200°-250°) (Tavildaroff, B. 9, 1442).

Oxim Pr₂C:NOH. (190°-195°) (Meyer a. Warrington, B. 20, 501). Yields an acetyl

derivative.

Di-isopropyl-ketone Pr₂CO. S.G. $\frac{9}{4}$ ·8230; $\frac{39}{4}$ ·8063 (Poletéeff, J. R. 20, 672). H.C. 1,044,559 (L.). R_{∞} 33·46. Formed by distilling calcium isobutyrate (Popoff, B. 6, 1255; Münde, B. 7, 1370; A. 180, 327). Formed also by heating tetra-methyl-phloroglucin with HClAq at 200° (Spitzer, M. 11, 288). Oil, with ethereal odour, miscible with alcohol and ether. Does not unite with NaHSO. Reduces ammoniacal AgNO. Yields isobutyric and accessed acids and CO₂ on oxidation, and secheptyl alcohol (c. 140° cor.) on reduction.

Oxim Pr₂C:NOH. [8°]. (c. 183°). Liquid, converted by AcCl into Pr₂C:NOAc, which on heating produces PrCO.NHPr (M. a. W.).

Dipropyl-diketone v. DIBUTYBYL.

Reference. - CHLORO-DI-ISOPROPYL-KETONE.

PROPYL-LUPETIDINE v. DI-METHYL-PROPYL-PYRIDINE HEXAHYDRIDE.

PROPYL-LUTIDINE v. DI-METHYL-PROPYL-PYRIDINE.

C,H1,0, PROPYL-MALONIC ACID Mol. w. 146. [84°] (8.); [96°] CHPr(CO,H)

(F.). H.C.p. 675,000. H.C.v. 674,700. H.F.

234,000 (Stohmann, J. pr. [2] 40, 211).

Ethyl ether Et.A.". (222°) (Fürth, M. 9, 309). S.G. 35 9931; 25 9854. M.M. 10:367 at 18° (Perkin, C. J. 45, 514). Formed from malonic ether, PrI, and zinc or NaOEt.

Isopropyl-malonic acid CPrH(CO₂H)₂. [87°]. H.C.v. 674,900. H.C.p. 675,200. H.F. 233,800 (Stohmann). Formed by saponifying its ether, which is made from sodium-malonic ether and PrI (Conrad a. Bischoff, B. 13, 595; A. 204, 144). Prisms, sol. water, alcohol, and ether.

Yields iso-valeric acid at 180°.—Ag₂A".

Ethyl ether Et₂A". (214°). S.G. 20 997 (C. a. B.); 15 9927; 25 9852 (Perkin). M.M. 10 482 at 17°.

References .- DI-BROMO- and OXY- PROPYL-

MALONIC ACID.

ISOPROPYL-MALONIC ALDEHYDE TRILE PrCH(CN).CHO. (137°). V.D. 3.83 (calc. 3.87). S.G. 15.911. Formed by heating iodoisovaleric aldehyde with KCy or AgCy (Chautard, A. Ch. [6] 16, 188). Oil which reduces Fehling's solution. Miscible with alcohol and ether.

n-PROPYL MERCAPTAN PrSH. Mol. w. (68°) (Roemer, B. 6, 784; Schatzmann, A. 261, 7). Formed from PrBr and alcoholic

KSH. Oil, smelling like mercaptan. Hg(SPr)₂.

Isopropyl mercaptan PrSH. (57°-60°) Isopropyl mercaptan PrSH. (57°-60°) (Henry, B. 2, 495; Claus, B. 5, 659; 8, 532). Formed from PrI and alcoholic KSH. Hg(SPr),: white plates (from alcohol).

PROPYL-METHANE is BUTANE.

Di-propyl-methane v. HEPTANE. DI-PROPYL-TRIMETHYLENE TRISUL-

PHONE $CPr_2 < SO_2 \cdot CH_2 > SO_2 \cdot [297^\circ]$. Formed from sodium trimethylene trisulphone and PrI in alcohol (Camps, B. 25, 244). Slender needles, m. sol. boiling alcohol.

Hexa - propyl - trimethylene trisulphone CPr₂ SO₂ CPr₂ SO₂. [133°]. Formed from the trisulphone, PrI, and alcoholic NaOH (Camps). Prisms.

ISO.PROPYL-NAPHTHALENE C18H14 C₁₀H₂Pr. (265°). V.D. 5.85. Formed by heating naphthalene with PrBr and AlCl, (Roux, Bl. [2] 41, 879; A. Oh. [6] 12, 289). Oil, sol. alcohol and benzene. Yields (\$\beta\$)-naphthoic acid on oxidation.—B'C₂H₂N₃O₃. [90°]. Lemon-yellow needles.

PROPYL NITRATE PrNO. (110.5°) (Perkin, C. J. 55, 683). S.G. 15 1.0681; 25 1.0531. M.M. 4.085. Formed by distilling n-propyl alcohol with HNO. and a little urea (Wallach a. Schulze, B. 14, 420). Forms with anthracene O₁₄H₁₀PrNO₂ [92°] crystallising in four-sided prisms (Perkin, jun. a. Mackenzie, C. J. 61, 866).

Isopropyl nitrate PrNO. (102°). S.G. 2

1.054 (Silva, A. 154, 256)

Reference.—CHLORO-ISOPROPYL NITRATE. PROPYL NITRITE Pro.NO. (43° (43°-46°) (Cahours, C. R. 77, 749); (53°-60°) (Pribram a. Handl, M. 2, 655). S.G. 21 ·935 (C.); § ·998 (P. a. H.). Formed by passing nitrous acid gas into n-propyl alcohol.

Isopropyl nitrite PrNO. (44°). S.G. 2 .856; 25 .844 (Silva, Bl. [2] 12, 227; Kissel, S.G. º

J. R. 1882, 226).

PROPYL \u03c4-NITROLE C3H6N2O2 i.e. CH₃.C(NO₂)(NO).CH₃. Mol. w. 118. [70°] (Bewad, B. 24, 976). Formed by adding KNO₂ and dilute H2SO, to an alkaline solution of isonitro-propane CHMe₂.NO₂ (V. Meyer a. Locher, B. 7, 670; A. 175, 120). Monoclinic crystals, insol. water and alkalis. Forms a blue liquid when fused. Its solutions in alcohol and chloroform are blue.

PROPYL-NITROLIC ACID C₃H₆N₂O₃ CH₃.CH₂.CH(NO₂)NO or CH₃.CH₂.Č(NO₂):NOH. [60°]. Formed by adding potassium nitrite and dilute H₂SO₄ to an alkaline solution of nitropropane CH₃.CH₂.CH₂.NO₂. Formed also from Et.CBr₂.NO₂ and hydroxylamine (V. Meyer, B. 7, 670; 9, 395). Prisms, with sweet taste, v. sol. water, alcohol, and ether. Alkalis form a deep-red solution. Conc. H_2SO_4 forms propionic acid and N.O.

PROPYL OCTYL OXIDE Pr.O.C. H17. (207°).

S.G. % 8039. C.E. (0°-10°). ·00101. S.V. 272·4 (Dobriner, A. 243, 7).

DI-PROPYL OXIDE Pr₂O. Propyl ether. (90.7° i.V.). S.G. $\frac{9}{6}$.7633 (Zander, A. 214, 163). C.E. (0°-10°) .00125 (Dobriner, A. 243, 20). S.V. 150.9. Formed from PrI and KOPr (Chancel, A. 151, 304). Got also from PrI and Ag₂O (Linnemann, A. 161, 37) and by heating n-propyl alcohol with H₂SO₄ at 135° (Norton a. Prescott, Am. 6, 243).

Di-isopropyl oxide Pr_2O . (68·8° i.V.). S.G. 7435. C.E. (0°-10°) 00130. S.V. 151·6 ·7435. Got from PrI and Ag₂O (Erlenmeyer, (Zander).

A. 126, 306).

PROPYL OXYBUTYL KETONE

C₃H₇.CO.CH(OH).C₃H₇. Butyroin. (180°-190°). Formed from di-n-propyl diketone and alcoholic KOH (Klinger a. Schmitz, B. 24, 1273). Yields a phenyl-hydrazide [135°].

PROPYL-PHENOL v. CUMENOL

PROPYL-PHENOL CARBOXYLIC ACID v. OXY-CUMINIC ACID.

Isopropyl-phenol dicarboxylic acid

 $C_0H_2Pr(OH)(CO_2H)_2$. [295°]. Formed, together with oxy-cuminic acid, by the action of Na and CO₂ on isopropyl-phenol (Fileti, G. 16, 126). Tables or needles, sol. water and alcohol.

p-PROPYL-PHENYL-ACETIC ACID C₂H₇.C₆H₄.CH₂CO₂H. [52°]. Formed by saponifying the nitrile, which is got from ω -chloro-cymene, alcohol, and KCy at 100° (Rossi, A. Suppl. 1, 139). Small needles (from hot water).—AgA': slender needles.

References.—Amido- and Oxy-Propyl-PHENYL-

ACETIC ACID.

PROPYL-PHENYL-AMINE v. AMIDO-PHENYL-PROPANE and PROPYL-ANILINE.

ISOPROPYL-PHENYL- v. CUMYL-

ISOPROPYL-PHENYL-CINNAMIC ACID Pr.C₆H₄.C(CHPh).CO₂H. [184°]. Formed from Pr.C.H.CH2.CO2Na, benzoic aldehyde, and Ac2O (Magnanini, G. 15, 509). Needles (from dilute alcohol).—CaA',.—AgA'. ISOPROPYL-PHENYL-p-COUMARIC ACID.

Methyl derivative Pr.O.H., O(CO.H.): OH.O.H., OMe. [199°]. Formed from Pr.O.H., OH., CO.Na, anisic aldehyde, and Ao.O (Magnanini, G. 15, 511). Prisms (from

alcohol).—AgA'.
DI-p-PROPYL-DI-PHENYL-CYANAMIDE C(N.C,H,Pr), [168°]. Formed by heating CS(NH.C_sH₄Pr)₂ with benzene and PbO (Francksen, B. 17, 1228). Needles.

PROPYL-BENZONITRILE v. Nitrile of Cu-

MINIC ACID.

ISOPROPYL - DI - PHENYL - ETHYLENE C_eH₃, CH:CH.C_eH₄, Pr. [84°]. Formed by heating phenyl-acetic acid with cuminic aldehyde and NaOAc at 250° (Michael, Am. 1, 314). Scales (from alcohol), v. sl. sol. hot water.

DI-p-PROPYL-DIPHENYL-GUANIDINE
NH:C(NH.C,H,Pr), [113°]. Formed by heating
di-propyl-di-phenyl-thio-urea with alcoholic NH,
and PbO (Francksen, B. 17, 1225). Needles, v.
sol. warm alcohol and ether.—Platinochloride (C₁₈H₂₂N₃), H₂PtCl₄: yellowish-brown pp.
Tri-propyl-tri-phenyl-guanidine

C.H.Prisc(NH.C.H.Pr). Formed by heating CS(NH.C.H.Pr). with C.H.Pr.M., alcohol, and PbO. Amorphous resin, v. sol. alcohol.—

B'2H2PtCl6: brown powder.

pn-PROPYL-PHENYL METHYL KETONE CH₃.CO.C₆H₄Pr. (259° i.V.). S.G. ¹⁶ '979. Formed from n-cumene, AcCl, and AlCl₄ (Widmann, B. 21, 2224). Colourless liquid. Yields an oxim [54°], which melts at 44° after fusion. The phenyl-hydrazide [92°] forms six-sided hatchet-shaped tables.

Isomeride CH₃.CO.C₆H₄Pr. (253°). Yields an oxim [71°] and a phenyl-hydrazide [82°].

ISOPROPYL-PHENYL PHOSPHATE
PO(O.C_oH₄Pr)_s. (375°-380° at 280 mm.). Got
from isopropyl-phenol and PBr₅ (Fileti, G. 16,
130). Viscid liquid, insol. water, sol. alcohol.

p-ISOPROPYL-(Py. 3)-PHENYL-QUINOL-INE C₁₈H₁₇N i.e. C₄H₄CHCH_{N=C,C₄H₄Pr. [60°]. Formed by heating its carboxylic acid with sodalime (Döbner. 4. 249. 102). Needles. sl. sol.}

lime (Döbner, A. 249, 102). Needles, sl. sol. water. — B'₂H₂PtCl₆ 2aq. — B'₂H₂Cr₂O₇. — Picrates: [195°]; plates (from alcohol).

Carboxylic acid $C_0H_4 < C(CO_2H): CH_4 Pr.$ [201°]. Formed from cuminic aldehyde, pyruvic acid, and aniline (Döbner). Yellow plates (from HOAo).—AgA': white powder.

p-PROPYL-PHENYL-THIOCARBIMIDE C.H.Pr.N:CS. (263°). Formed by heating dipropyl-di-phenyl-thio-urea with syrupy H₂PO₄ (Francksen, B. 17, 1223). Oil, volatile with steam, v. sol. alcohol and ether.

p-PROPYL-PHENYL-THIO-UREA C₁₀H₁₄N₂S i.e. NH₂CS.NH.C₆H₄Pr. [159°]. Formed from amido-phenyl-propane hydrochloride and ammonium sulphocyanide (Francksen, B. 17, 1222). Needles, sol. alcohol and ether.

Di-propyl-di-phenyl-thio-urea

CS(NH.C₆H₄Pr)₂. [138°]. Formed from amidophenyl-propane, CS₂, and alcohol (F.). Plates.

PROPYL-PHENYL-UREA C₁₀H₁₄N₂O i.e. NH₂CO.NHC₆H₄Pr. [143°]. Formed by the action of potassium cyanate on amido-phenyl-propane hydrochloride (Francksen, B. 17, 1225). Plates, sol. hot alcohol. insol. water.

Di-p-propyl-di-phenyl-urea CO(NHC₆H₄Pr)₂. [205°]. Formed by heating amido-phenyl-propane with urea. Formed also by the action of COCl₂ on amido-phenyl-propane in benzene, and by heating amido-phenyl-propane sulphate with a saturated solution of potassium cyanate. Needles, v. sol. ether and hot alcohol.

PROPYL PHOSPHATE PO(OPr). Formed, together with PrCl and PO(OH). (OPr) (which is sol. water) by the action of PCl, on n-propyl alcohol (Winssinger, Bl. [2] 48, 111). Oil. V. sl. sol. water, its insolubility being greatest at 75°. Cannot be distilled, even in vacuo.

ISOPROPYL-PHOSPHINE PrPH. (41°).

ISOPROPYL-PHOSPHINE PrPH. (41°). Formed, together with Pr_PH (118°) and Pr_P, by heating PrI with PH_I and ZnO for 6 hours at 100° (Hofmann, B. 6, 292). Liquid, with penetrating odour. Readily absorbs oxygen, taking fire on a hot day.—*B'HI: decomposed by water.

Tri-isopropyl-phosphine Pr₂P. Oil. Forms red crystals with CS₂.—Pr₃PHI. Large crystals, v. sol. water.—Pr₄PI. Cubes or octahedra.

Di-chloro-isopropyl-phosphine PrPCl₂. (c. 135°). Formed by heating HgPr₂ with PCl₂ (Michaelis, B. 13, 2175). Liquid.

Reference.—OXY-TRI-PROPYL-PHOSPHINE.

PROPYL PHOSPHITE P(OPr)_s. (240°). S.G. 16 1.004. Formed from POl, and NaOPr (Jaehne, A. 256, 282). Liquid.

ISOPROPYL-ISOPHTHALIC ACID

 $C_8H_4Pr(CO_2H)_2$ [1:3:5]. [285°]. Formed by heating pyruvic acid with isobutyric aldehyde and Ba(OH)₂ (Doebner, B. 23, 2380; 24, 1748). Plates (from dilute alcohol, v. sl. sol. cold water). —BaA" $2\frac{1}{2}$ aq.—CaA" $2\frac{1}{2}$ aq.—Ag₂A" aq.

PROPYL-PHYCITE. According to Fauconnier (C. R. 107, 629), the substance described under this name by Carius (A. 134, 71) is

glycerin.

B. 23, 680).

PROPYL-PIPERIDINE v. PROPYL-PYRIDINE HEXAHYDRIDE.

PROPYL-PROPANE v. HEXANE.

(a)-PROPYL-PYRIDINE C₈H₁₁N i.e. N CH. CH CH CH. Conyrine. (168°). Formed

by distilling coniine hydrochloride with zinc-dust (Hofmann, B. 17, 825; Ladenburg, A. 247, 20). Light oil, with blue fluorescence. Yields picolinic acid [134°] on oxidation. Reduced by conc. HIAq at 290° to coniine.—B',H_PtCl_s, [160°] and [172°]. Orange monoclinic tables. a:b:c=1.0622:1:1:5856; \(\beta=87^\circ 18'\)—B'MeI. Oil.—B',Me,PtCl_s. Crystals, sl. sol. water.

Tetrahydride NH CH2.CH2 CH2 or CH3.CH2.CH: CH2.CH2 CH2. This is (?)-coniceine (vol. ii. p. 248). Yields coniine on reduction with tin and HCl (Lellmann a. Müller,

Hexahydride NH CH2 CH2 CH2. Inactive coniine. (167°). S.G. 2 · 8626. Formed by reducing (a)-allyl-pyridine (derived from (a)-methyl-pyridine and paraldehyde) in alcoholic solution with sodium (Ladenburg, B. 19, 2679; A. 247, 80). Oil, optically inactive. Can be separated by means of the acid tartrate into a dextro-modification (coniine) and a lævo-modification.—B'HCl. [203°] (L.); [213°] (L. a. M.). V. sol. water.—B'2H2PtCl₄.—B'H,Cdl₄. [118°].

(β)-Propyl-pyridine N CH: CH: CH: Collidine. (170°). Got by passing nicotine through a red-hot tube (Cahours a. Etard, J. 1881, 928). Yields nicotinic acid on oxidation.

¥ \$

(a) - Isopropyl - pyridine $N \stackrel{CH . CH}{\stackrel{CP}{\sim}} CH$. (159°). S.G. 2 .9342. Formed, together with the (γ) -isomeride, by heating pyridine with PrI and PrI at 290° (Ladenburg, B. 17, 772, 1121; 18, 1587; A. 247, 22). Liquid, with unpleasant odour, sl. sol. water. Gives picolinic acid on oxidation with KMnO₄.—B'₂H̄₂PtCl₈. [170°]. Hexagonal rhombohedra, a:c=1:.9489.B'HAuCl₄. [91°].—B'C₆H₂N₃O₇. [116°]. Needles. Mercuric chloride double salt: [90°] .-

B'MeAuCl. [128°].

Tetra-hydride C₈H₁₃N. (164°). S.G. ²
896. Formed from the hexahydride, Br, and NaOHAq (Ladenburg, B. 20, 1646).—B'₂H₂PtCl_s. [189°]. Tables.

Hexahydride C₈H₁₇N i.e.

 $NH < CH_2$. $CH_2 > CH_2$. (159°). S.G. \circ .8668. Formed by reduction in alcoholic solution by Na (Ladenburg, A. 247, 73). Liquid, more sol. cold than hot water. Converted by MeI into CS_2 yields $(C_aH_{17}N)_2CS_2$ [105°]. — B'HCl. [210°]. Trimetric prisms; a:b:c=814:1:419. — B'HBr. (C₈H₁₇N)₂CS₂ [105°]. — B'HCl. [243°]. — B'2H2PtCl6. [233°].—B'HI. [193°]. Monoclinic prisms; α:b:c= 977:1:1·385; β = 89° 38'.—*B'₂H₂CdI₄. [133°].

v-Propyl-pyridine hexahydride

 $NPr < \stackrel{CH_2 \cdot CH_2}{CH_2 \cdot CH_2} > CH_2$ (149°-150°). Formed from piperidine and PrI (Ladenburg, B. 14, 1348). Liquid.—B'2H2SnCI6: monoclinic crystals.

v-Isopropyl-pyridine hexahydride

NPr CH₂.CH₂ CH₂ CH₂ -B'₂H₂SnCl₆: monoclinic crystals (Hjortdahl, J. 1882, 1085).—B'2H2PtCl3.

(178°). S.G. 2 9439. Formed as above, being separated by means of the platinochloride, which is more soluble than that of the (a)-isomeride. Oil. Yields isonicotinic acid [305°] on oxidation.—B'2H2PtCl6. [205°]. Plates.

NH CH2 CH2 CHPr. Hexahydride (171°). Formed by reducing (γ) -isopropylpyridine in alcoholic solution by Na. Fuming liquid, v. sol. cold water.—B'H2PtCl. [172°]. Golden prisms, sl. sol. water.

References. - OXY-PROPYL-PYRIDINE and OXY-

PROPYL PYRIDYL KETONE v. PYRIDYL PROPYL KETONE.

PROPYL-PIPERIDINE.

PROPYL - PYROGALLOL $C_9H_{12}O_8$ C₆H₂Pr(OH)₃. [80°]. Got by heating its dimethyl ether with conc. HClAq at 130° (Hofmann, B. 8, 67; 11, 329). Prisms (from benzene), v. e. sol. water and alcohol. FeSO, colours

its aqueous solution blue (Pastrovitch, M. 4, 182).

Methyl ether C₆H₂Pr(OH)₂(OMe). (290°
cor.). S.G. ¹² 1.023. Occurs, together with the di-methyl ether, in beech-wood tar (Pastro-vitch). Oil, with smoky smell. Gives an intense bluish-green colour with FeCl, in alcohol.

—C.H.2Pr(OK)2(OMe). Pearly crystals (from -C_eH₂Pr(OK)₂(OMe). Pearly crystal dilute alcohol). -C_eH₂Pr(OAc)₂(OMe). ì88°7. Needles (from alcohol).—C.Br.Pr(OAc)2(OMe). [79°]. Golden needles.

 $C_6H_2Pr(OH)(OMe)_2$. Di-methyletherPicamar. (c. 286° cor.). Occurs in beech-wood tar (Reichenbach, A. 8, 224; Niederist, M. 4, 487). Oil, with bitter, burning taste. Reduces salts of Au and Ag. Yields the di-methyl ether of di-oxy-quinone on oxidation.—C₁₁H₁₈O₃K. Pearly leaflets (from spirit).—C₁₁H₁₈AcO₃. [87°]. Monoclinic prisms; a:b:c=39:1:547; = 96° 29'.—C₁₁H₁₃Br₂AcO₅ [101°]. Trimetric prisms; a:b:c='99:1:1'93.—C₁₁H₁₃BzO₅. [91°].

Tri-methyl ether C₅H₂Pr(OMe)₃. (164°).

Formed from propyl-pyrogallol and MeI (Will,

B. 21, 2020).

n-PROPYL-PYRROLE C.H., N i.e.

NPr CH: CH: (147°). Formed from potassium pyrrole and PrI (Zanetti, B. 22, 2518). Liquid. Isopropyl-pyrrole C₄H₄PrN. (174°). Formed by boiling pyrrole with acetone and ZnCl₂ (Dennstedt a. Zimmermann, B. 20, 851; 21, 1480). Liquid. Dry HCl passed into its ethereal solution forms (C,H,1N)2HCl, a crystalline salt yielding a liquid base (c. 287°) and a picrate (C,H₁₁N)₂C₆H₅N₅O₇ [146°]. Ac₂O forms C₄H₅Pr:NAc (222°-232°) and C₄H₂AcPr:NH [64°]

(251°).

ISOPROPYL-PYRRYL STYRYL KETONE NHC₄H₂Pr.CO.CH:CHPh. [143°]. Formed by boiling NHC₄H₂Pr.CO.CH₁ with benzoic aldehyde and dilute KOHAq (Dennstedt a. Zimmer-

mann, B. 20, 853). Yellow crystals.

(B. 3)-ISOPROPYL-QUINOLINE C12H13N i.e. CH:CH.C.CH:CH Cumoquinoline. Formed CPr:CH.C.N = CHby heating (Py. 3)-chloro-isopropyl-quinoline with a solution of HI in HOAc (Widman, B. 19, Oil, easily volatile in steam.—Salts: B'₂H₂PtCl₆2aq. [220°]. — Picrate: [206°]. — Chromate: [c. 92°]. Large red prisms.

Methylo-iodide B'MeI. [c. 200°].

(Py. 2)-Isopropyl-quinoline $C_6H_4 < \begin{array}{c} CH:CPr\\ N=CH \end{array}$ [c. 10°]. (275°-280°) at 715 Formed by distilling its (Py. 3)-carboxylic acid (Spady, B. 18, 3383). Liquid, v. sol. alcohol and ether. B'2H2PtCl8. B'2H2Cr2O,.-B'CaHaNaO7: long slender needles.

(Py. 3)-Isopropyl-quinoline C_eH₄ CH:CH N=CPr (255°). Formed by heating its (Py. 1)-carboxylic acid with soda-lime (Doebner, B.20, 279; A. 242, 279). Oil, smelling like quinoline. B'₂H₂PtCl₅2aq: yellow needles.—B'C₆H₅N₅O₇. [150⁵]. Yellow plates (from alcohol).

Reference.—CHLORO- and OXY- ISOPROPYL-

(Py. 2)-ISOPROPYL-QUINOLINE (Py. 3)-CARBOXYLIC ACID C.H. CH:CPr [189°]. Formed by oxidation of (Py. 2,3)-isopropyl-isobutyl-quinoline with CrO, and dilute H₂SO, (Spady, B. 18, 3379). Plates (from dilute alcohol).—AgHA'2HNO3.—(HA')2H2PtCl3: prisms.

(Py. 3)-Isopropyl-quinoline (Py. 1)-carboxylic acid $C_eH_e < \stackrel{C(CO_eH):CH}{OPr}$. [146°]. Formed, together with C10H20N2O [222°], by adding aniline to an alcoholic solution of isobutyric aldehyde and pyruvic acid (Doebner, A. 242, 276; B. 20, 279). Prisms (containing 1 aq).—

HA'HCl. — (HA')₂H₂PtCl₆aq. — (HA')₂HAuCl₄:

lemon-yellow needles .- AgA

PROPYL SILICATE Si(OPr). (226°). S.G. 12 '915 (Cahours, C. R. 76, 1383). On heating with SiCl, at 160° it yields (PrO), SiCl (209°) and (PrO),SiCl, (187°)

DI-PROPYL-DI-STYRYL KETONE

(C_sH_r.C_sH₄.CH:CH)₂CO. [106°]. Formed from cuminic aldehyde, acetone, alcohol, and NaOHAq (Claisen a. Ponder, A. 223, 148). Prisms.

ISOPROPYL-STYRYL-PHENOL. Methylether. C.H.Pr.CH:CH.C.H.OMe. [152°]. Got by heating C.H.Pr.C(CO.H.):CH.C.H.OMe (Magnanini, G. 15, 513). Pearly scales (from alcohol).

PROPYL-SUCCINIC ACID C,H12O4 CO₂H.CH₂.CHPr.CO₂H. Oxy-hexic acid. [92°]. Formed by heating pentane tri-carboxylic acid (Waltz, B. 15, 608; A. 214, 59). Got also by reducing oxy-hexic acid with zinc and H2SO, (Gorboff, J. R. 1887, 605; Walden, B. 24, 2036).

Iso-propyl-succinic acid v. Pimelic Acid.

acid $C_{10}H_{18}O_4$ i.e. [168°]. The ether is Di-isopropyl-succinic CO₂H.CHPr.CHPr.CO₂H. [168°]. formed from a-bromo-isovaleric ether and finelydivided silver (Hell a. Mayer, B. 22, 48). Dendritic crystals (from water).—BaA" 5aq. ta 17°.—CuA" aq. S. 099 at 24°.—SrA" 3aq.—CaA"4aq.—MgA"7aq.—CoA"7aq.—NiA" 5aq.—CdA"4aq.—MnA"7aq. S. 4·71 at 21°.—ZnA".—PbA".—Ag₂A". S. 062 at 20°.

Isomeride C₁₀H₁₈O₄. [200°]. Accompanies Granular crystals. preceding acid. BaA"2aq.—SrA"4aq.—CaA"2aq.—MgA"5aq. — MnA" 3aq. — NiA" 4aq. — CuA" 3aq. — CdA" 3aq.

-ZnA".-PbA".-Ag₂A".

Reference.—OXY-PROPYL-SUCCINIC ACID.

TETRA-PROPYL-SUCCINIMIDINE C16HaiN, i.e. CH:C(NPr₂) N. Got from succin-imidoethyl ether hydrochloride, NHPr2, and alcohol at (Pinner, B. 23, 2930).—B'2HNO₃. (53°].—H₂PtCl_s. [174°]. Needles, sl. sol. hot water. **PROPYL SULPHATES**. B',H,PtCl

Propyl sulphuric acid Pro.SO2.OH. Formed from propyl alcohol and H₂SO₄ (Chancel, C. R. 37, 410). — KA': needles, v. sol. water. — BaA'₂ 3aq (Schmidt, Z. 1870, 576).

Di-propyl sulphate Pr₂SO₄. Formed from

propyl alcohol and CISO2.OH (Mazurowska, J. pr.

[2] 13, 162). Oil.

SULPHIDE Pr.S. DI-n-PROPYL (142°)(Winssinger, Bl. [2] 48, 109). S.G. 17 814 (C.) Occurs in raw petroleum (Mabery a. Smith, B. 22, 3303). Formed by heating K2S with PrCl or Prl in alcohol (Cahours, C. R. 76, 133). Fetid oil. Bromo-acetic acid forms the acid Pr₂SBr.CH₂.CO₂H, which yields Pb₃A'Br₅ and Pb₂A'Br₅ (Letts, Tr. E. 28, 586).

Compounds with platinum sal (Blomstrand, J. pr. [2] 38, 354, 498). $(Pr_2S)_2PtCl_2$. Occurs in three varieties: (a) [46°] S. (alcohol) 15.6 at 15°, (b) [86°], and (7) [63°]. Alcoholic KOH (1 mol.) acting on the (a)-compound forms crystalline (Pr₂S)₂PtCl(OH).—
(Pr₂S)₂EtCl₂: syrup.—(Pr₂S)₂PtCl₄. [185°].—(Pr₂S)₂PtBr₂. [139°].—(Pr₂S)₂PtBr. [141°].—(Pr₂S)₂PtBr. [110°].—(Pr₂S)₂PtBr. [141°].—(Pr₂S)₂PtBr. [110°].—(Pr₂S)₂PtBr. [110°].

[210°] and [195°].—(Pr₂S)₂Pt(NO₂)₂. Occurs in two varieties, an oil and a crystalline body [70°] which yields (Pr₂S)₂Pt(NO₃)(OH) [145°].-(Pr₂S)₂PtCrO₄. Red prisms (from chloroform). (Pr₂S)₂PtC₂O₄: orystalline pp.—(Pr₂S)₂Pt(SCN)₂: yellow pp.—(Pr₂S)₂PtCl₂HgCl₂. [82°]. Trimetric crystals, a:b:c = .554:1:.591.

Propylo-iodide SPr.I. Yields the com-

pound (SPr,Cl)2PtCl4 (Cahours).

Isopropyl sulphide Pr.S. (121° i.V.). Formed by distilling PrI with alcoholic K2S (Henry, B.

23, 495; Beckmann, J. pr. [2] 17, 459). Oxidised by KMnO, to Pr₂SO₂ [36°]. Yields Pr₂SHgOl₂.

Compounds with salts of platinum (Blomstrand):—(Pr₂S)₂PtCl₂. [163°].—
(Pr₂S)₂PtBr₂. [174°].—(Pr₂S)₂PtL₂. [176°].—(Pr₂S)₂PtCl₃. [176°].—(Pr₂S)₂PtCl₃. [176°].—(Pr₂S)₂PtCl₃. [176°].—(Pr₃S)₃PtCl₄. $(Pr_2S)_2Pt(SCN)_2$. $[102^\circ]$. $(Pr_2S)_2Pt(NO_2)_2$. Prisms, decomposing at 210°. $(Pr_2S)_2PtI_4$. $[139^\circ]$.

Di-propyl disulphide Pr₂S₂. (193°). (Spring

a. Legros, B. 15, 1940).

Di-isopropyl disulphide Pr.S. (175°). A product of the action of Na and Mel on isopropyl mercaptan in ether (Obermeyer, B. 20, 2923)

PROPYL SULPHOCYANIDE PrSCN. (168°). Liquid (Schmidt, Z. 1870, 576).

Isopropyl sulphocyanide PrSCN. (Henry, B. 2, 496); (153°) (Gerlich, A. 178, 90). S.G. 20 963. Formed from PrI and potassium sulphocyanide. Liquid, decomposed by boiling water. H₂S yields NH₂.CS₂Pr [97°].
DI-n-PROPYL SULPHONE Pr₂SO₂.

Formed by oxidising Pr₂SO (Winssinger, Bl. [2] 48, 111). Scales, sol. water, alcohol, and ether

Volatile with steam.

Di-isopropyl sulphone Pr₂SO₂. [36°]. Got by oxidising Pr₂S with KMnO₄ (Beckmann, J. pr. [2] 17, 459). V. sol. water and HClAq. Not

affected by reducing agents.
DI-PROPYL SULPHONE DICARBOXYLIC ACID SO₂(CHEt.CO₂H)₂. Sulpho-dibutyric acid. [152°]. Formed from SO₂(CH₂.CO₂Et)_e, EtI; and NaOEt (Lovén, B. 17, 2817). Dimetric octahedra.

Di-isopropyl sulphone dicarboxylic acid

SO₂(CMe₂CO₂H)₂. [188°]. Formed from SO₂(CH₂CO₂Et)₂ (1 mol.), MeI (4 mols.) and NaOEt (4 mols.) (Lovén, B. 17, 2824). DI-PROPYL SULPHOXIDE Pr₂SO. [15°].

Got by oxidising Pr₂S with dilute HNO₂ (S.G. 1.2) (Winssinger, B. 16, 329; Bl. [2] 48, 110). Needles, sol. water, alcohol, and ether. Decomposed by heat. Readily reduced to Pr2S. Calcium nitrate forms (Pr2SO)85Ca(NO3)2, a fibrous mass [80°

β-ISOPROPYLTHIENYL ETHYL KETOME C.H.Prs.CO.C.H. (251°). Formed from (β)isopropyl-thiophene, propionyl chloride, and AlCl, (Thiele, A. 267, 136). Oil, volatile with steam. Appears to yield isopropyl-thienylglyoxylic acid on oxidation with alkaline

KMnOPROPYL-THIENYL-GLYOXYLIC ACID

C₄H₂PrS.CO.CO₂H. Formed by oxidising propyl thienyl methyl ketone with alkaline KMnO₄ (Ruffi, B. 20, 1745). Solid.—AgA

PROPYL-THIENYL METHYL KETONE C.H.PrS.CO.CH. (255°). Formed by the action of AcCl and AlCl, on a solution of (a)-propylthiophene in ligroin (Ruffi, B. 20, 1744). Liquid. Yields an oxim [55°] and a phenyl-

Isopropylthienyl methyl ketone

C₄H₂FrS.CO.CH₂. (237°). Formed from isopropyl-thiophene, AcCl, and AlCl₂ (Thiele, A. 267, 187). Oil. Yields a crystalline oxim and

phenyl-hydrazide.

PROPYL-DI-THIO-BIURET C₅H₁₁N₃S₂ i.e. NH₂.CS.NH.CS.NHPr. [121°]. Formed from sodium cyanamide and propyl-thiocarbimide (Hecht, B. 25, 754). Crystalline mass, v. sl. sol. cold water.

PROPYL-THIOCARBIMIDE PrN:CS. (153°) S.G. $\frac{9}{0}$ ·9909; $\frac{99}{0}$ ·8924. Formed from propylamine and CS₂ followed by HgCl₂ (Hecht, B. 23, 281, 1662). Pungent liquid, sl. sol. water, miscible with alcohol and ether. Sodium cyanamide and alcoholic MeI form NHPr.CS.NCyMe [115°]. In the same manner may be prepared NHPr.CS.NCyEt [56°], NHPr.CS.NCyC₂H₃ [50°], NHPr.CS.NCyPr [56°] crystallising in needles, and NHPr.CS.NCy.CH₂Ph [113°].

Isopropyl-thiocarbimide PrN:CS. (137°).

Pungent liquid (Jahn, M. 3, 168).

PROPYL DI-THIO-CARBONATE Pro.CS.SH. Unstable oil.—Pro.CS.SK. Formed from CS2 and KOH dissolved in propyl alcohol (Scala, G. 17, 78). Silky needles. Its aqueous solution gives an orange pp. with CuSO4. Iodine forms oily (PrO.CS)₂S₂ decomposing at 180°.— PrO.CS.SMe. (202°). Oil.—PrO.CS.SEt. (216°).

(a)-PROPYL-THIOPHENE C4HsPrS. cor.). S.G. 10 .974. Formed from iodo-thiophene, PrBr, and Na (Meyer a. Kreis, B. 17, 1561). Oil. Yields (a)-thiophenic acid on oxidation.

Isopropyl-thiophene C_4H_3PrS . (154° cor.). S.G. $\frac{16}{16}$ 9695. Formed from PrBr, thiophene, and AlCl₃ (Schleicher, B. 19, 672). Oil with powerful odour. With HOAc, phenanthaquinone, and H2SO4 it gives a deep violet colour.

 β -Isopropyl-thiophene $\stackrel{CPr:CH}{CH:CH}>S$. (158°).

Formed by distilling sodium isopropyl-succinate with P2S5 (Thiele, A. 267, 133). Oil, smelling like benzene, insol. water, miscible with alcohol, ether, and benzene. Forms C.H., C.H., S.HgCl [187°] crystallising from alcohol in white needles, and C.H..C.HS(HgCl)₂ as white grains, insol. alcohol (Volhard, A. 267, 183).

References. - Bromo-, Iodo-, and Nitro-

PROPYL-THIOPHENE.

PROPYL-THIOPHENE CARBOXYLIC ACID C₄H₂PrS.CO₂H. [57°]. Formed from iodopropyl-thiophene, ClCO₂Et, and sodium-amalgam; the product being saponified by alcoholic potash (Ruffi, B. 20, 1748). Needles (from warm water), or plates (from dilute alcohol).
PROPYL THIOSULPHATE. The

Prs.SO₂Na 5aq is made from Na₂S₂O₃ and PrI

(Spring a. Legros, B. 15, 1938).

PROPYL-THIO-UREA NHPr.CS.NH2. [110°]. Formed from propyl-thiocarbimide and NH, Aq in the cold (Hecht, B. 23, 283). Four-sided plates, sol. alcohol.

Isopropyl-thio-urea NHPr.CS.NH. [1579].

Plates (Jahn, M. 3, 168).

Di - propyl - thio - urea CS(NHPr) [71°]. Formed in the preparation of propyl thiocarbimide. Pearly plates (from water)

Di-isopropyl-thio-urea CS(NHPr)2. [161°].

Needles (from hot water) (Jahn).

PROPYL-TOLUENE v. CYMENE.

PROPYL-TOLUIC ACID v. CYMENE CARBOXYLIC

S.G. 23 .0000 $n ext{-PROPYL-}p ext{-TOLUIDINE}$ C.H.Me.NHPr. (232°)..9296. $R_{\infty} = 82.5$. Formed = 1.5367. S.V. 197.53. by heating p-toluidine (1 mol.) with PrI (1 mol.) for two days at 160°; the yield being 90 p.c. of the theoretical amount (Hori a. Morley, C. J. 59, 36). Purified by means of the nitrosamine. Oil. Salts.—B'HCl. [151°]. Needles (from boiling benzene), v. sol. water and alcohol.—B'H₂C₂O₄. [173°]. S. (alcohol) 1.4 at 21°. Ppd. on mixing alcoholic solutions of the base (1 mol.) and oxalic acid (1 mol.). Sl. sol. cold water and alcohol.—B'₂H₂C₂O₄. [117°]. Occurs in the filtrate from the preceding salt, and formed also when a large excess of base is added to a cold alcoholic solution of the acid oxalate. Crystals, m. sol. cold water and alcohol. Decomposed by boiling water, B'H₂C₂O₄ being ppd.—B'₂H₄FeCy₆.
Nitrosamine C₆H₄Me.NPr.NO. Oil, not

solid at -20° . Decomposes below 100° .

Isopropyl - p - toluidine C₆H₄Me.NPr.NO. (220° uncor.). S.G. 23 9129. $\mu_{\rm p}=1.5322$. S.V. 199.57. R_{∞} 81.4. Formed by heating PrI with p-toluidine for two days at 130° (Hori a. Morley, C. J. 59, 34). Got also by heating diazotoluene toluide C,H,,N₂,NHC,H, with alcoholic NaOEt and PrI for 5 hours on the water-bath, distilling off the alcohol, pouring into water, and decomposing the pp. with HCl. Oil.

Salts.-B'HCl. [171°]. Wax-like crystals (from alcohol) or needles (from benzene), v. sol. (from alconol) or needles (Hold Bellett), r. sol. water. — B'₂H₂C₂O₄. [130°]. S. (alcohol) 5.76 at 22°. Crystals (from dilute alcohol). The acid oxalate is not easily crystallisable. — B'₂H₄FeCy_e. Nitrosamine C₆H₄Me.NPr.NO. [59°]. S. (alcohol) 65 at 22°. Yellowish crystals (from

alcohol), not volatile with steam.

Reference. -- OXY-PROPYL-TOLUIDINE.

NH2.CO.NHPr. PROPYL-UREA. [107°]. Formed from propyl cyanate and NH_s, and from potassium cyanate and propylamine sulphate (Chancel, Bl. [3] 9, 101). Long needles, sol. water and alcohol. Its oxalate and nitrate are v. sol. water. Decomposed by HClAq in a sealed tube at 160° into NH₃, NH₂Pr, and CO₂. Butyryl derivative NHPr.CO.NH(CO.C₂H₇). [99°]. Formed by the action of cold KOHAq on a mixture of butyramide (2 mols.) and bromine (1 mol.) (Hofmann, B. 15, 757). Colourless plates, sl. sol. water.

Isopropyl-ures. Isobutyryl derivative NHPr.CO.NH.COPr. [86°]. Formed in like manner from the amide of isobutyric acid. Tables, sol. alcohol and ether.

NH2.CO.NPr2. u-Di-propyl-urea Formed from dipropylamine and KCyO. Needles (from alcohol). Its oxalate crystallises in needles, sl. sol. water.

s-Di-propyl-urea CO(NHPr)₂. [105°]. (255°). Got by boiling the corresponding thio-urea with HgO and water (Hecht, B. 23, 285), or from PrCyO and aqueous NH_Pr. White plates (from water). Sl. sol. cold water, v. sol. alcohol and ether. Its salts are decomposed by water.

Di-propyl-urea C,H,eN2O i.e. NH2.CO.NPr [76°]. Formed from propylamine sulphate and potassium cyanate (Von der Zande, R. T. C. 8, 228). Needles. An ethereal solution of aldehyde forms CH, CH(NH.CO.NPr2)2 [113°]. Chloral

hydrate gives CCl₃.CH(OH).NH.CO.NPr₂ [128°] and in aqueous solution (C₇H₁₉N₂O)C₂HCl₃O aq Enanthol in ethereal solution gives C,H₁₄(NH.CO.NPr₂)₂ [113°].—B'₂H₂C₂O₄. [103°]. _B'C,H,N,O,. [135°]._B'2HNO.

s-Di-isopropyl-urea CO(NHPr)2. Formed as a by-product in the production of isopropyl cyanate from isobutyric bromo-amide and Na₂CO₃ (Hofmann, B. 15, 756). Needles, sol. alcohol, insol. ether.

u-Di-isopropyl-urea NH2.CO.NPr2. [103°]. Formed by evaporating a solution of isopropylamine sulphate with potassium cyanate (Von der Zande, R. T. C. 8, 231). Crystals (from ether). An ethereal solution of aldehyde forms CH₃.ĆH(NH.CO.NPr₂)₂[147°]. Chloral in ethereal solution gives CCl₃.CH(OH).NH.CO.NPr₂ [121°]. $-B'_{2}H_{2}C_{2}O_{4}$. [111°]. Needles. - [134°]. Needles. $-B'HNO_{3}$. [79°]. $-B'C_6H_3N_8O_{7}$

Reference.—DI-BROMO-PROPYL-UREA.

PROTAGON v. LECITHIN.

PROTAMINE C9H21N5O3 or C16H31N9O4. Occurs in the seminal substance of the salmon in November (Miescher, B. 7, 376; Piccard, B. 7, 1714). Gummy mass, insol. alcohol and ether, sol. water with alkaline reaction. Its salts give with K₄FeCy₅ a milky pp. composed of minute drops. Salts. — (C₆H₂₁N₅O₅)₂H₂PtCl₆ (M.); C₁H₂₁N₁O₁H₂PtCl₆ (P.). Yellow powder.

PROTEIDS. (Πρωτείον, 'pre-eminence.') This name is given to a large group of organic compounds which are also known as 'albuminous.' The word 'proteid' itself has been retained simply as a convenient general term, and not as implying any adherence to the theory, first advanced by Mulder, that they all contain a complex molecule to which he gave the name of 'protein.' The word albumin is now limited to a certain class of proteids, and the word albuminoid, though it is still largely used synonymously with proteid, is restricted by physiological chemists to a class of nitrogenous substances (which includes such substances as gelatin and mucin) which are not proteids, but still in certain reactions resemble those substances closely.

Proteids are never absent from the protoplasm of active living cells, whether animal or vegetable, and they are indissolubly connected with every manifestation of organic activity. A definition of proteids is not possible in the logical Gamgee (Physiol. Chem. p. 4) gives in the following sentences a terse description of these substances, which must take the place of a definition: 'Proteids are highly complex, and for the most part non-crystallisable compounds of carbon, hydrogen, nitrogen, oxygen, and sulphur, occurring in a solid viscous condition or in solution in nearly all the solids and liquids of The different members of the the organism. group present differences in physical, and to a certain extent in chemical, properties; they all possess, however, certain common chemical reactions, and are united by a close genetic relationship.

In vegetables the proteids are constructed out of the simpler chemical compounds which serve as their food. In animals such a synthesis never occurs, but the proteids are derived directly or indirectly from vegetables. By the action of certain digestive juices all proteids are capable of being converted into closely allied substances called peptones, which after absorption undergo reconversion into proteids.

Although much work has been done in at tempting to unravel the constitution of the proteids, there is at present but little positive to be said on this point. All we can do is to give the result of the experiments that have been performed which show the classes of products yielded by the proteids, and to enumerate the various theories held by different observers concerning their constitution.

The various proteids differ somewhat in elementary composition within the limits of the following numbers (Hoppe-Seyler, Handbuch d. Phys. und Path.-Chem. Analyse, 4te Aufl. p.

223):

The various decompositions that proteids undergo may be conveniently stated under the following heads:

a. Decomposition in the body. In the alimentary canal the proteids are converted into proteoses (albumoses) and peptones; this change is probably due to hydration. Under the influence of the pancreatic ferment, a certain class of peptones called hemi-peptones are further acted upon, resulting in the formation of leucine, tyrosine, aspartic acid, ammonia, and proteinchromogen (a substance coloured purple by bromine). Putrefactive processes due to bacteria in the small intestine also occur; these result in the formation of indole, skatole, phenol, and oxyacids. Ethereal hydrogen sulphates can be detected in the urine as a result of these putrefactive changes (Baumann, H. 10, 123); when putrefaction is hindered by the administration of large doses of iodoform in dogs, these products do not appear in the urine (V. Morax, H. 10, 318). One of the sources of hippuric acid in the urine of flesh-feeders is the phenyl-propionic acid that results from the putrefaction of proteids in the alimentary tract (Salkowski, B. 12, 648; Tappenier, Z. B. 22, 236).

After the proteids have been absorbed from the alimentary canal, they become assimilated by the tissues, and there undergo combustion or metabolism, the chief ultimate products being water, carbonic acid, and urea. It is probable that glycocine, leucine, creatine, and ammonium carbonate are intermediate products in this change. Urea had not been obtained from proteids by experiments performed outside the body until quite recently, when Drechsel (B. 23, 3096) has succeeded in obtaining a crystalline base lysatin by acting on casein with zinc and hydrochloric acid. Lysatin yields urea when boiled with baryta water. It has also been demonstrated, by experiments on animals, that proteid food gives rise to glycogen in the liver, and to fat in the subcutaneous and other tissues. That proteids can be converted into fats is also shown by the occurrence of adipocere in the muscular tissues after death.

b. Action of heat. By dry distillation the proteids yield an oily liquid called Dippel's oil,

which contains ammoniacal salts of the fatty acids, amines, and aromatic compounds.

c. Action of putrefaction. The chief products

are ammonia, ammonium sulphide, carbonic acid, volatile fatty acids, lactic acid, amines, leucine, tyrosine, and other complex aromatic

compounds.

d. Action of acids and alkalis. Prolonged boiling with strong sulphuric or hydrochloric acid, or fusion with caustic alkalis, gives rise to a large number of products, of which the following are the most important: leucine, tyrosine, aspartic acid, and glutamic acid. Prolonged heating with dilute acids gives rise to hydrolytic decomposition (Hermann) and the formation of albumoses and peptones (Neumeister, Z. B. 23,

381).

The effect of heating proteids with caustic baryta was investigated by Nasse, and subsection of the subsection of quently by Schützenberger (Bl. Feb. 15, March 5 and 15, 1875). Ammonia was evolved, and carbonic acid remained in combination with barium; these two products occurred in the same ratio as would result from urea similarly treated. addition oxalic acid, acetic acid, tyrosine, amidoacids of the fatty acid series (especially leucine, butylamine, and amido-butyric acid), acids allied to glutamic and aspartic acids, and lastly a dextrin-like substance, were found in the residue. Schützenberger (A. Ch. [5] 16, 289) explains the mechanism of the reaction as follows: the proteid molecule, losing ammonia and carbon as carbonic oxalic, and acetic acids, and assimilating water, is converted into a mixture of glucoproteins C_nH_{2n}N₂O₄, containing as its principal term C₂H₁₈N₂O₄; by the prolonged action of baryta at a high temperature these split up partly into leucines or amido- acids of the acetic series $C_nH_{2n+1}NO_2$ and leuceïnes or amido-acids of the acrylic series C_nH_{2n-1} . NO₂, and partly into double compounds formed by the union of glucoproteins and leuceines with the leucines. For Schützenberger's recent attempts to synthesise proteids see C. R. 112, 198.

e. Action of hydrochloric acid and stannous chloride. Ammonia, aspartic acid, glutamic acid, leucine, and tyrosine are the chief products (Hlasiwetz a. Habermann, Anz. Wien. Acad. 1872, 1873; A. 159, 304; 169, 150; J. pr. 7,

897).

f. Action of oxidising agents. By means of manganese dioxide and sulphuric acid, or potassium chromate and sulphuric acid, many fatty and aromatic compounds are formed. Nitric acid produces first a yellow insoluble substance, xantho-proteic acid, which dissolves on further action, and yields ultimately para-oxybenzoic

and oxybenzoic acids.

Mulder (J. pr. 16, 129; 17, 312; A. 31, 129) obtained by the action of potash on proteids a substance which is now called alkali-albuminate. but which was regarded by Mulder to be the base of all proteids; he gave it the name 'protein.' Various definite compounds of protein were described, such as oxyprotein, trioxyprotein, sulphoproteic acid, &c. Liebig (A. 57, 182) was the first to disprove these assertions, and the only remnant of this theory now is the term 'proteid.' For recent work on this subject and the related one of sulphur in proteids see Danilewski (H. 7, 440), and Krüger (Pf. 43, 244).

Gerhardt was of opinion that all the proteids are identical in constitution and molecular arrangement, but differ from one another in the nature of the mineral substance with which they are associated. He designated the common organic element by the name 'albumin,' and it may be mentioned that in the elaboration of his theory he regarded egg albumin as the neutral albumate of sodium, and fibrin as a compound of albumin with earthy phosphates. Proteids, however, have been separated from all but traces of mineral matter, and still exhibit their peculiar characteristics; careful elementary analysis also has shown that there is a difference of the percentage amounts of carbon, nitrogen, sulphur, &c., in various proteids. Nevertheless, it is still a fact that some of the distinguishing features of certain proteids are due to the adhering salt; for instance, the coagulation of caseinogen by rennet occurs only in the presence of calcium

salts (Hammarsten).

Gautier (Chimie appliquée à la Physiologie, i. 251) states that the proteids behave as amides of the higher homologues of lactic and tartaric acids, and residues of aromatic acids; hence when proteids are oxidised there is a simultaneous production of fatty acids, of aromatic compounds, and doubtless of bodies analogous to urea; the products furnished by different proteids are not in the same proportions. It must, therefore, follow that the different radicles they contain differ not only in arrangement but in relative proportion, and sometimes even in their nature. It was Nasse who first observed (Pf. 6, 589) that the nitrogen in proteids appears to exist in two conditions, a certain fraction of it being more unstable and apparently more feebly combined than the rest. This view was fully confirmed by Schützenberger in the experiments already The latter observer regards the alluded to. proteids as complex ureides, i.e. combinations in various proportions of urea with amido-acids, some of which belong to the leucine series, others to the aspartic series. Representing albumin by the empirical formula $C_{72}H_{112}N_{18}O_{22}S$ (Lieberkühn), it yields, when decomposed by caustic baryta, urea, acetic acid, a sulphur-containing body, and a substance with the formula C₈₈H₁₃₂N₁₄O₂₄, which may on further hydration be split up into the substances already mentioned (glucoproteins, leucines, leuceïnes, &c.).

Pflüger's view relative to the constitution of the proteids (Pf. 10, 251) may be thus summarised. One of the most striking features of the proteids is the difference that exists between nonliving proteid matter, such as white of egg, and that which forms part of living protoplasm. The former may remain for years without undergoing decomposition, and is indifferent to neutral oxygen; while the latter is continually decomposing, undergoing intramolecular changes, and lives by breathing oxygen. The assimilation of proteid food is looked upon by Pflüger as an ether-like combination of a molecule of living proteid with one of the non-living proteid, water being eliminated. This union of isomeric molecules, which forms a heavy mass but still remains a simple molecule, may be conceived to go on indefinitely. In this process it seems probable that the nitrogen of the food proteid, which exists there in combination as amides, passes into combination with carbon, so that in living proteid it is combined in the form of cyanogen. This view is strengthened by the fact that sub-

stances like uric acid, creatine, and guanine, are products of proteid metabolism, while such compounds are never obtained from non-living proteid material. O. Loew (Die chemische Kraftquelle im Lebenden Protoplasme, Munich, 1882, B. A. 1885, p. 1075) has also found that a striking difference between living and non-living proteids is the fact that the former reduce silver nitrate, probably owing to some radicle of the nature of aldehyde, whereas the latter do not behave in this way. He suggests that albumin is a condensation product of aspartic aldehyde. The simplest formula for Schützenberger's leuceine $C_4H_1NO_2$ is also that of aspartic aldehyde.

Latham (Brit. Med. Journ. [1] 1886, 629) regards albumin as a compound of cyan-alcohols united to a benzene nucleus. The cyan-alcohols resemble proteids in readily undergoing condensation and intramolecular changes; and, moreover, the various products of the disintegration of proteids can also be obtained as the decomposition products of these cyan-alcohols. See also Drechsel's art. in Ladenburg's Handw. d. Chem. 3, 541; Beilstein's Handb. 3, 1258.

GENERAL PROPERTIES OF PROTEIDS.

1. Indiffusibility. Solutions of proteids are non-diffusible. They belong to Graham's class of colloid substances. Peptones, and to a less extent albumoses, are, however, diffusible. This property of indiffusibility enables us to separate proteids from saline admixtures, and also to separate various proteids from one another: e.g. if a mixture of albumin and globulin in a saline solution be dialysed, the salts pass out, the albumin remains within the dialyser in solution, while the globulin, which is insoluble in water, is precipitated.

2. Action on polarised light. Proteids all rotate the plane of polarised light to the left. The following table gives the specific rotatory power for the yellow line D of some of the chief

proteids:

of various proteids is usually characteristic, but it can be varied somewhat by additions of small quantities of acid or alkali; the former lowers, the latter heightens, the coagulation temperature. Alkali albumin in the latter case is not coagulable by heat, and forms so readily that it is advisable to have the solution either perfectly neutral or faintly acid before performing the experiment. Rendering the liquid faintly acid with weak (2 p.c.) acetic acid also renders the flocculi of precipitate larger and easier to filter off. The alkalinity of a liquid is always increased, or its acidity diminished, after the separation of a heat coagulum. Various neutral salts in excess also cause the coagulation temperature to vary; sodium chloride lowers the temperature. while magnesium sulphate has little or no effect. A solution of proteid enters into an opalescent condition a few degrees below that at which the separation of flocculi begins. In the dry state proteids are rendered insoluble, like coagulated proteid, at a temperature of 110°C. On the influence of salts on coagulation temperatures, see Limbourg (H. 13, 450). Haycraft criticises the method of fractional heat coagulation (Pr. Roy. Soc. Edin. 1888-9, 361). It is defended by numerous observers (for references see Halliburton, J. Physiol. 13, 806; Hewlett, ibid. 798).

The following table gives the heat coagulation temperatures of some of the most important proteids:

Proteid				erature of	heat
Egg albumin		•	•	72-3°C.	
Serum albumin	α	•	•	72-3°	
Vitellin .			•	75°	
Serum globulin			•	75°	
Fibrinogen	•	•	•	56°	
Myosin .	•	•	•	56°	
Crystallin .	•	•	•	93°	
Hæmocyanin	•		•	65°	

4. General tests for proteids. Solutions of the proteids are precipitated by strong mineral

•				1
Proteid				Observer Value of [a] _p
Serum albumin			•	Hoppe-Seyler (Zeit. f. Chem. und
				Pharm. 1864, 737)56°
				Starke (Maly's Jahrs. 11, 17) -60°
Egg albumin .	•	•	•	Hoppe-Seyler
				Haas $(P. 12, 378)$ -38.08°
Fibrinogen .	•	•	•	Haas (C. C. 1876, 295, 811, 824)59.75°
				Hermann $(H. 11, 508)$ -43°
Sodium albumate			•	Haas
Syntonin (from egg	g albu	min)	•	Haas63·12°
Syntonin (from m	yosin)		•	Hoppe-Seyler \cdot \cdot \cdot \cdot -72°
Casein (in weak Mg	SO, s	olutio	a)	Hoppe-Seyler80°
Serum-globulin	•			Haas
Various albumoses				Kühne a. Chittenden $(Z. B. 20, 51)$. -70° to 80°

According to Haas, these numbers are unaltered by the salts adherent to the proteids.

3. Heat coagulation. Most of the native proteids are coagulated by heating their solutions, but more especially the globulins and albumins. By this means the character of the original proteid is lost, and all are converted into an exceedingly insoluble substance, called 'coagulated proteid.' When several proteids are mixed in a solution, if their temperature of coagulation differs, it is possible to separate them by a process of 'fractional heat coagulation.' The heat coagulation temperature

acids, acetic acid and potassium ferrocyanide, by acetic or oxalic acid together with a large addition of concentrated solutions of neutral salts, by basic lead acetate, by mercuric chloride, by copper sulphate, by tannic acid, by saturation with powdered potassium carbonate, ammonium sulphate or sodio-magnesium sulphate, and by alcohol.

Strong acids and salts of the heavy metals produce a precipitate of coagulated proteid, or of metallic albuminate; in the latter case the metal can be removed by treatment with sulphuretted hydrogen, and the proteid again ob-

tained unaltered. Neutral salts like ammonium sulphate, or sodio-magnesium sulphate, completely precipitate all proteids but peptones, and in an uncoagulated condition—i.e. they can be re-dissolved in water or weak saline solutions, and exhibit their characteristic properties. Alcohol precipitates proteids, and after long contact with the precipitate converts them into coagulated proteid.' The following reactions are those most frequently employed in the de-

tection of proteids in solution:

(a) The xanthoproteic reaction. Nitric acid is added; this usually produces a precipitate, but in the case of peptones, or if the quantity of proteid is very small, it does not. The mixture is then boiled, when the liquid and the precipitate turn yellow (xanthoproteic acid); if it is boiled sufficiently long with excess of acid the precipitate dissolves. Addition of ammonia causes the yellow colour to become orange. A similar reaction is given by gelatin and keratin. Analyses of xanthoproteic acid and its compounds made by Mulder and others are not of any value, as it has since been shown that this yellow substance is not constant in composition. The reaction is due to the aromatic radicle undoubtedly present in all proteids. On colour reactions of proteids see Salkowski (H. 12, 215), Gnezda (Pr. 47, 202), Pickering (J. Physiol. 14).

(b) The liquid is rendered strongly acid with acetic acid, and solution of potassium ferrocyanide added, which throws down the proteid

as a white flocculent precipitate.

(c) Millon's reaction. Millon's reagent 1 causes a white precipitate of the proteid. On boiling this precipitate and the surrounding liquid, both turn a brick-red colour. If only a trace of proteid is present there may be no initial precipitation. This test is given also by tyrosine, and it is in all probability owing to the aromatic radicle in proteids that those substances give it.

(d) Addition of a trace of copper sulphate and excess of potassium hydroxide causes a violet solution. In the case of albumoses and peptones the colour produced is a pink one (biuret reaction). See also Gnezda a. Pickering (loc. cit.) for similar reactions with nickel and cobalt.

(e) Proteids when dissolved in glacial acetic acid and treated with strong sulphuric acid exhibit a violet colour and faint fluorescence, and if sufficiently concentrated show an absorption spectrum very like that of urobilin (Adamkiewicz, B. 8, 161).

The reactions which are applicable to proteids which will not dissolve in water or neutral saline liquids, are the xanthoproteic, Millon's, and the Adamkiewicz colour reactions.

QUANTITATIVE ESTIMATION OF PROTEIDS.

A large number of different methods of estimating quantitatively the amount of proteid in a solution have been proposed and used; the polarimetric method has been much employed (Hoppe-Seyler, Virchow's Archiv, 11, 547). Among gravimetric methods various precipitants

¹ Millon's reagent. Equal quantities of mercury and strong nitric acid are mixed and gently warmed till the secroury is dissolved. The solution is diluted with twice its bulk of water, and the copious precipitate which forms allowed to settle. The supernatant liquid is the reagent.

of proteids have been suggested, e.g. tannin and sodium chloride (Girgensohn, N. R. P. 22, 557); hydrochloric acid and potassio-mercuric iodide (Brücke, Sitz. W. 1871), &c.; but the following methods are those most frequently em-

ployed:

(a) Clinical method. The rough method generally used for the comparison of albuminous urines consists in acidulating the urine, if it is as alkaline, with acetic acid and then boiling it in a graduated tube. The flocculi which form are allowed to settle, and the proportion of the precipitate to the total volume of the liquid read off and expressed in fractions, such as a half, a tenth, and so on. Esbach's tubes are graduated to correspond to percentages of proteid; the precipitant being picric acid.

(b) Precipitation by heat. 50 to 100 c.c. of water are boiled and an accurately-weighed amount of albuminous fluid, about 15 or 20 g., poured in. The fluid is boiled for some minutes, a few drops of two per cent. acetic acid added, until the precipitate separates in flakes from a clear liquid. The precipitate is collected on a weighed filter, washed with water, alcohol, and ether, and finally dried, weighed, incinerated, and the ash deducted. This method is obviously applicable only to such proteids as are coagu-

lable by heat.

(c) Precipitation by alcohol. An accurately measured or weighed quantity of the liquid is mixed with three or four times its volume of spirits of wine, and set aside for some hours; the precipitate is then collected, washed with spirit, hot absolute alcohol, ether, warm water, and finally dried and weighed as before (Hoppe-Seyler, Handbuch, 3te Auft. 312). Or the liquid may be neutralised with acetic acid and mixed with ten times its volume of strong alcohol, and then boiled, the precipitate being treated as before (A. Schmidt, Pf. 11, 10; A. Hoffmann, Virchow's

Archiv, 1879, 255).

(d) Precipitation with ammonium sulphate. Since Kühne (Z. B. 22, 423) has fully shown that saturation with this salt causes complete precipitation of all proteids but peptones, this method may be employed as follows for the quantitative estimation of these proteids:—The liquid is shaken or stirred with excess of the powdered salt, filtered, and the precipitate washed with saturated solution of ammonium sulphate. The filter with adherent proteid and salt is dried at 110°, by which means the proteid is rendered insoluble, and the salt can then be washed away with distilled water. The precipitate is then washed with alcohol and ether, dried, and weighed as before.

CLASSIFICATION OF PROTEIDS.

I. Albumins. These are proteids which are soluble in water, and not precipitable from their solutions by saturation with sodium chloride or magnesium sulphate. They are coagulated by heat. The important members of the group are egg albumin, serum albumin, and lactalbumin.

II. Globulins. These are proteids which are insoluble in water; they are soluble in dilute solutions of neutral salts; and are precipitated in an uncoagulated condition by saturation with sodium chloride and magnesium sulphate. They are coagulated by heat. The most im-

portant members of the group are: serum globulin, fibrinogen, myosin, crystallin, and globin.

III. Albuminates. This name is applied to the metallic compounds of proteids, and also to acid albumin or syntonin, and alkali-albumin. Restricting the term to the two latter substances, they may be defined as proteids insoluble in water or in solutions of neutral salts, but readily soluble in dilute acids or alkalis. Their solutions are not coagulated by heat.

IV. Proteoses. These are proteids which are not coagulable by heat, and most of them are precipitable by saturation with certain neutral salts. They are precipitated by nitric acid, the precipitate dissolving on the application of heat and reappearing when the solution is cooled. They resemble peptones in being slightly diffusible, and in giving the biuret reaction. They are formed from other proteids as the result of the action of proteolytic ferments on them, being an intermediate stage in the formation of paptones. They are also found in certain animal and vegetable tissues. The best-known members

of the group are the albumoses.

V. Peptones. These are proteids which are very soluble in water; they are not precipitated by heat, by saturation with any neutral salt, nor by nitric acid. They are completely precipitated by tannin, by excess of absolute alcohol, and by potassio-mercuric iodide; incompletely by phosphotungstic acid, phosphomolybdic acid, and picric acid. They give the biuret reaction. Peptones are subdivided into hemipeptones, those which yield leucine and tyrosine as the further result of pancreatic digestion, and antipeptones, those which do not.

VI. Insoluble proteids. This class includes a number of proteids varying in their reactions which cannot be included in any of the foregoing classes, but which all resemble one another in their extreme insolubility in various reagents. This class includes fibrin, coagulated proteid, lardacein, antialbumid, and gluten.

DESCRIPTION OF THE PROPERTIES OF INDIVIDUAL PROTEIDS.

It will now be convenient to take the most important members of the various foregoing groups, and describe the characteristic properties of each. The animal proteids will be taken first.

CLASS I., ALBUMINS.—Egg albumin. This is the albumin occurring in white of egg; it may be separated from the globulin, which also occurs in that situation, by precipitating the latter with dilute acetic acid, or by saturation with magnesium sulphate. It is coagulated at a temperature of 70°-73°C. By fractional heat coagulation further pps. are obtained at 76° and 82° (Corlin a. Berard, Arch. de Biol. 9, 1). Its specific rotation is -35.5°. It is precipitated by agitation with ether. Crystalline egg albumin may be obtained by addition of an equal vol. of sat. sol. of ammonium sulphate and filtration to remove oviglobulin. On slow evaporation of the filtrate, crystals of egg albumin form (Hofmeister, H. 14, 165; Gabriel, 15, 456).

Lieberkühn ascribed to albumin the formula $C_{72}H_{112}N_{18}SO_{25}$, Schützenberger $C_{846}H_{802}N_{65}O_{75}S_{5}$, Harnack (H.~5,~207) $C_{204}H_{322}N_{52}O_{66}S_{2}$. Since Lieberkühn attempted to establish the molecular weight of albumin by preparing and analysing

the copper compound, various investigations have been published bearing on the nature and composition of the compounds of albumin with the heavy metals, and especially with copper (F. Rose, Mitscherlich, Harnack, Bielitzki, Ritthausen, Lassaigne, &c. Further particulars will be found, with references to literature, in a paper by R. H. Chittenden and H. H. Whitehouse in Studies from the Laboratory of Physiol. Chem., Yale University, 2, 95). The results obtained are most contradictory, and illustrate that numerous compounds may be obtained by slightly modifying the conditions of precipitation; they are also very liable to dissociation. The only other proteid the metallic compounds of which have been studied is myosin; the percentage amount of the various metals in the compounds obtained differ from those formed from egg albumin (Chittenden a. Whitehouse,

Uses of albumin. It is much used for clarifying vinous and syrupy liquids, as when boiled with it the heat coagulum carries down colouring matter and other impurities. In cookery white of egg is employed for this purpose, but in large operations, such as sugar refining, blood serum is used. Albumin is also largely used in photography, and in calico-printing for fixing colours. Its property of forming a hard compound with lime renders it useful as a cement in repairing earthenware. Spoilt albumin in print works may be recovered for practical purposes by the action of pepsin (Wagner a. Witz, Ď. P. J. 119, 166).

Serum albumin. This is the albumin occurring in serum. Its specific rotation is -56° . Fractional heat coagulation, and elementary analysis show that it is probably not a single proteid but consists in warm-blooded animals of three, α , β , and γ , coagulating respectively at 73°, 77°, 84° (v. Bloop). Aronstein a. A. Schmidt (*Pf.* 2, 1) stated that serum albumin completely freed from salts by prolonged dialysis does not coagulate by heat. This statement has not been confirmed by subsequent observers, who have found it impossible to free serum albumin from ash by this method, and have attributed the non-coagulation to a slight alkaline residue (Heynsius, Pf. 9, 514; Winogradoff, 2, 605; Hinzinga, 2, 392; Haas, 2, 378). On ash-free albumin see also Harnack, B. 22, 3046; 23, 3745. Serum albumin is not coagulated by ether. It is the proteid which most frequently occurs in morbid urine.

Serum albumin is not precipitated by saturating the serum with magnesium sulphate; if, however, the globulin precipitated by this method be filtered off, and the filtrate saturated with sodium sulphate, the albumin is precipitated; this is due to the formation of sodiomagnesium sulphate (MgSO, Na, SO, 6H, O) in the liquid (Halliburton, Journ. of Physiol. [5] 152). Albumin having nearly the same characters as serum albumin is found also in milk, muscle, lymphatic glands, and other tissues in small quantities.

CLASS II., GLOBULINS. — Serum globulin. (Synonyms, paraglobulin, fibrinoplastin, serumcasein.) This is the globulin occurring in serum. It is soluble in weak solutions of sodium chloride; it is partially precipitated when the

percentage of sodium chloride reaches 0.03 to 0.07, re-dissolved on the addition of more salt, and again precipitated, though never completely, when the amount of salt present is 20 p.c. It is completely precipitated by saturation with magnesium sulphate. It does not coagulate on the addition of fibrin ferment, and probably takes no part in the formation of fibrin (v. art. BLOOD; v. also Hammarsten, Pf. 17, 447; 18, 39). Its heat coagulation temperature is 75°C.

The following methods have been suggested for the separation of the globulin from the

albumin of serum:

(a) Dialysis. When the salts are dialysed out, the globulin is precipitated, the albumin

remains in solution.

(b) Dilute the serum to 10 or 15 times its volume with water, and pass a stream of carbonic acid through it, the globulin is precipitated (A. Schmidt, Arch. f. Anat. u. Physiol. 1862, 429); it is also partially precipitated by the addition of very weak acetic acid when the serum is similarly diluted (Panum, Arch. f. Pathol. Anat. 4); it was on account of this that a special alkali albumin or serum casein was formerly believed to exist in the blood.

(c) Saturation with sodium chloride precipitates the globulin, leaving the albumin in

solution (A. Schmidt).

(d) Saturation with magnesium sulphate (Hammarsten). This is the only method commonly employed which effects a complete separation. Kauder has stated that half-saturation with ammonium sulphate will act similarly, and Halliburton (loc. cit.) that sodium acetate

and sodium nitrate may also be used.

Serum globulin may be estimated quantitatively in serous liquids in the following way: a weighed or measured quantity of the liquid is saturated with magnesium sulphate, the precipitate collected on a weighed filter and washed with a saturated solution of magnesium sulphate, and then dried at 110°C. for some hours; the proteid is thus rendered insoluble, and is freed from the salt by washing with water and alcohol. The precipitate is again dried on the filter to constant weight, weighed, and subsequently ignited and the ash deducted. In such a liquid as serum, viz. one containing a mixture of globulin and albumin, the latter may be estimated by deducting the amount of globulin from the amount of total proteids present.

Cell globulin found in lymph cells and

Cell globulin found in lymph cells and other cellular structures coagulates at 47°-50°. A globulin (myoglobulin) which coagulates at 63°C. but otherwise resembles serum globulin

exists in muscle plasma.

Fibrinogen. This proteid, which occurs in blood plasma, very closely resembles the preceding; it differs from serum globulin in being completely precipitable by half-saturation with sodium chloride, and may thus be separated from that proteid (Hammarsten, Pf. 19, 563). It is precipitable by a stream of carbonic acid, but it is necessary to dilute the plasma with at least 20 times its volume of water before precipitation occurs, whereas serum globulin is precipitated when the dilution is less. It is coagulated by heat at 56°C. The heat coagulum is a very sticky one; the low temperature at which fibrinogen coagulates forms the basis of a method for esti-

mating it quantitatively: the coagulated proteid formed from the fibrinogen at 56°C. is collected on a weighed filter, washed with distilled water and alcohol, dried at 110°C., and finally weighed, ignited, and the amount of ash deducted (Fredericq). Under the influence of the fibrin ferment, fibrinogen is converted into fibrin; it is due to this change that blood coagulates when drawn, the fibrin entangling the blood corpuscles to form the clot. The term fibringen has been recently extended by Wooldridge to other ill-defined proteid substances which he considers to be precursors of fibrin. These are probably nucleo-albumins mixed with lecithin. For latest views on blood coagulation with references see Halliburton (J. Physiol. 13, 829; Goulstonian Lecture No. 3, Brit. Med. Jl. March 1893).

Plasmine. The term 'plasmine' was given by Denis (Mémoire sur le Sang, Paris, 1859) to the precipitate produced by saturating blood plasma with common salt; this has since been shown to consist of the two proteids, fibrinogen and serum globulin; serine was the name given to the proteid left in solution, or, as it is now

called, serum albumin.

Myosin. This is the proteid which is formed after the death of muscle; the muscle plasma separates into a clot composed of myosin, and muscle serum, as blood plasma separates into fibrin and blood serum. This produces a stiffening of the muscles called rigor mortis. The name given to the proteid precursor of myosin in the living muscle is myosinogen. The clot of myosin formed by diluting with water a saline extract of flesh is not so coherent as that of fibrin; it is readily soluble in 10 p.c. sodium chloride solution, from which it is precipitable by saturation with that salt or with magnesium sulphate; it forms a heat coagulum in such a solution at 56°C. Myosin is converted with great readiness into acid albumin or syntonin by weak hydrochloric acid; A. Danilewsky regards the compound so formed as being not ordinary syntonin but a combination of HCl with myosin, from which by suitable treatment myosin is recoverable. After long-continued dialysis, myosin can be freed from salts, and is thus rendered exceedingly insoluble, both in saline solutions and in dilute hydrochloric acid (A. Danilewsky, H. 5, 158; Halliburton, Journ. of Physiology, 8, 136). (See also Muscle.)

Vitellin. A proteid present in yolk of egg, which, although it resembles globulins in certain particulars, differs from them in not being precipitable by saturation with sodium chloride. Solutions coagulate at 70°-75°C. It is prepared by treating the yolk of hen's egg with ether, by which reagent most of the fatty and colouring matters are dissolved; vitellin remains undissolved as a white granular substance, which is insoluble in water, but readily soluble in dilute saline solutions. Nuclein appears to be united closely to, and may be obtained from, vitellin.

Crystallin. This was once called globulin, but that term has been extended to the whole group. It is obtained from the crystalline lens of the eye, composing, according to Berzelius, 35.9 p.c. of that structure. Its saline solutions become opalescent at 73°, but a coagulum is not deposited till 93°C. is reached. According to

Lehmann (Physiol. Chem. 1, 366) the formation of a heat coagulum is accompanied by the development of an acid instead of an alkaline reaction, as is the case with other proteids; he accounts for this by supposing that crystallin is united to ammonio-sodic phosphate, which, on the application of heat, is resolved into ammonia and acid phosphate of soda. Valenciennes and Fremy (C. R. 44, 1122) state that other proteids are found in the lens, and have given the name phaconin to one in that of fishes (v.

also Béchamp, C. R. 90, 1255).

Ræmoglobin. When a solution of hæmoglobin, the red pigment of the blood, is heated, the liquid becomes turbid, and a coagulum of a reddish-brown colour separates out between 70° and 80°C. Heat decomposes the hæmoglobin into two substances insoluble in water, one coagulated proteid, the other a colouring matter, hæmatin, which contains all the iron of the hæmoglobin. A similar decomposition occurs when hæmoglobin is acted on by strong acids or alkalis, or by excess of alcohol. Preyer described the proteid substance under the term globin, which is insoluble in water, and which swells in

the proteid substance under the term globin, which is insoluble in water, and which swells in solutions of sodium chloride and sodium hydrate without dissolving. Kühne concludes from the action of reagents that a mixture of proteids rather than a single proteid results from the decomposition of hæmoglobin; these would, however, seem to belong to the globulin class from the fact that hæmoglobin is precipitable from its solutions by saturation with such neutral salts as precipitate other globulins, and also by a stream of carbonic acid (v. art. Hæmoglobin acid (v. art. Hæmoglobin).

Hæmocyanin. This is a proteid associated with a colouring matter which takes the place of hæmoglobin as an oxygen-carrier in the blood of certain crustacea, arachnida, gastropoda, and cephalopoda. It was first described by Fredricq in the blood of the octopus; the blood which comes to the gills being colourless, and that leaving them being blue: the blue pigment is due to a loose combination of the proteid with oxygen, which, like oxy-hæmoglobin, parts with its oxygen to the tissues. Copper takes the place of the iron of hæmoglobin in this compound. It has not been separated into a pigment and a proteid, as hæmoglobin has. coagulates by heat at 65°, and is precipitated by saturation with neutral salts, and by dialysis like

other globulins (for references see Halliburton,

Journ. of Physiol. 6, 300; Griesbach, Pf. 50,

473).

GLOBIN).

CLASS III., ALBUMINATES.

Acid albumin. If a small quantity of dilute acid (hydrochloric or acetic) be added to serum or egg-albumin, no precipitation occurs, but after a short time it will be found that the albumin no longer coagulates on heating, and that its dextrorotatory power is increased. The albumin has been converted into acid albumin. On carefully neutralising such a solution with weak alkali, the whole of the proteid matter is thrown down as a white precipitate, which is soluble in excess of the alkali, and may be re-precipitated by neutralisation with weak acid, and again dissolved by excess of the acid, and so on. Acidalbumin is soluble in solutions of alkaline carbonates; it is precipitated, like globulin, by saturation with sodium chloride or magnesium

Acid-albumin may also be formed in a similar way from any globulin; myosin, for instance, is converted into acid albumin or syntonin, as it was first termed by Liebig, when prepared from muscle. The parapeptone of Meissner, which is formed in the first stages of peptic digestion, is also identical with acid albumin.

Alkali albumin. If, instead of dilute acid, dilute alkali be added to an albumin or globulin, coagulation by heat is similarly prevented, and the whole of the proteid is thrown down by neutralisation, and is soluble in excess of either acid or alkali. The name alkali-albumin is given to the proteid thus altered by alkali. Like acid albumin it is soluble in solutions of alkaline carbonates, and precipitated by saturation with sodium chloride or magnesium sulphate. In the presence of alkaline phosphates it is necessary to add more acid to obtain a precipitate than if the alkaline phosphate were absent; the acid is just used to convert the alkaline into neutral phosphate. When heated with strong solution of potassium hydrate, potassium sulphide is not formed.

Lieberkühn's jelly. When either tribasic phosphoric acid, acetic or tartaric acids, which do not precipitate moderately concentrated solutions of albumin, is added in excess to concentrated serum or white of egg, a jelly is produced which liquefies like gelatin on heating, and again solidifies on cooling. A similar jelly is formed by adding a strong solution of potash to serum or white of egg. This substance, called Lieberkühn's jelly, is regarded as a variety of acid or alkali albumin respectively; it is soluble with difficulty in water; the aqueous solution remains, however, clear when boiled, and is precipitated by neutralisation.

Compounds of albumin with acids. On account of the present uncertain state of our knowledge concerning the constitution of the proteids, the statements concerning compounds which have been described of albumin with various acids must be received with caution, as is also the case with metallic albuminates (see

 $Egg\ albumin).$

Trinitro-albumin C₇₂H₁₀₅(NO₂)₃N₁₈SO₂₂ is prepared by triturating or shaking dried and powdered albumin with 14 or 15 times its weight of nitric acid (HNO₃); a gelatinous mass is the result, and on pouring this into water the insoluble light-yellow tinted compound separates out. If the gelatinous mass be left to itself at a temperature below 10°, a small quantity of nitrogen tetroxide is given off, and a solution formed which, when diluted with water, deposits a dark-yellow tasteless hydroxytrinitro-albumin, $C_{72}H_{100}(NO_2)_{5}N_{16}(OH)_{2}(SO_2OH)O_{27}$. Both these compounds dissolve without alteration in dilute alkalis, from which solution they are precipitated in yellow flocculi on addition of acid (O. Loew, J. pr. [2] 5, 433). If, instead of using nitric acid, a mixture of concentrated nitric and sulphuric acids be employed, a bitter yellow powder, insoluble in water, alcohol, or dilute acids, but soluble in dilute alkalis, is obtained which is called hexnitro-albumin sulphonic acid, C₇₂H₁₀₁(NO₂)_e(SO₂.OH)N₁₈SO₂₂; on treating this with ammonium sulphide, hexamido-albumin

sulphonic acid C₂₂H₁₀₁(NH₂)_e(SO₂.OH)N₁₈SO₂₂ is formed; this is decomposed by strong alkalis with evolution of ammonia, and by nitric acid with evolution of red fumes. It is insoluble in dilute acids, but readily dissolves in concentrated

ones (O. Loew, J. pr. [2] 3, 180).

Gelatinous compounds of albumin with acids are prepared by leaving it twenty-four hours in a parchment paper dialyser and a dilute acid in the outer vessel. For a quantity of albumin represented by Lieberkühn's formula, $C_{72}H_{112}N_{18}SO_{22}$, the compounds contain 2 mols. nitric, 2 mols. hydrochloric, 1 mol. sulphuric, and $1\frac{1}{2}$ mols. phosphoric acid (G. S. Johnson, O. J. 27, 734).

Casein and Caseinogen. Caseinogen is the chief proteid in milk. By the action of rennet it is converted into a curd called casein. Caseinogen is thrown down by the addition of acetic or mineral acids, especially if the milk be first diluted; this precipitate when freed from fats is insoluble in water, soluble in dilute acids or alkalis (and alkaline carbonates), and re-precipitable by acidulation. It may also be prepared from milk by saturating it with magnesium sulphate, followed by washing the precipitate with saturated solution of that salt till the washings contain no albumin, and then re-dissolving the precipitated caseinogen by adding water. salt still adherent to the precipitate enables it to dissolve: the fat remains undissolved on the filter. Such a neutral solution of caseinogen becomes cloudy on heating, and the cloudiness disappears on cooling if the heating has not lasted too long. Lubavin states that caseinogen differs from alkali albumin in the following particulars: (1) when caseinogen is treated with caustic potash potassium sulphide is produced; (2) on digestion with artificial gastric juice, caseinogen yields a substance containing phosphorus, whereas an albuminate can be prepared from white of egg which contains no phosphorus (Lubavin, Hoppe-Seyler's Med. Chem. Untersuch. 1871, 463). C. Wurster (B. 20, 263) states that when hydrogen peroxide is passed through a solution of egg albumin in the presence of sodium chloride and an acid (lactic, acetic, phosphoric, or sulphuric) a proteid insoluble in water is thrown down, which he states to be a mixture of gelatinous acid albumin with a large quantity of proteid similar to caseinogen; he further supposes that hydrogen peroxide plays an important rôle in the living tissues of the body, and that the caseinogen of milk may be produced from the proteids of the blood by a similar means. The precipitate produced by adding ether to yolk of egg is also stated to have properties similar to caseinogen (Bopp, A. 79, 16).

Most of the older writers use the word casein to include both it and caseinogen. Different varieties of casein have been described by different observers; Berzelius, Braconnot, and others described soluble casein, soluble in water, and msoluble casein, insoluble in water (v. also Bopp, l.c.). The soluble casein probably consisted of the other proteids present in milk in addition to casein. More recently the view has, however, been again advanced that the only proteid in milk is casein; and that the apparent presence of albumin and other proteids is due to the action of reagents or ferments (Duclaux, C. R. 98, 373, 438, 526;

E. Pfeiffer, Maly's Jahrber. 14, 177). H. Struve (J. pr. 29, 70, 110) distinguishes between casein which forms the envelopes of the fat globules, and that which exists in solution in the milk; both, however, have the same properties. Danilewsky (H. 7, 433) states that casein can be separated into caseo-protalbin, soluble in hot 50 p.c. alcohol, and caseo-albumin, insoluble in that reagent. Hammarsten (H. 7, 227) has, however, shown that this behaviour of casein depends in part upon its content of calcium phosphate, the presence of which impurity depends upon the use of hydrochloric acid in the precipitation of the casein, which acid does not favour the removal of the salt as well as acetic acid. Hammarsten, moreover, showed by elementary analysis that casein is a single body; this has been confirmed by Chittenden and Painter (Studies from the Lab. of Physiol. Chem., Yale Univ. 2, 156). The two last-named observers obtain the following as the percentage composition of casein: C, 53-30; H, 7.07; N, 15-91; S, 0.82; P, 0.87; O, 22.03. Hammarsten (H. 7, 269) and Ritthausen (Maly's Jahrsber. 1873, 28) have also published elementary analyses of casein.

It is owing to the presence of caseinogen that when rennet is added to milk a curd is produced, which is composed of the clotted casein and entangled fat globules; the whey or liquid residue contains the sugar, albumin, and salts of the milk. The spontaneous coagulation that sometimes occurs in milk is due either to the precipitation of caseinogen by lactic acid formed from lactose by fermentation, or to aerobic organisms which act like rennet. The coagulation that occurs when milk is boiled is due, not to the casein, but to the albumin in the milk. The ferment coagulation is a process which is entirely different from the precipitation by acid. A solution of pure cascinogen in dilute alkali will not coagulate on the addition of rennet; if, however, a small quantity of calcium phosphate or chloride be added, coagulation readily occurs. If caseinogen, precipitated by acetic acid, be well washed with water till the reaction is nearly neutral, and then neutralised with solid calcium carbonate, a pasty mass is formed which, on the addition of rennet thereto, sets into a solid clot: but here again if the caseinogen has been washed free from calcium phosphate rennet has no such This dependence of the rennet coagulation on the presence of earthy phosphate has been worked out by Hammarsten. The ferment causes the change from caseinogen to casein. The lime salt combines with the latter and precipitates it. Hammarsten showed that on the coagulation of caseinogen by rennet a whey albumin easily soluble in water is formed in addition to the clot of casein. Soxhlet (J. pr. [2] 6, 1) states that alkaline solution of artificially prepared alkali albumin also coagulates on the addition of rennet, but this statement has not been confirmed. Metacasein is a name given to an early stage of the action of rennet and pancreatic juice on caseinogen. It coagulates by heat (Roberts, Proc. Roy. Soc. 1881; Edkins, Journ. Physiol. 12, 193).

Casein in the milk of different animals is stated by Simon to vary slightly; thus human casein is yellowish-white, and very friable; it is incompletely precipitated from its aqueous solu-

tion by alum or by acetic acid. Casein from cow's milk is less soluble in water, and becomes viscid and horny on drying. Canine milk gives a casein which does not become viscid and horny when dried, and is less soluble in water. Human casein is much more easily digested than that of cow's milk, which is more viscid, and sets into a more solid clot under the influence of the rennet ferment of the gastric juice. For recent papers on casein see Ringer, Journ. Physiol. 11, 464; 12, 164; Halliburton, 11, 459; Sebelien, 12, 95; Hewlett, 13, 798.

CLASS IV., PROTEOSES.—Under the name proteose it is convenient to include the large number of products which are intermediate in the formation of peptones from other proteids. Similar substances are formed by the action of both the pentic and tryptic ferment; those formed from fibrin or egg albumin have received the name of albumose; from globulin, globulose; from casein, caseose; from vitellin, vitellose, &c. These various substances resemble each other in their general reactions, and differ from one another in origin, slightly in percentage composition, and in a few minor reactions. The albumoses are those which have been most studied. It will be sufficient here to describe the properties of the varieties of albumose. The globuloses, vitelloses, &c., are divided into similar varieties, with corresponding prefixes.

The properties of these substances have been chiefly worked out by Kühne and Chittenden. The names first given to them were propertone (Kühne) and α peptone (Meissner). Kuhne then adopted the name albumose, and subdivided the albumoses into hemi-albumoses, those which are converted into hemi-peptones, and anti-albumoses, those which are converted into anti-peptones. Ampho-albumose is a term which includes both hemi- and anti- varieties.

The following are the chief albumoses:

(i.) Proto-albumose, soluble in distilled water and weak saline solutions, partly precipitated by saturating its acidified solutions with sodium chloride. It is also precipitated by mercuric chloride and by copper sulphate.

(ii.) Hetero-albumose, insoluble in distilled water, soluble in weak saline solutions, and completely precipitated therefrom either by dialysing out the salt or by saturation with sodium chloride. It is precipitable by mercuric chloride only in acid solutions; it is precipitated by copper sulphate. By heating its saline solutions a cloudiness is produced, which is not caused by the formation of coagulated proteid, since it readily disappears on the addition of weak acid or alkali. By the action of alcohol hetero-albumose is partially converted into an insoluble product (dys-albumose).

(iii.) Deutero-albumose. This is the albumose most nearly allied to peptone. It is soluble in distilled water, and in saline solutions; it is not precipitated by saturation with sodium chloride. It is precipitable by nitric acid only in the presence of excess of salt, and then does not dissolve so readily on heating as in the case of the other albumoses. It is precipitable by mercuric chloride, but not by copper sulphate. It can be separated from peptones by saturation with ammonium sulphate, which precipitates

deutero-albumose but not peptones. For latest work on this see Kühne, Z. B. 29, 1.

The above reactions are those in which the albumoses differ from one another; they resemble one another in their characteristic behaviour with nitric acid (already described), and in giving the biuret reaction. Neumeister says that in the formation of peptones from albumin, in the first stage of digestion, proto-albumose, chiefly a hemi-product), hetero-albumose, and anti-albumid (an insoluble product only partially further changed, and that into antipeptone) are formed. Deutero-albumose is in all cases an intermediate product between these and the final peptones.

The following are the chief papers published in reference to this subject. On albumoses: Kuhne and Chittenden, Z. B. 20, 11; R. Neumeister, ibid. 23, 381; Chittenden and Bolton, Studies from the Physiol. Lab., Yale Univ. 2, 126. On globuloses: Kühne and Chittenden, Z. B. 22, 409. On vitelloses: Neumeister, ibid. 23, 402. On caseoses: Chittenden and Painter, Studies from the Physiol. Lab., Yale Univ. 2, 156. On the digestion of casein: H. Thierfelder, H. 10, 577. On benzoylated ethers of the albumoses: Schrötter, B. 22, 1950.

CLASS V., Peptones.—The general reactions of these substances and of their chief varieties, hemi- and anti- peptone, have been already described. It is only since the introduction of the method of separating them from other proteids by saturating a solution containing such a mixture with ammonium sulphate that they have been obtained free from proteoses. Most of the peptones of commerce consist almost entirely of proteoses. Peptones have also been described as occurring in milk, and in certain animal and vegetable tissues, whereas these proteids are in reality proteoses artificially formed during manipulation. Certain properties previously ascribed to peptones, e.g. a power of preventing blood from coagulating after intravenous injection, are now known to be due to proteoses.

When peptones are formed artificially a bitter taste is developed; what this is due to is unknown. Pure peptone has a meaty or cheesy taste; the albumoses are also free from bitterness. It is exceedingly difficult to dry pure peptone, and when dry it hisses and froths, with evolution of heat on the addition of water.

Anti-peptone—and the same appears to be true of other anti-products of digestion (e.g. anti-albumid)—do not only not yield leucine and tyrosine on further pancreatic digestion, but appear not to contain the tyrosine radicle; they do not give Millon's reaction, and do not yield tyrosine on boiling with sulphuric acid.

(For the most recent paper on pertones, methods of purification, and ultimate analysis, see Kühne and Chittenden, Z. B. 22, 423.)

CLASS VI., INSOLUBLE PROTEIDS.

Fibrin. This is the proteid formed from fibrinogen, probably under the influence of a ferment derived from the white corpuscles of the blood. It is owing to its formation that blood clots after withdrawal from the body (see BLOOD). It may be prepared pure by washing the clot free from corpuscles by a stream of water, or better, by allowing the plasma freed from corpuscles to coagulate, and then it is

washed free from other proteids by water. It is a white elastic solid, usually exhibiting fibrillation under a high magnifying power. soluble in water, partially soluble in dilute saline solutions, the proteid which goes into solution having the properties of serum globulin. Fibrin possesses the power of liberating oxygen from solutions of hydrogen peroxide. Fibrin swells up in cold weak hydrochloric acid (0.1 p.c.); it dissolves, forming acid albumin, on heating it with weak hydrochloric acid. It is readily digested by pepsin or trypsin, yielding albumoses and peptones. Hasebroek (H. 11, 348) and A. Herrmann (ibid. 11, 508) state that a first product in the digestion of fibrin is a substance with many of the properties of fibrinogen. (For elementary analyses and varieties of fibrin see Hammarsten, Pf. 22, 484; 80, 437. See also Fibrinogen.)

Coagulated proteid. This is proteid converted by heat or alcohol into a coagulum. It is insoluble in water or dilute acids or alkalis; it is, however, like fibrin, readily digestible by

proteolytic ferments.

Lardacein (so-called amyloid or albuminoid substance). This is formed in the disease known as waxy degeneration, the cells of the liver, spleen, and other organs being replaced by this substance. It is coloured brownish-red by iodine, hence the name amyloid (Virchow). Kekulé and Schmidt showed that it was a proteid. It is insoluble, like coagulated proteid, and it has been usually stated that it is not dissolved by gastric juice. Kostiurina (C. C. 1887, 120) finds, however, that it is soluble in pepsin solution acidified with hydrochloric acid.

The vegetable proteids have the same general reactions, and may be divided into the same classes as the animal proteids. Their percentage composition is also approximately the same, but varies with individual members of the group, as is the case with the animal proteids. tyrosine, asparagine, and other amides are found in plants, which may perhaps be stages in the formation of proteids from the union of inorganic nitrogen with certain non-nitrogenous compounds in the plant tissues (C. O. Müller, L. V. 1886, 326). Various observers have found in vegetable tissues a crystallisable proteid, often spoken of as an albumin, but which has since been shown to consist of a globulin (vitellin). These crystals were first observed by Hartig (Botan. Zeitung, 1885, 881). Maschke obtained crystals from Para nuts by extracting them with water at 50°; the crystals formed on evaporating this extract at the same temperature (J. pr. 74, 436). Similar crystals have been obtained by Schmiedeberg (H. 1, 205), by Drechsel (J. pr. [2] 19, 331), Ritthausen (J. pr. 181, 481). G. Grübler ob-tained octahedral crystals from pumpkin seeds and castor-oil seeds by cooling to 7° a sodium chloride extract of the seeds made at 70° (J. pr. 131, 105). The aleurone grains of plants have been investigated by S. H. Vines (Pr. 30, 387;31, 62); in the peony, castor-oil plant, blue lupin, and many other plants. Vines found that the grains are either wholly or partially composed of proteids, and classifies them according to whether (1) they are soluble in water, in which case they consist of vegetable peptone or hemi-albumose, (2) soluble in 10 p.c. sodium chloride solution, and (8) partially soluble in 10 p.c. sodium chloride solution. In the two latter cases globulins and albumins are also present. For recent work on crystallised vegetable proteids see Osborne, Am. 14, No. 8.

CLASS I., VEGETABLE ALBUMINS.-In addition to the crystalline body just described, the position of which in our classification is properly in the next group, there are in vegetable juices true albumins. These have been prepared and analysed from rye by Jones (A. 40, 66), from wheat flour by Dumas and Cahours (A. Ch. [3] vi), by Boussingault (ibid. [2] 63, 225), from potatoes by Rüling (A. 58, 306), in the latex of several caoutchouc-yielding plants by J. R. Green (Pr. 40, 28), and in papaw juice by Martin (J. Physiol. 6, 336). Such albumins are coagulated by heat at about 70°, and are not precipitated by saturation with sodium chloride, magnesium sulphate, nor by acetic acid. myrosin of mustard seeds also resembles vegetable albumin, and myco-protein from yeast and bacteria is also stated to be an albumin

(Schaffer, J. pr. 131, 302).

CLASS II., VEGETABLE GLOBULINS.—These were first described by Vines (L.c.), and have since been classified by Martin (Proc. Physiol. Soc. 1887, 8), who has found them in the flour of wheat, rye, and barley, in papaw juice, and in the seeds of Abrus precatorius (jequirity). He divides them into two classes: vegetable myosins and vegetable paraglobulins. The myosins coagulate at 55°-60°, are precipitated by dialysing the salt away from their solutions; but this precipitate is no longer a globulin, being insoluble in saline solutions. It has the properties of an albuminate, i.e. soluble in weak acid or alkali and precipitated by neutralisation. similar transformation occurs when a solution of the myosin, especially a dilute solution, is placed in an incubator at 35°-40° for twelve to eighteen hours. The vegetable paraglobulins coagulate at 70°-75°C., and are not transformed into albuminates, either by dialysis or by a long exposure to a temperature of 35°-40°. This classification does not include plant vitellin (phytovitellin), the crystalline form of which has been already alluded to. The crystals are membraneless, doubly refracting, and the proteid of which they consist has all the reactions of vitellin prepared from yolk of egg. This vitellin was found to be free from nuclein and lecithin, and may be regarded as the purest globulin yet known. Elementary analysis gave C 5243, H 7·12, N 18·1, S 0·55 p.c. (For further analyses of vegetable proteids see Barbière, J. pr. 126, 114.)

CLASS III., VEGETABLE ALBUMINATES.—Legumin and conglutin. Legumin, or vegetable casein, was discovered by Einhof in 1805, and called by him végéto-animale (N. allgem. J. D. Ch. v.; A. Gehlen, vi. 126, 548), in peas, beans, and lentils. Proust, H. Vogel, Boullay, Braconnot, and Liebig considered it to be identical with the casein of milk. Dumas and Cahours found that it contained less carbon and more nitrogen than casein (Gerhardt, Traité, 4, 491). The composition and properties of this proteid have been more recently studied by Ritthausen. He prepared it by extracting pul-verised peas, beans, lentils, and vetches with dilute alkali, precipitating the strained liquid

with acetic acid, washing the precipitate with alcohol on a filter, and finally drying over sulphuric acid. It dissolves both in cold and boiling water; on analysis the following percentages were obtained: From peas, lentils, vetches, and field beans—C 51:48; H 7:02; N 16:77; O 24:32; S 0:40. From garden beans—C 51:48; H 6:92; N 41:71; O 26:36; S 0:45. Treated with sulphuric acid it yields leucine, tyrosine, glutamic, and aspartic acids. The legumin of almonds and lupines, called by Ritthausen conglutin, is richer in nitrogen, more glutinous, and more soluble in acetic acid (Ritthausen, J. pr. 103; 65, 78, 193, 273; [2] 26, several papers; Z. [2] iv. 528, 541; vi. 126; Gm. xviii. 427, 437). Legumin differing slightly in properties and composition has been prepared from many other sources.

In spite, however, of the vast amount of work which has been done on this subject, and which is only briefly alluded to in the foregoing paragraph, the existence of this substance is now pretty generally denied. Fresh seeds or vegetable tissues and juices contain no proteids of the casein or albuminate group, and all the substances hitherto designated plant casein are artificial products produced by the alkali used to extract them from the various globulins and albumins which exist there normally (Hoppe-Seyler, Physiol. Chemie, Theil i. 75). Although Ritthausen defends his views (C. C. 1877, 567, 586) on the ground chiefly of the concordance of numerous elementary analyses, the recent work of Vines, Green, and Martin confirms Hoppe-Seyler's statement. For proteids of oats see Osborne, Am. 13, 327, 385; 14, 212; of maize, Osborne a. Chittenden, ibid. 13, 453, 529; 14, 20; of flax, Osborne, ibid. 14, No. 8.

CLASS IV.—VEGETABLE PROTEOSES.— These were first described by Vines (l.c.) in aleurone grains, and spoken of as hemi-albumose by him. They were previously regarded as peptones. Martin (J. Physiol. 6, 336) has given the name phytalbumose to these substances. In papaw juice he found two phytalbumoses: a - phytalbumose, with which the ferment papain is associated; this resembles the proto-albumose of Kühne a. Chittenden, but differs from it in the fact that copper sulphate and saturation with sodium chloride do not precipitate it; \(\theta\the

also under GLUTEN).

CLASS V. — VEGETABLE PEPTONES. — True peptone does not apparently exist as such in vegetable tissue; as the result of peptic or pancreatic digestion, however, peptones are formed with intermediate products (proteoses), as in the case of animal proteids. Some, moreover, are further acted on yielding leucine and tyrosine. It is interesting to notice that in the digestion of the proteids of papaw juice by papain, a proteolytic ferment present in that juice, no true peptones, but only phytalbumoses, are formed. Papain, however, which acts like trypsin best in Vol. IV.

an alkaline medium, forms true peptones with animal proteids (Martin, J. Physiol. 5, 213; 6, 236). For ferments in pineapple juice see Chittenden, Trans. Connecticut Acad. 8, 1. It appears probable that such ferments are widespread in the vegetable kingdom to convert the proteid of the cotyledons into peptone-like substances, and so render it available for food in the growing plant.

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CLASS VI.—INSOLUBLE VEGETABLE PROTEIDS.— Gluten. This is the most important of the vegetable proteids which falls under this head. When wheat flour is made into a paste its stickiness is due to the presence of gluten, which can be obtained free from starch by washing the paste with water; this is most conveniently done by kneading the paste in a muslin bag under running water. Gluten is tenacious. almost tasteless, of a light brownish-grey colour; on being burnt it emits the smell of burnt horn, and on destructive distillation yields the same products as animal proteids. It dissolves in strong acids and alkalis; it also dissolves, but very slowly, in 0.2 p.c. hydrochloric acid (Bouchardat, C. R. 14, 962). Gluten has been stated to possess amylolytic powers, converting starch first into dextrin and then into dextrose. Taddei (Giornale fisica di Brugnatelli, 12, 360) first showed that gluten consists of two substances—one soluble, the other insoluble in alcohol. Ritthausen called the part insoluble in alcohol vegetable fibrin, and the part soluble in alcohol, and to which the stickiness of the gluten is due, he subdivided into two substances, mucedin or vegetable mucin, and glutin, gliadin or vegetable gelatin. These are extracted with boiling alcohol of 70-80 p.c. This extract becomes turbid on cooling, and after half the alcohol has been distilled off mucedin is deposited in flocculi. It may be purified by redissolving it in hot alcohol, and re-precipitating by cooling. The gliadin remains in solution in cold alcohol. Mucedin is soluble in cold dilute acetic acid, but when precipitated from this solution by ammonia, or left in contact with or boiled with water or dilute alcohol, it is converted into an insoluble substance like fibrin. Gliadin forms a solution in alcohol which resembles varnish; on evaporating the alcohol it is obtained in a form resembling animal gelatin. It is more soluble in hot than cold water, and is precipitated by tannic acid, basic lead acetate, and mercuric chloride. The analyses of crude gluten, and of glutin, give approximately the same percentage composition as other proteids (Ritthausen, J. pr. 74, 193, 384).

Gunsberg (J. pr. 85, 213) regards mucedin as consisting simply of fragments of suspended fibrin; he states that gliadin is also not a distinct proximate principle, for cold water extracts from it a brown substance containing nitrogen and sulphur, and the residue has nearly the same composition as animal gelatin. Martin (Brit. Mat. Jour. 1886, 2, 104) has shown that gluten does not exist in flour as such, but is formed by the action of water (perhaps also by a ferment action) on the proteids pre-existent in the flour. The doctrine of a ferment action is supported by the fact that washing flour with water at a low temperature (2°C.) does not lead to the formation of gluten. Johannsen (Ann. Agronom. 14,

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420) has advanced evidence against the ferment theory. Gluten is insoluble in cold water, and in 10 to 15 p.c. sodium chloride solution. It dissolves partially in alcohol and in boiling water. Both these extracts contain the same substance, which is an albumose (insoluble phytalbumose), and it corresponds to Ritthausen's mucedin and gliadin. The insoluble residue, which is not sticky, may still be called gluten fibrin, and it is soluble in 0.2 per cent. hydrochloric acid. The flour itself contains two proteids: (1) A myosin coagulating between 55° and 60°, precipitated by sodium chloride and magnesium sulphate; and (2) a soluble phytalbumose. Both can be extracted from flour by 10-15 p.c. sodium chloride solution. They are considered to be the precursors of gluten, according to the following scheme:

Gluten = {Gluten-fibrin-precursor: myosin | Insoluble albumose - precursor: soluble albumose The more wheaten flour has been extracted with a 10 or 15 p.c. sodium chloride solution the less is the yield of gluten when it is subse-

quently treated with water.

The proportion of gluten in wheat flour varies considerably according to climate, soil, temperature, &c.; the best flour containing 10-11 p.c., inferior kinds 8-9 p.c. moist state gluten weighs about three times as much as when dry. The flours of barley, rye, and oats contain only a trace of gliadin, or albumose, as it may now be regarded; hence these flours, containing mostly gluten-fibrin or its precursor, cannot be made into such a sticky paste as can be made with wheat flour. Gliadin also does not exist in leguminous seeds; it is said to be present in the juice of the grape and other fruits, and is held in solution there by tartaric or other vegetable acids.

APPENDIX A.—Proteids in urine. Normal urine contains no proteid. The chief facts concerning proteids in morbid urine are treated fully in medical works (MacMunn's Clin. Chem. of Urine, Halliburton's Chem. Physiol.), but may

be here briefly summarised as follows:

When urine contains blood (which may be detected by the microscope or spectroscope) or pus (which may be also detected microscopically), it of necessity contains a certain amount of proteid derived from these contaminations. In paroxysmal hæmoglobinuria, the urine may contain hæmoglobin or methæmoglobin independently of the presence of blood corpuscles. In the various forms of Bright's disease, the unhealthy kidney allows the proteids of the blood to diffuse into the urine, the most common and abundant being serum albumin. globulin is usually also present, and is recognised by the fact that saturation with magnesium sulphate in the neutralised urine precipitates it (A. Ott, C. C. 1886, 540). A crystalline globulin was found in urine by Paton, Lab. Rep. R. Coll. Phys. Edin. 4, 47.

Clinical observers do not as a rule attempt to distinguish, however, between the various forms of proteid that occur in urine, but speak of them all under the name of albumin. The most commonly applied tests for its detection are:—(1) To boil the upper part of a test tube full of urine; any cloudiness produced is then seen in contrast to the clear urine below. If any precipitate occurs it is insoluble in acetic acid, so distin-

guishing it from phosphates. If the urine is neutral or alkaline it is necessary always to acidify with acetic acid either before or after boiling, in order to obtain a precipitate. The clinical method of estimating the proteid quantitatively has been already described. (2) Nitric acid causes a precipitate in the cold. If urine contains but little proteid, it may be detected by pouring the urine on to the surface of some nitric acid in a narrow test tube, the ring of precipitate at the junction of the two liquids is then clearly seen (Heller). The difficulty of carrying nitric acid about has led to the adoption more recently of methods in which test papers saturated with various reagents, picric acid, potassio-mercuric iodide, &c., are added to the urine. A committee of the Clinical Society have recently presented a report in which the relative advantages of various methods of detecting albumin in urine are discussed (Clin. Soc. Trans. 19, 339). They have investigated Dr. Oliver's test papers just mentioned; Dr. Pavy's pellets of citric acid and potassium ferrocyanide; Dr. Johnson's picric acid solution; Sir W. Roberts's acid brine test; acetic acid and boiling nitric acid; and a solution of potassio-mercuric iodide with citric acid. The last named is the most delicate reagent in the list; the test papers and pellets are not so delicate but are more convenient; the picric acid test has the advantage of being also applicable for recognising (after the addition of potash) sugar as well as albumin.

Besides albumin and globulin in urine, other proteids may sometimes occur; viz. (1) egg albumin—this occurs after a very large ingestion of eggs as food; (2) Bence-Jones albumin, so called after its first observer, has been shown to be a form of hemi-albumose—it occurs in cases of osteomalacia; (3) peptones or deutero-albumose occurs in many suppurative diseases, and doubtless originates from the disintegration of pus cells; (4) casein has been stated to occur in chylous urine, but there are considerable doubts as to the accuracy of this statement; (5) mucin occurs not only in suspension in the mucus from the urinary tract, but it appears to be occasionally present dissolved in the urine-it is precipitable therefrom by acetic acid. It is probably not true mucin, but a nucleo-albumin.

APPENDIX B.—Proteids as poisons. For effects on blood pressure, &c., of albumoses and peptones, v. Pollitzer, J. Physiol. 7, 283. The poison of venomous snakes has also been shown to depend on the proteids contained therein (Weir Mitchell), and not to any alkaloid, ptomaine, or cobric acid (W. Blyth, An. 1, 204). Wolfenden has separated a globulin, albumin, syntonin, and albumose from the venom of the cobra and daboia; for the action of these v. J. Physiol. 7, 327; see also Kanthack, ibid. 13, 272. The poisonous principle abrin of Abrus or Jequirity is also proteid (Martin, Pr. 42, 331). The subject of proteid poisons has recently risen into prominence, as it is found that the micro-organisms of many diseases produce these toxalbumoses. Koch's tuberculin owes its activity to similar substances. The poisons produced in anthrax, diphtheria, &c., are also proteid in nature. For references see Halliburton's Chem. Physiol. German translation; Martin, Brit. Med. Journal, March and April 1892. On the other hand,

certain proteids (globulins or nucleo-albumins) are protective and destroy micro-organisms. They are termed alexines by Daremberg, Buchner, and Hankin. For Hankin's most recent paper, see *Centr. Bacteriol.* 12, Nos. 22 and 23.

APPENDIX C.—Albuminoids. The term 'albuminoid' is still used by some synonymously with proteid; it will be here restricted to a number of substances which resemble proteids in many points, but which differ from them in others.

Collagen. This is the substance of which the white fibres of connective tissue are composed, and which by the action of boiling water is converted into gelatin. Collagen is prepared from tendons by Rollett's process as follows: the finely-divided tissue is soaked in water to remove proteids, and then for some days in lime water to dissolve the mucin-holding cementing substance between the fibres. The insoluble matter is washed first with water, then with weak acetic acid, and then again with water. The residue consists of collagen, mixed, however, with small quantities of elastin and nuclein. With very dilute acids or alkalis collagen swells and becomes transparent.

The organic material which composes onethird of the substance of bone consists mainly of a substance identical with collagen, and which has been termed ossein by some writers. The organic basis of dentine (but not of the enamel

of tooth) is also collagenous.

Gelatin. When the white fibres are subjected to the action of boiling water, or of water heated under pressure, as in a Papin's digester, or to the long-continued action of dilute acids at the ordinary temperature, they dissolve, and the solution contains a substance called gelatin. Gelatin may be similarly prepared from bones. Isinglass is gelatin prepared from the swimming bladder of the sturgeon; an inferior kind is made from fish bones. Glue is the crude product obtained by boiling down hides, bones, &c. (v. Ure, Dictionary of Arts, &c., [2] 324, 376). Pure gelatin is prepared from commercial gelatin by soaking the latter in distilled water for some days to remove salts; it is then dissolved in hot distilled water, and filtered while hot into The gelatin separates in the 90 p.c. alcohol. form of white thready masses, which can be subsequently dried. Thus prepared it contains only 0.6 p.c. of ash.

Gelatin is insoluble in cold, but soluble in hot water; on cooling the hot watery solution it sets into a jelly (gelatinises); this property is possessed by solutions as weak as 1 p.c., it is lost by prolonged boiling, or instantly by heating to 140° in sealed tubes. Gelatin is insoluble in alcohol, ether, and chloroform. Aqueous solutions are powerfully lævorotatory, the rotatory power being influenced by the temperature and reaction of solution; at 30° [a] = -130 (Hoppe-Gelatin is not precipitated by acetic acid nor by a solution of lead acetate; it may be thus distinguished from chondrin. It is not precipitated by acetic acid and potassium ferrocyanide, nor by the majority of metallic salts which precipitate proteids. Tannic acid even in very dilute solutions precipitates it; it is on the formation of the last-named precipitate that the conversion of hides into leather is brought about.

Schützenberger and Bourgeois ascribe to gelatin the formula $C_{10}H_{124}N_{24}O_{29}$. Hofmeister (H. 2, 315) finds that by heating gelatin it loses water, and is converted into collagen, which he therefore considers an anhydride of gelatin; his

formula for gelatin is $C_{102}H_{151}N_{31}O_{39}$.

By the action of boiling water, gelatin loses after 25 hours its power of gelatinising, and it is split up into two peptone-like bodies, semiglutin (sparingly soluble in 70-80 p.c. alcohol and precipitated by platinum tetrachloride) and hemicollin (soluble in 70-80 p.c. alcohol, not precipitated by platinum tetrachloride). Similar substances are formed by the action of the peptic or pancreatic ferment, but ultimately true gelatin peptones (not precipitable by saturation with ammonium sulphate), leucine, glycocine, and volatile fatty acids are formed. (The foregoing account of gelatin is very largely taken from Gamgee's Physiol. Chem., p. 252.) Many bacteria liquefy gelatine. On digestion of gelatin see Chittenden, J. Physiol. 12, 23, 34. The most recent work on the decomposition products of gelatin has been done by Schützenberger. On heating gelatin at 200° with barium hydroxide, one-fifth of the total nitrogen is converted into ammonia; carbonic and oxalic acids are also formed, these products being in the ratio of the products of decomposition of urea and oxamide. The other products are amidoacids of the acetic series, the most important being glycocine, alanine, amido-butyric acid and leucine, and acids of an homologous series $C_nH_{2n}N_2O_5$, the value of *n* varying from 8 to 10. These latter acids are difficult to isolate; at 100°-120° they are converted into anhydrides; they do not yield derivatives with acetic anhydride, but with ethyl iodide in the presence of an alkali they yield di-ethyl derivatives. The general results lead to the conclusion that gelatin is formed by the combination with elimination of water of 1 mol. urea or oxamide with 2 groups $C_nH_{2n}N_2O_5$ and 4 groups $C_nH_{2n+1}NO_2$, n being 2, 3, 4, or 6, with a mean value of 3.5 (v. Schützenberger a. Bourgeois, C. R. 82, 262; Schützenberger, C. R. 102, 1296). Both gelatin and albumin yield on treatment with alcoholic hydrochloric acid a diazo-compound C_bH₆N₂O₃ (Buchner a. Curtius, B. 19, 850).

The question of the part played by gelatin, which is an easily-digestible substance in nutrition, is very important practically, jellies especially being given to invalids. The question was first investigated by J. Etzinger (N. R. P. 23, 535), and subsequently by Voit (Z. B. 8, 297). Voit's chief result showed that gelatin will not entirely replace proteids, but that animals rapidly waste which are fed on it alone, but, in conjunction with a certain small amount of proteid, it is capable of maintaining nitrogenous equilibrium as well as if the only nitro-genous food taken was proteid in nature. These results have been since very generally confirmed (v. S. Pollitzer, Pf. 37, 301). Voit distinguishes between circulating and organic albumin; gelatin can never yield the latter, but it may replace the former in so far as it prevents the conversion of organic into circulating albumin. Gelatin also diminishes the waste of fat in the body.

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This body forms the chief con-Mucin. stituent of the ground substance of connective tissue, the other organic constituent being a globulin very like serum globulin. Mucin is especially abundant in the 'jelly like' tissue of the umbilical cord, and in the vitreous humour. The cement-substance of epithelia behaves similarly to microscopic reagents (especially silver nitrate, by which it is stained brown owing to a deposit of metallic silver), and is probably of the same chemical nature as the ground substance of connective tissue; both are soluble in alkalis. Mucin forms the chief constituent of mucus, and gives the sliminess to the secretion of mucous membranes. In mucus it is suspended in an alkaline exudation from the blood and nixed with the debris of epithelium cells, and a lew white blood corpuscles. The mucin itself is here formed by the protoplasm of certain cells of the epithelium becoming altered, so that it becomes swollen and brightly refracting; the globule of mucin so formed is discharged, leaving a so-called goblet cell. In mucous glands, such as the submaxillary salivary gland, a very similar replacement of protoplasm by mucin (or mucigen, as it is called when inside the secreting cells) takes place. Mucin is also largely contained in the surface secretion of several invertebrate animals, e.g. the snail. Mucin is contained in submaxillary saliva; the metalbumin and paralbumin (q. v.) of ovarian cysts is a very similar substance. The substance which confers sliminess to the bile and the synovial fluid, formerly considered to be mucin, has been shown to consist of a nucleo-albumin (q. v.).

The methods of preparation of mucin from its various sources differ; from tendon (v. Rollett, Sitz. W. 30, 308; Löbisch, H. 10, 40); from submaxillary gland (Obolensky, Pf. 4, 336; Hammarsten, H. 12, 163); from bile (Gautier, Chimie Appliquée à la Médecine, 2, 126; Paijkull, H. 12, 196). Eichwald, Hammarsten, and others have prepared it from snails (Pf. 36, 373), and Giacosa (H. 7, 40) from the membranes of frog's eggs. These methods depend upon the fact that mucin is soluble in weak alkalis, e.g. lime water or dilute baryta water, and can be precipitated from this solution by acetic acid, in excess of which it is not soluble. Hammarsten finds, however, that submaxillary mucin is easily decomposed by lime water and similar weak alkaline fluids; he, therefore, uses water to extract the mucin. Hammarsten has shown that considerable differences exist in mucin according to its origin; this was previously suspected from the divergencies in elementary composition. Tendon mucin (C, 48.3; H, 6.44; N, 11.75; S, 0.81 p.c., Loebisch) and submaxillary mucin (C, 48.84; H, 6.8; N, 12.32; S, 0.8, Hammarsten) are very much alike in elementary composition, but differ in their reactions; e.g. tendon mucin is not easily decomposed by weak alkalis, nor is it so easily soluble in weak hydrochloric acid as submaxillary mucin. These forms of mucin differ still more from those obtained from Helix pomatia; Hammarsten, indeed, has shown that two distinct varieties of mucin, from the foot and mantle respectively, can be obtained from this snail.

Mucin gives the xanthoproteic, Millon's, and | yeast (Hoppe-Seyler) and mildew; in tea leaves the Adamkieowicz reactions; it is precipitated, | (Kossel), in poppy, earth nut, rape and cotton

but not rendered insoluble, by saturation with sodium chloride or magnesium sulphate, and by alcohol. It is precipitated by acetate of lead, but by no other metallic salt (except the submaxillary mucin, which is precipitated by several), it is not precipitated by tannic acid nor by boiling; acetic acid gives a characteristic stringy precipitate. Mucin yields, when boiled with strong sulphuric acid, leucine and tyrosine, and when boiled with caustic soda pyrocatechin (Obolensky). When boiled with dilute sulphuric acid for a few hours, mucin yields a reducing sugar, but one which is not capable of the alcoholic fermentation, and albumin is also formed. Landwehr regards mucin as a compound of a proteid and a non-reducing carbohydrate ($C_6H_{10}O_8$) which has most of the properties of plant gum; he calls it animal gum (Pf. 39, 193). At one time this carbohydrate was regarded as a variety of glycogen which gave no colour with iodine (achrooglycogen).

Met-albumin and paralbumin. These two proteid-like substances are fairly constantly found in the fluids removed from ovarian cysts (Scherer), and occur occasionally in other cysts and in ascitic fluid also. Metalbumin is a form of mucin (pseudomucin); paralbumin differs from metalbumin by giving a precipitate when it is boiled; it is probably a mixture of pseudomucin with albumin, and can indeed be prepared by mixing these two substances. Both yield reducing sugars when boiled, and both contain animal gum (D. Hammarsten, Maly's Jahresb 11, 11; Landwehr, H. 8, 114; Pf. 39, 193; Oerum, Maly's Jahrsb. 14, 459).

Nuclein. This is the name givento the sub-

stance which composes the nuclei of cells. It was prepared by Lauder Brunton from the red blood corpuscles of birds and snakes; the nuclei were freed from adhering stroma and hemoglobin by repeated agitation with ether and water (Journ. of Anat. and Physiol. 2nd series, 3 91). Nuclein resembles mucin in its solubilities. Plosz found, however, that it contained phosphorus (Hoppe-Seyler, Med. Chem. Untersuchungen, Heft 4, 460), and considered it was identical with the nuclein which Miescher separated from the nuclei of white corpuscles (ibid.). Miescher separated the nuclein from the other constituents of the cells by subjecting them to artificial gastric digestion, nuclein, like mucin, being indigestible by pepsin. Miescher ascribes to it the formula C₂₉H₄₉N₉P₂O₂₂. Nuclein has also been separated from the brain (Jaksch, Pf. 13, 469) to the amount of 0.14 p.c. (Geoghegan, H. 1, 330), and from the liver; it is probably present in all parts where nucleated cells occur. Miescher has also described nuclein as occurring in the yolk of hens' eggs; it has also been described in milk. A. Kossel (H. 10, 248) has shown that the nuclein of yolk and milk differ from that of cell nuclei in containing iron, and by not yielding guanine and hypoxanthine when decomposed at a high temperature by weak acids. Adenine C₈H₈N₅ is described by Kossel as being an intermediate product between cell nuclein and hypoxanthine. present in vegetable cells also, and adenine can It is present in also be obtained from this. yeast (Hoppe-Seyler) and mildew; in tea leaves

cake; also in palm nut, but the proportion P:N is different in the nuclein prepared from this source (Klingenberg a. A. Stutzer, B. C. 1883, Miescher's formula for nuclein must, indeed, be received with caution; there are either several varieties of nuclein-for elementary analyses show great discrepancies-or else, as Worm Müller (Pf. 8, 190) supposes, it is no definite chemical individual, but the different nucleins are mixtures of an organic phosphorus compound with varying quantities of proteids. Recent work on nuclein has shown that this phosphorus compound is nucleic acid (Altmann, Arch. f. Anat. u. Phys. 1888. 524). For artificial nuclein made by adding phosphoric acid to albumin see Liebermann, B. 21, 598; Pohl, H. 13, 292; Malfatti, B. d. Naturwiss. Med. Ver. Innsbruck, 1891-2; Kossel, Verh. physiol. Ges. Berlin, Oct. 21, 1892. See also numerous papers by Kossel, Kruger, and others on adenine and nuclein bases in last few volumes of H. For iron compounds of nuclein see Bunge's Physiol. Chem. This hematogen is probably the normal iron-containing food. The chromatin and plastin of histologists are respectively phosphorus-rich and phosphorus-poor nucleins. For iron in the nucleus see Macallum, P. 49, 488.

Nucleo-proteids. Nucleo-albumin. E. Merck (D. P. J. 261, 316) calls nucleo-proteids substances which, when boiled with water under pressure or treated with acids, alkalis, or ferments, are resolved into nuclein and albumin: for instance, vitellin and casein (v. also Lubavin, Hoppe-Seyler, Med. Chem. Unters. p. 447), Hammarsten gives the name nucleo-albumin to a class of proteid-like bodies which are very like globulins, being precipitated from their solutions by saturation with sodium chloride or magnesium sulphate; they are also precipitated by acetic acid, but, unlike mucin, are soluble in excess. On gastric digestion of a clear solution, an insoluble phosphorus-containing substance, like nuclein, is formed. Many nucleo-albumins have the physical characters of mucin, and the sliminess of bile and synovia is due to such bodies (Hammarsten, H. 12, 173, on synovia; Maly's Jahrsber. 12, 1882, on bile-mucin; Paijkull, H. 12, 196). The chief proteid constituent of all protoplasm is nucleo-albumin (Halliburton's Goulstonian Lectures, Brit. Med. Journ., March 1893). Wooldridge's tissue fibrinogens are also nucleo-albumins (for ref. see above lectures).

Chondrin. This is the substance obtained from cartilage by boiling; the mother substance of chondrin in the matrix of the cartilage is termed chondrigen. Hot aqueous solutions of chondrin gelatinise on cooling like those of gelatin. Aqueous solutions are precipitated by the same reagents that precipitate solutions of gelatin and of mucin. It is strongly laworotatory; its rotatory power in solutions of different strengths has been studied by de Bary (Med. Chem. Unters. i. 71). Very great discrepancies exist between the results of various analyses (see table comparing analyses by Mulder, Fischer a. Bödecker, Schützenberger a. Bourgeois, and v. Mehring in Gamgee's Physiol. Chem. p. 270), hence considerable doubts have been entertained of its being a chemical individual. It gives the reactions both of mucin and gelatin; on being boiled with dilute acids it

yields a lævogyrate reducing sugar, called by de Bary chondriglucose (v. also Fischer Bödecker, A. 117, 111), but is now regarded as identical with that obtained from mucin, Landwehr considering that animal gum is contained in chondrin as in mucin (Pf. 39, 204, 40). Morochowitz was the first to arrive at the conclusion that chondrin is a mixture of gelatin and mucin (Verhandl. d. naturhist, med. Vereins zu Heidelberg, 1, Heft 5); mucin can be extracted from it with lime or baryta water, and pure gelatin is left behind. On this assumption chondrigen = collagen + mucin. It must, however, be mentioned, in opposition to this view, that Hoppe-Seyler (J. pr. 56, 129) and Otto (Z. [2] 4, 628)state that on treating chondrin with dilute sulphuric acid it yields leucine, but no tyrosine or glycosine. The most recent work on chondrin shows that it is chondroitic acid rather than mucin which is present (C. T. Mörner, H. 12, 396; Skand, Arch. Physiol. 1, 210; Krukenberg, Z.B. 20, 307; Schmiedeberg, Arch. exp. Path. u. Pharm. 1891, 355).

Elastin. This substance is the very insoluble material of which the elastic fibres of connective tissue are composed. It may be prepared from the ligamentum nuchæ by boiling it with ether and alcohol to remove fatty matters, then for 36 hours with water to remove the collagen. The residue is boiled with strong acetic acid, and afterwards with concentrated caustic soda till the fibres begin to swell; it is then treated with weak acetic acid, water, and lastly for 24 hours with hydrochloric acid. The acid is removed by washing with water, and the residue has still all the characters of fresh elastic tissue, and has the following percentage composition: C, 55-45; H, 7-41; N, 16-19; O.20-89 (Müller, Zeit.f. Nat. Med. 3, 10, Heft 2). For recent analyses see Chittenden, Z. B. 25, 368.

Elastin is not soluble in any liquid which does not decompose it. It is soluble in boiling concentrated caustic potash, in concentrated nitric and sulphuric acids. It is digestible by both pepsin and trypsin; the former ferment being the more active (A. Ewald a. W. Kühne, Die Verdauung als histol. Methode). Peptones are ultimately formed; there are also intermediate bodies of the nature of albumoses (Horbaczewski, C. C. 1885, 843). Elastin when treated with sulphuric acid yields leucine but no tyrosine.

Keratin. This substance replaces the protoplasm in the cells of certain epidermal structures, viz. the superficial layer of the epidermis itself, nails, horns, hoofs, feathers, and the cuticle and fibrous substance of hairs. It is prepared by successively boiling the tissue with ether, alcohol, water, and dilute acids; the insoluble residue is keratin.

Subjected to the prolonged action of water under pressure at 150°-200° it yields a turbid solution. It is also dissolved by boiling with alkalis, and on the addition of acids to this solution sulphuretted hydrogen is given off, the sulphur in keratin, which varies considerably in amount, being very loosely combined. Horn swells in dilute acetic acid, dissolves in boiling glacial acetic acid, and in nitric acid; it yields aspartic acid, volatile fatty acids, leucine and tyrosine when boiled with dilute sulphuric acid;

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when burnt it gives off a characteristic smell. The following are the chief analyses that have been made (Hoppe-Seyler, *Physiol. Chem.*, Th. 1, p. 90):—

_	Hair	Nails	Horn	Hoof
	(v. Laer)	(Mulder)	(Tilanus)	(Mulder)
C	50·60	51·00	51·03	51·41
H	6·36	6·94	6·80	6·96
N	17·14	17·51	16·24	17·46
O	20·85	21·75	22·51	19·49
S	5·00	2·80	3·42	4·23

Neurokeratin. This is a substance which forms an irregular framework in the medullary sheath of nerve fibres, which resembles keratin in its general behaviour, but differs from it in being less easily soluble in boiling solutions of caustic potash (Ewald a. Kuhne, Verhandl. d. naturhist. med. Vereins zu Heidelberg, vol. i. Heft 5; Kühne a. Chittenden, Z. B. 26, 291). This substance is interesting, as both the epidermis and the nervous system are derived from

the epiblast of the embryo.

Chitin. This substance forms the chief constituent of the skeletal octodermal tissues of invertebrate animals, especially arthropoda. list of situations in which it has been described is given in Gamgee's Physiol. Chem. p. 299; to these must be added the pen of cuttlefishes (Krukenberg), the cartilages and other mesodermal tissues of sepia and the king-crab (Halliburton, Pr. 38, 75). In crustacea it is often impregnated with calcareous matter, and in the odontophore of molluscs with silica. It is prepared from the wing-cases or shells by boiling them with caustic soda. The chitin remains insoluble. It may be dissolved in cold concentrated hydrochloric acid, and the solution precipitated by the addition of water. It is colourless, amorphous, insoluble in water, alcohol, ether, acetic acid, dilute mineral acids, and solutions of the alka-It is dissolved by concentrated mineral acids. (On the solubilities of chitin v. Krukenberg, Z. B. 22, 480.)

The formula for chitin is C₁₅H₂₅N₂O₁₀ (Led-

The formula for chitin is $C_{13}H_{25}N_2O_{10}$ (Ledderhose, H. 2, 213; 4, 139). Berthelot (C. R. 47, 227) stated that it yields a fermentable sugar on boiling it with sulphuric acid; and Sandwick considered it to be an amine derivative of a carbohydrate with the formula $N(C_{12}H_{20}O_{10})$. Ledderhose showed, however, that the reducing substance is a nitrogenous body, glucosamine and when chitin is heated with acids it takes up the elements of water, and yields glucosamine

and acetic acid.

 $2C_{1b}H_{2c}N_2O_{1o} + 6H_2O = 4C_bH_{11}NO_b + 2C_2H_1O_2$. Glucosamine is an amido- derivative of grape sugar $(C_0H_{12}O_0 - OH + NH_2 = C_0H_{18}NO_b)$; it forms salts, of which the hydrochloride is formed by boiling chitin with hydrochloric acid; this is a crystalline substance soluble in water, and in solution dextrorotatory $(\alpha_D = +70^{\circ}C)$. The pure base prepared by the action of barium hydrate on the sulphate of glucosamine crystallises from alcohol in the form of needles. It is not fermentable. (The foregoing account of elastin, keratin, and chitin is largely taken from Gamgee's *Physiol. Chem.*, which see for fuller details.)

Skeletins. This term is applied by Krukenberg (Z. B. 22, 241) to a number of nitrogenous but sulphur-free substances, including conchiolin, spongin, &c., found in the skeletal tissues of invertebrates; they are probably all like chitin amido-derivatives of carbohydrates. The substances are all very insoluble.

Conchiolin $(C_{30}H_{48}N_9O_{11})$ forms the organic basis of the shells of mussels and snails. On decomposition it yields leucine, perhaps glycocine, but no tyrosine nor reducing substance. It does not give the xanthoproteic, Millon's, nor the Adamkiewicz reactions. The cementing substance between the eggs of various molluscs, whose shells and egg capsules contain conchiolin, is coloured red by heating with Millon's reagent, and contains a body allied to keratin. Corneïn (from corals) $(C_{30}H_{41}N_9O_{13})$ differs from conchiolin by giving a red colour with Millon's test.

Spongin, the organic basis of the common sponge, yields as decomposition products leucine and glycocine (Städeler), but no tyrosine. It does not give any of the colour reactions mentioned above; it also resembles conchiolin by yielding on digestion peptone-like substances which differ from true peptones and albumoses by not giving the three colour reactions just mentioned; they thus differ from keratin, which is not digestible.

Fibroin, the chief constituent of insects' cocoons and spiders' threads, behaves to all three tests like an ordinary proteid, and on decomposition yields leucine, glycocine, and tyrosine. Fibroin is soluble when heated in sealed tubes in glacial, acetic, and other organic acids (A. Ledow, Maly's Jahrsbericht, 13, 32). For

silk see Weyl, B. 21, 1407, 1529.

Hyalins and Hyalogens. The term 'hyalin' is applied to the chief constituent of the walls of hydatid cysts. Krukenberg states (Z. B. 22, 261) that the substance is present in the cyst wall as hyalogen, an insoluble substance which by the action of alkalis is changed into hyalin, which is easily soluble in water. Hyalogen is also converted into hyalin by heating with water (under pressure) at 150°C. (For elementary composition v. Lücke, Virchow's Archiv, 19, 189). By heating with sulphuric acid hyalin yields a sugar, probably glucose, which is dextrorotatory and capable of the alcoholic fermentation, and in this hyalin resembles mucin.

Krukenberg has extended the terms hyalogen and hyalin to other similarly related bodies obtained from various sources. The edible bird's nest has properties very like those of mucin (v. J. R. Green, J. Physiol. 6, 40), and is the result of the activity of certain glands described by Bernstein (Journ. Ornithologie, 1859. 111) as being remarkably developed in the nest-building season. Krukenberg finds that it is chiefly composed of a hyalogen (neossine), which yields as its hyalin, neossidine. Chondrosine is a hyalogen, obtained from the sponge Chondrosia reniformis. The vitreous humour from the eyes of oxen and pigs was found to contain a hyalogen, whereas the cornea did not yield one, but was found to consist chiefly of collagen, and a proteid allied to myosin. Spirographin, which largely composes the skeletal tissues of the worm Spirographis is also a hyalogen, but differs from others by yielding when acted on by alkalis not only a hyalin, spirographidin, but also pyrocatechin. The hyalogens are not acted on by gastric juice; some are, some are not attacked by trypsin. W. D. H.

PROTEÏN v. PROTEÏDS.

PROTEOSES v. PROTEÏDS.

PROTO-ALBUMOSE v. PROTEÏDS.

PROTOCATECHUIC ACID C,H,O, $C_6H_3(OH)_2.CO_2H[4:3:1].$ Mol. w. 154. [199°] (Hlasiwetz); [194°] (Barth a. Schmidt, B. 12, 1265). S. 1.9 at 14°; 10 at 60° (Tiemann a. Nagai, B. 10, 211). Occurs in the fruit of Illicium religiosum (Eykman, R. T. C. 4, 47).

Formation.—1. By potash-fusion from piperic acid (Strecker, A. 118, 280), catechin (Kraut, A. 128, 285), maclurin (Hlasiwetz a. Pfaundler, A. 127, 351), guaiac resin (Hlasiwetz a. Barth, A. 130, 346), kino (Stenhouse, C. J. 28, 6), caffeïc acid (Hlasiwetz, A. 142, 219), cinchona-red (Rembold, A. 143, 273), asafætida (Hlasiwetz a. Barth, A. 138, 61), angelica-resin (Brünner, N. R. P. 24, 641), many other resins, the methyl-derivative of sulpho-p-oxy-benzoic acid (Malin, A. 152, 109), sulpho-m- and sulpho-p-oxy-benzoic acids (Barth, C. J. 24, 829), and iodo-p-oxy-benzoic aldehyde (Tiemann a. Herzfeld, B. 10, 213).—2. By heating piperonylic acid with HClAq at 150° (Fittig a. Remsen, A. 159, 129).—3. By the action of Br and water on quinic acid (Hesse, A. 112, 52; 122, 221; Fittig, A. 168, 111).—4. By heating pyrocatechin with water and ammonium carbonate at 140° (Miller, C. J. 41, 400).

Properties.—Tufts of monoclinic needles

(containing aq), sol. water, alcohol, and ether, nearly insol. boiling benzene. FeCl, colours its aqueous solution bluish-green, the colour changing to red on addition of alkalis. FeSO, colours solutions of its salts violet. $Pb(OAc)_2$ gives a pp., sol. acetic acid. Crystallises with p-oxybenzoic acid as (C,H,O,)C,H,O, 2aq, which yields PbC₁₄H₁₀O₇ 2aq (Hlasiwetz, A. 134, 276). Reduces aqueous AgNO, on heating or on adding NH₃Aq. Does not reduce Fehling's solution.

Reactions.—1. Split up by dry distillation into CO₂ and pyrocatechin.—2. Soda-fusion gives 55 p.c. of the theoretical amount of pyrocatechin .- 3. Bromine in the cold forms bromoprotocatechuic acid. Br at 100° yields tetrabromo-pyrocatechin.-4. N₂O₃ passed into the ethereal solution forms oxalic acid (16.5 p.c.), carboxy-tartronic acid (10 p.c.), di-nitro-di-oxyquinone (5 p.c.), trinitrophenol (4 p.c.), (a)-dinitro-phenol [114°] (3·5 p.c.), and nitro-p-oxybenzoic acid (1 p.c.) (Gruber, B. 12, 514).—5. H₂SO₄ (25 pts.) at 140° forms a very small quantity of tetra-oxy-anthraquinone (rufiopin) (Nölting, Bl. [2] 37, 395).—6. Heating with As O at 160° or exposing a solution in Na CO Aq to the air forms catellagic acid C₁₄H₁₀O₂? which resembles ellagic acid and forms an orange solution in nitric acid.

Salts.—BaA'₂5aq.—Ba₃(C,H₃O₄)₂ (dried at 130°) (Barth, A. 142, 246).—CaA'₂4aq.—CaA'₂3aq.—Pb₃A'₂O₂.—PbA'₂2aq.

Acetyl derivative C₆H₃(OAc)₂.CO₂H.

[153°]. Crystals (Herzig, M. 6, 872).

Methyl ether Mah'. [134·5°]. Needles.

Ethyl ether Eth'. [134° cor.]. Prisms
(Matsmoto, B. 11, 129).

m-Methyl derivative $C_6H_3(OH)(OMe).CO_2H[4:3:1].$ Vanillic acid. [207°]. S. ·12 at 14°; 2·5 at 100°. Formation (Tiemann, B. 8, 509, 1123; 9, 52, 419; 10, 59, 202; 11, 122).—1. By exposing moist powdered vanillin to the air.—2. By oxidising coniferin with dilute KMnO₄.—3. From its acetyl derivative, which is got by the action of KMnO₄ on acetyl-eugenol, acetyl-ferulic acid, and acetylcreosol.-4. Together with isovanillic acid by heating C₆H₃(OMe)₂CO₂H with HClAq at 140°. Properties.— Needles (from water), v. e. sol. alcohol, m. sol. ether. Has no smell. May be sublimed. Gives no colour with FeCls. Reactions.—1. Split up by HClAq (S.G. 1.1) at 150° into protocatechuic acid and MeI.-2. Potashfusion yields protocatechuic acid.—3. By heating with MeI (2 mols.) and KOH (2 mols.) it is converted into C₆H₃(OMe)₂.CO₂Me.—4. Yields guaiacol on distillation with lime.—5. The Ca salt yields vanillin on distilling with calcium formate.

Methyl ether of the m-Methyl deriva-

tive MeA'. [63°]. (286°).

Ethyl ether of the m-Methyl derivative EtA'. [44°]. (292°).

Acetyl derivative of the m-Methyl

derivative C_eH₃(OAc)(OMe).CO₂H. Needles (from dilute alcohol).

Benzoyl derivative of the m-Methyl derivative. [178°] (Tiemann, B. 15, 2068).

p-Methyl derivative
C₀H₃(OMe)(OH).CO₂H [4:3:1]. Isovanillic ac

Isovanillic acid. [250°]. S. 06 at 15°; 7 at 100°. Formed with other products by heating protocatechuic acid with MeI and KOH at 150°. Got also by heating hemipic acid with HClAq (Matthiessen a. Foster, A. Suppl. 2, 378), and, together with vanillic acid, by digesting C₆H₃(OMe)₂CO₂H with dilute HClAq at 140° (Tiemann). The acetyl deriva-tive is got by oxidation of the acetyl derivative of isoferulic acid (hesperetic acid) by KMnO, (Tiemann a. Will, B. 14, 963), and likewise by oxidation of the acetyl derivative of betelphenol C₆H₃(OMe)(OH).C₃H₅ [4:3:1] (Bertram a. Gildemeister, J. pr. [2] 39, 349). Prisms, v. sol. alcohol and ether. May be sublimed. Its aqueous solution is not coloured by FeCl₃. Yields a nitro- derivative [173°] and an acetyl derivative C_sH_s(OMe)(OAc).CO₂H [207°]

Di-methyl derivative C₆H₂(OMe)₂.CO₂H. Veratric acid. [174°] (T.); [181°] (G.). S. 05 at 14°; 6 at 100°. Occurs in small quantity in sabadilla seeds (Merck, A. 29, 188). Formed by heating at 140° protocatechnic acid (1 pt.) with MeI (4 pts.) and KOH (1 pt.) dissolved in MeOH (Kölle, A. 159, 241). Produced also by oxidation of C_sH_sMe(OMe)₂ (Tiemann, B. 8, 1138), of methyl-eugenol (Graebe, A. 158, 282), of pspaverine (Goldschmidt, M. 6, 378), and of di-methyl-caffere acid (Tiemann a. Will, B. 14, 962). Formed also by saponifying its amide, which is made by the action of Cl.CONH, on C₈H₄(OMe)₂ in CS₂ in the presence of AlCl₄ (Gattermann, A. 244, 71). Veratric acid is a product of the action of alcoholic sods on pseudaconitine, on veratrine (Wright a. Luff, C. J. 33, 160, 353), and on hexa-methyl-quercetin (Herzig, M. 5, 83), and of fused potash on papaveraldine (Goldschmidt, M. 7, 493). Veratric acid is also got by the action of KOH on pseudoopianic acid (W. H. Perkin, jun., C. J. 57, 1068).

Properties.—Slender needles (containing aq). From hot solutions (above 50°) it separates in anhydrous crystals. V. e. sol. alcohol and ether. FeCl, gives no colour. Distillation with lime forms C₆H₁(OMe)₂. Potash-fusion yields protocatechuic acid. HIAq at 160° forms MeI and protocatechuic acid. HClAq forms vanillic and also iso-vanillic acids .- Salts .- NaA' 2aq. BaA'2 6aq. - AgA'. - Ethers. - MeA' [60°] (c. 300°).—EtA'. [44°]. (296°).

Di-ethyl derivative C.H. (OEt)2.CO2H. [166°]. Formed by saponifying its ether, which is made from protocatechuic acid, alcoholic potash, and EtI (Kölle, A. 159, 240). Got also by heating hexa-ethyl-quercetin with alcoholic potash at 150° (Herzig, M. 5, 78). Needles (from alcohol).—KA'½aq.—BaA'23aq.—AgA'.—Ethyl ether EtA'. [57°]. Crystals.

Methyl-ethyl derivative C₅H₃(OMe)(OEt).CO₂H [3:4:1]. [194°]. Formed by oxidation of ethyl-vanillin (Tiemann, B. 8, 1180), of ethyl-eugenol (Wassermann, A. 179, 379; Balbiano, G. 11, 416), and of di-ethyl-eur-eumin (Jackson a. Mencke, Am. 4, 90). Needles, almost insol. cold water.—BaA'₂ 4aq: needles.

Methyl-propyl derivative $C_nH_n(OMe)(OPr).CO_nH[3:4:1]$. Formed by oxidation of the propyl derivative of eugenol (Cahours, Bl. [2] 28, 314). Needles.

Methyl-carboxymethyl derivative C₆H₃(OMe)(O.CH,CO₂H).CO₂H [3:4:1]. [256°]. Formed from C₆H₃(OMe)(O.CH₂:CO₂H).CHO by oxidation (Elkan, B. 19, 3056). Yields CuA" as a green insoluble pp.

Methylene derivative CH₂:O₂:C₆H₃.CO₂H. Piperonylic acid. [228°]. Occurs in para-coto bark (Hesse a. Jobst, A. 199, 63). Prepared by oxidation of piperonal (Fittig a. Mielck, A. 152, 40). Got by heating protocatechuic acid with potash and CH₂I₂ (Fittig a. Remsen, A. 168, 94). Piperonylic acid is also formed by the action of KMnO, on cubebin, on methystic acid (Pomeranz, M. 8, 468; 10, 790), on safrole (Eykman, R. T. C. 4, 39; Poleck, B. 19, 1096), and on the (a)-dihydride of piperic acid (Regel, B. 20, 415). Needles (from alcohol). May be sublimed. Nearly insol. cold water, m. sol. boiling alcohol. FeCl, gives a brown pp. in neutral solutions. Dilute HClAq at 170° forms protocatechnic acid. HNO, forms nitro-piperonylic acid [172°]. Salts.—NaA' aq.—KA'aq.—BaA', aq.—CaA', 3aq. S. 625 at 15°.—PhA', aq.—CuA', aq.—AgA'. Quinine salt -PbA'₂ aq. -CuA'₂aq. -AgA'. Quinine salt C₂₀H₂₄N₂O₂HA'aq: needles, sol. hot water. Cinchonidine salt C₁₀H₂₂N₂OHA': needles (Hesse, A. 243, 147). — Ethyl ether EtA'. Oil.— Nitrile CH₂O₂:C₆H₃:CN. [95°]. Formed from the oxim of piperonal and Ac₂O (Marcus, B. 24, 2656). 3656). Needles, v. sol. alcohol. Alcoholic hydroxylamine at 60° forms the amidoxim CH₂O₂:C₆H₈.C(NH₂):NOH [151°] which yields B'HCl [193°], and is converted by Ac₂O into $CH_2O_2:C_6H_3.C < NO > CMe [110°].$

Ethylene derivative C2H4O2:C6H3.CO2H. [184°]. Formed by heating protocatechuic acid with ethylene bromide and KOH (Fittig, Z. [2] 7, 289; A. 168, 99). Needles (from hot water). May be sublimed. POL, yields a product whence water reproduces ethylene-protocatechuic acid.

PCl₅ at 130° followed by water yields the acid C₂H₂Cl₂O₂:C₆H₅.CO₂H [121°].—BaA'₂ 2aq.—CnA'₂ 2aq: monoclinic crystals.—Ethyl ether

Di-methyl-ethylene ether $C_2H_4(O.C_6H_2(OMe).CO_2H)_2$. Formed by oxidising the ethylene ether of eugenol with KMnO. (Cahours, Bl. [2] 29, 270). Amorphous.

Reference.-NITRO-PROTOCATECHUIC ACID.

Homo-protocatechuic acid v. DI-OXY-PHENYL-ACETIC ACID.

Diprotocatechuic acid C₁₂H₄(OH)₄(CO₂H)₂. Formed by potash-fusion from divanillin (Tiemann, B. 18, 3494). Amorphous, sl. sol. water and alcohol. Coloured bluish-green by FeCl,

PROTOCATECHUIC ALDEHYDE C, H, O, i.e. C₆H₃(OH)₂.CHO [4:3:1]. Mol. w. 138. [150°].

Formation. -1. From piperonal by successive treatment with PCl_s and water at 100° (Fittig a. Remsen, Z. [2] 7, 100; A. 159, 148; 168, 97).— 2. By heating piperonal with dilute HClAq at 200° (F. a. R.).—3. By heating a solution of pyrocatechin in dilute NaOH with chloroform (Tiemann, B. 9, 1269; 14, 2020).-4. By heating vanillin with dilute HCl at 200° (Tiemann, B. 7, 620).-5. By heating opianic acid with dilute HCl at 170° (Wegscheider, M. 3, 792).

Properties.—Flat needles (from water), v.sol. alcohol, ether, and hot water. FeCl, colours the aqueous solution green, changing to red on addition of Na₂CO₃. Oxidised by KMnO₄ and by potash-fusion to protocatechuic acid. Gives a mirror with ammoniacal AgNOs. Combines with

NaHSO3.

C,H,O, m-Methyl derivative. C₆H₃(OMe)(OH).CHO. Vanillin. Mol. w. 152. [81°]. (285°). S. 1 at 14°; 5 at 80° (Tiemann a. Nagai, B. 10, 211). The fragrant constituent of the pods of Vanilla aromatica (Gobley, J. 1858, 534; Stokkebye, J. 1864, 612). Occurs also in the seeds of Lupinus albus (Campani a. Grimaldi, G. 17, 545), in raw beet-root sugar (Weger, D. P. J. 237, 146; Scheibler, B. 13, 335; Lippmann, B. 13, 662), in gum benzoïn from Siam (Jannasch a. Rump, B. 11, 1635), and in small quantity in wood (Singer, M. 3, 409). Formation.-1. Together with isovanillin, by heating guaiacol with chloroform and NaOHAq (Reimer, B. 9, 424; Tiemann, B. 14, 2023).-2. By oxidation of coniferin or coniferyl alcohol with chromic acid mixture (Tiemann, B.7, 613) .-3. By oxidation of eugenol by KMnO, (Erlenmeyer, B.9, 273).—4. By heating calcium vanillate with calcium formate (Tiemann, B. 8, 1124).-5. By the action of chloroform and potash on vanillic acid (Tiemann, B. 9, 1280).—6. By the action of emulsin or of boiling dilute acids on glucovanillin (Haarmann a. Reimer, C. J. 46, 1348). -7. From C₆H₃(OMe)(NO₂).CHO by reduction followed by the diazo- reaction (Ulrich, B. 18, 2573). Properties.—Monoclinic needles, v. sol. alcohol, ether, CHCl₂, and CS₂; v. sl. sol. cold, v. sol. hot, ligroïn. Smells and tastes like vanilla. May be sublimed. Acid in reaction, and decomposes carbonates. FeCl, gives a blue colour. Boiling aqueous HgCl₂ containing KNO₂ gives a violet colour (Nickel, Fr. 28, 247). Reduces silver solution. Oxidised by moist air to vanillic acid C₅H₅O₄. Reactions.—1. Bromine forms C₅H₇BrO₅ [161°].—2. Dilute HCl at 190° yields MeCl and protocatechuic aldehyde .-8. Potash-fusion gives protocatechuic acid.-4. Sodium-amalgam reduces it in alcoholic solution to vanillyl alcohol C₈H₁₀O₂ as a yellowish oil, and also hydrovanilloin C16H18O6 [c. 225°] (Tiomann, B. 8, 1123).—5. By heating with di-methyl-aniline and $ZnCl_2$ it is converted into $C_{21}H_{28}N_2O_2$ [136°] (Fischer a. Schmidt, B. 17, 1895)—6. Vanillin (1 mol.) mixed with pyrogallol (2 mols.) is converted by conc. HClAq into 'pyrogallo-vanillein' C20H18O8, which forms colourless crystals, insol. water, v. sol. alcohol, giving isomeric bluish-violet crystals by the further action of HCl (Etti, M.3, 637) .-7. Vanillin (1 pt.) boiled with diacetonamine acid oxalate (1 pt.) forms the oxalate of vanillodiacetonamine C₁₄H₁₉NO₃ (v. vol. i. p. 28).—8. Heating with Ac O forms the acetyl derivative of vanillin, and also C₈H₂(OMe)(OAc).CH(OAc)₂[89°] crystallising in six-sided tables (Tiemann a. Nagai, B. 8, 1143).-9. Boiling aqueous FeCl, forms divanillin C₁₆H₁₄O₆ [304°], whence NaOEt and MeI form $C_eH_2(OMe)_2(CHO).C_eH_2(OMe)_2(CHO)$ [138°] (Tiemann, B. 18, 3493).—10. Chloro-acetic acid and KOHAq form C₆H₃(OMe)(OCH₂CO₂H).CHO [188°] (Elkan, B. 19, 3055). Salts.—*C_sH,NaO_s. Needles (from alcohol), sl. sol. NaOHAq.— Hg(C₈H₇O₃)₂: crystals, sl. sol. cold water, insol. alcohol.—Zn(C₈H₂O₃)₂: crystalline pp. Estimation.—Vanilla pods are extracted with ether, the extract shaken with a nearly saturated solution of NaHSO_s, the aqueous layer decomposed by H.SO., the vanillin thence extracted by ether, and the extract evaporated (Tiemann a. Haarmann, B. 8, 1115).

Acetyl derivative of vanillin C₈H₃(OMe)(OAc).CHO. [77°]. Formed slowly by the action of an ethereal solution of Ac₂O on sodium-vanillin in the cold (Tiemann a. Nagai, B. 11, 646). Flat needles. Combines with bi-

sulphites.

Oxim of vanillin $C_sH_s(OMe)(OH).CH:NOH.$ [122°] (Tiemann a. Kees, B. 18, 1664; cf. Lach, B. 16, 1786).

Phenyl-hydrazide of vanillin C_uH_u(OMe)(OH).CH:N₂HPh. [105°]. Plates.

Glucoside of vanillin $C_sH_s(OMe)(OC_sH_{11}O_s).CHO.$ Glucovanillin. [192°]. [a]_D = -88·63 at 20°. Prepared by slowly adding CrO_s (8 pts.) in water (100 pts.) to coniferin (10 pts.) dissolved in water (200 pts.), and leaving the mixture to stand for 5 days (Tiemann, B. 18, 1596, 1661). Colourless needles (containing 2 aq), v. sol. water, m. sol. alcohol, insol. ether. Readily split up by dilute H_sSO_s or by emulsin into vanillin and glucose. Does not reduce Fehling's solution in the cold. Yields $C_sH_s(OMe)(OC_sH_{11}O_s).CH:N_2HPh [c. 195°]$ and a hevorotatory oxim [152°], crystallising in slender yellow needles (containing aq).

p-Methyl-derivative
C.H₃(OMe)(OH).CHO[4:3:1]. Isovanillin. [116°].
Formed by oxidising acetyl-isoferulic acid with
KMnO, and saponifying the resulting acetyl
derivative (Tiemann a. Will, B. 14, 968). Formed
also by heating opianic acid (4 pts.) with water
(30 pts.) and HClAq (8 pts. of S.G. 1·17)
(Wegscheider, M. 3, 789). Monoclinic pyramids,
sl. sol. cold water, v. sol. alcohol and ether. May
be sublimed. When warm it smells somewhat
like vanillin. Its alkaline solutions are yellow.

FeCl, does not colour the aqueous solution. Reduces boiling ammoniacal AgNO. Forms a very soluble compound with NaHSO.

Di-methyl derivative C₃H₃(OMe)₂CHO. Methyl-vanillin. [43°]. (c. 283°). Formed from potassium vanillin and MeI (Tiemann, B. 8, 1135), and also by distilling opianic acid with soda-lime (Beckett a. Wright, C. J. 29, 164). Needles, v. sl. sol. hot water, v. sol. alcohol and ether. Smells like vanilla.

Methyl-ethyl derivative C₈H₃(OMe)(OEt).CHO. [65°]. Formed by boiling potassium-vanillin with EtI and alcohol (Tiemann, B. 8, 1129). Prisms, v. sl. sol. hot

water. Smells like vanilla.

Methylene $C_8H_6O_8$ i.e. Mol. w. 150. derivativeCH₂O₂:C₆H₃.CHO. *Piperonal*. Mol. w. 150. [37]. (263°). S. 2 in the cold. Formed by oxidation of piperic acid in neutral solution by KMnO, (Fittig a. Mielck, A. 152, 35). Formed also by oxidation of the (a)-dihydride of piperic acid by alkaline KMnO₄ (Regel, B. 20, 415). Transparent prisms (from water), m. sol. hot water, v. sol. alcohol and ether. Smells much like coumarin. Forms a crystalline compound with NaHSO₃. Not affected by aqueous alkalis. Reactions.-1. By oxidation, or by boiling with alcoholic potash, it is converted into piperonylic acid CH2O2:C6H3.CO2H.-2. Reduced by water and sodium-amalgam to piperonyl alcohol $C_8H_8O_8$, hydropiperoin $C_{18}H_{14}O_8$ [202°], and iso-C₈H₈O₈, hydropheroin C₁₈H₁₄O₅ [202], and 180-hydropheroin [138°]. AcCl converts the hydropheroins into C₁₈H₁₂Cl₂O₄ [198°] (Remsen a. Fittig, Z. [2] 6, 97; A. 159, 129).—3. An alcoholic solution of KCy forms piperonyloin CH₂O₂:C₈H₈.CH(OH).CO.C₈H₁:O₂CH₂ [120°], separating from alcohol in yellow crystals (F. M. Perkin C. J. 59, 164). A Vary dilute HClAs. Perkin, C. J. 59, 164).-4. Very dilute HClAq at 200° forms protocatechuic aldehyde.—5. PCl, forms liquid CH₂O₂:C₆H₃.CHCl₂ (c. 235°) and C.H.Cl.O., whence cold water forms dichloropiperonal C.H.Cl.O. [90°].—6. Gaseous HCl, passed into a mixture of phenyl mercantan and piperonal, forms $CH_2O_2:C_0H_3.CH(SPh)_2$ [48°] (Baumann, B. 18, 886).—7. Alcoholic ammonia, in presence of some HCy, forms $C_{24}H_{18}N_2O_{e}$, crystallising in prisms [213°], insolalcohol and ether. Piperonal, heated with alcoholic NH, at 70°, forms an isomeric body [172°], crystallising in yellow needles, sol. hot alcohol, insol. water and ether (Lorence, B. 14, 791).-8. Aqueous HCy at 65° forms a compound whence an alcoholic solution of NH, yields CH2O2:C6H3.CH(NH2).CN, converted by boiling HClAq into C, H,O,.—9. Antline forms, on heating, C14H11NO2, crystallising in colourless needles [65°] (L.).—10. p-Phenylene-di-methyl-diamine gives C_eH₃(O₂CH₂).CH:N.C_eH₄NMe₂ [110°] (Nuth, B. 18, 575).

Oxim of piperonal CH₂O₂:C₈H₄.CH:NOH. [110°]. Needles, v. sol. alcohol (Marcus, B. 24, 3656).

Phenyl-hydrazide of piperonal CH₂O₂:C₄H₃.CH:N₂HPh. [100°] (M.); [103°] (Rudolph, A. 248, 103). Yellow needles.

PROTOPINE C₂₀H₁₀NO₅. [202°]. An alkaloid occurring in very small quantity in opium, and obtained from the mother-liquors after the separation of morphine by Gregory's process (Hesse, Z. [2] 7, 653; A. Suppl. 8, 818). Separated from cryptopine by ppn. of the solution of

the mixed hydrochlorides with conc. HClAq, the protopine salt adhering to the sides of the vessel. Crystalline powder (from alcohol), insol. water, sl. sol. hot alcohol, m. sol. chloroform. The alcoholic solution is alkaline in reaction. Sl. sol. ether, separating in spherical groups of prisms. Sl. sol. KOHAq and NH₃Aq. FeCl₃ gives no colour. HNO₃ forms a colourless solution, which turns yellow on warming. H₃SO₄ containing ferric sulphate gives a dark-violet solution, turned dirty brownish-green at 150°. The salts have a bitter taste, and do not gelatinise.— B'₂H₂PtCl₆ 2aq: yellow crystalline pp.

PRUSSIAN BLUE v. FERRIC FERROCYANIDE, vol. ii. p. 334; and Potassium-ferrous ferri-

CYANIDE, vol. ii. p. 339.

PRUSSIC ACID v. CYANHYDRIC ACID, vol. ii. p. 300.

PSATYRIN v. HARTIN.

PSOROMIC ANHYDRIDE $C_{20}H_{11}O_{9}$? [264°]. Occurs in *Psoroma crassa*, a lichen growing in Sicily (Spica, G. 12, 431). Needles, sol. alcohol and ether, insol. benzene. Yields $C_{20}H_{13}AgO_{10}$ as a flocculent pp.

PSYCHOSINE. A substance got, according

to Thudichum (J. pr. [2] 25, 19), from brain. PTEROCARPIN $C_{20}H_{16}O_{6}$. [152°]. [a]₁ = -211° 14 · 6 p.c. chloroform solution. Obtained by mixing powdered sandal wood (Pterocarpus santalinus) with slaked lime, and extracting with ether. The residue is crystallised from alcohol, and the homopterocarpine dissolved in CS₂, which leaves the pterocarpine (Cazeneuve a. Hugounenq, A. Ch. [6] 17, 115). Square tables (from CHCl₃), insol. water, m. sol. hot alcohol and hot CS₂. Insol. acids and conc. KOHAq, even on boiling. Bromine forms $C_{20}H_{15}BrO_{6}$, crystallising in yellowish needles.

Homopterocarpin $C_{24}H_{24}O_8$. [82°-86°]. [a]₁ = -199° . Obtained as above (C. a. H.). Long needles (from boiling alcohol), insol. water,

sol. ether, v. sl. sol. cold alcohol.

Reactions.—1. Bromine gives $C_{24}H_{23}BrO_6$ and $C_{24}H_{18}Br_6O_6$ [270°].—2. On distillation it is largely split up into phenols resembling creosote, and a small quantity of pyrocatechin.—3. Distillation over zino-dust gives benzene, toluene, ethylene, and CO.—4. HCl gives MeCl and a resin.—5. HI gives MeI.—6. Potash-fusion gives phloroglucin.—7. Fuming HNO₈ gives oxalic acid and tri-nitro-orcin [162°]. Not acted upon by sodium-amalgam, phenyl-hydrazine, and Ac₂O.

sodium-amalgam, phenyl-hydrazine, and Ac.O. PTOMAINES. The name 'ptomaines' was first applied to poisonous organic bases obtained from dead bodies $(\pi\tau\hat{\omega}\mu\alpha$, a corpse), but is now usually given to poisonous organic bases formed in the putrefaction of any kind of animal matter. In extracting ptomaines no reagents that would decompose proteids may be used. The bases may be extracted by alcohol, and purified by ppn. with phosphomolybdic acid. In presence of a little HCl the ptomaines are fairly stable, and the solution can then be evaporated and the hydrochlorides extracted from the syrupy residue by absolute alcohol. By this method neuridine hydrochloride can readily be obtained. By the graphs: Ueber Ptomaine, Berlin, 1885; cf. Gautier, Bl. [2] 48, 10) obtained tri-methyl-vinyl-ammonium hydroxide (neurine) C,H₁₈NO or C.H. NMe.OH, muscarine C.H. NO. or

CH(OH)₂.CH₂.NMe₂OH (vol. iii. p. 444), a base isomeric with ethylene-diamine, neuridine O₅H₁₄N₂, gadinine C₇H₁₇NO₂, triethylamine, dimethylamine, and trimethylamine. Garcia (H. 17, 543) got hexamethylene-diamine C₆H₁₆N₂. Herring-pickle contains choline, NMe, NH2Me. By the putrefaction of the herring NMe,, NH2Me, cadaverine C, H16N2, putrescine C4H12N2, and gadinine C,H,,NO2 are formed (Bocklisch, B. 18, 1922). Gautier and Etard (C. R. 94, 1600) from putrid mackerel and putrid horse-flesh obtained a collidine dihydride C₈H₁₃N, parvoline C₉H₁₃N, and a base C₁₇H₃₈N₄. In the putrefaction of proteids poisonous bases are formed in the first five or six days, and are destroyed by further progress of putrefaction. The highly poisonous methyl-guanidine is formed by the action of putrefactive bacteria on beef-broth at 38° (Bocklisch, B. 20, 1441). From human corpses Brieger obtained tri-methyl-oxyethyl-ammonium hydroxide C₅H₁₅NO₂ or CH₂(OH).CH₂.NMe₃.OH (choline or neurine, vol. iii. p. 498), neuridine C₅H₁₄N₂, cadaverine C₅H₁₆N₂, putrescine C₄H₁₂N₂, saprine $C_3H_{14}N_2$, trimethylamine, methyl-guanidine, tetanine $C_{13}H_{20}N_2O_4$, mydine $C_3H_{11}NO$, mydatoxin C.H.3NO2, and mydaleme (Brieger, loc. cit.; cf. Selmi, Rend. Accad. Sci. Bologna, 1872; Schwanert, B. 7, 1332; Guareschi a. Mosso, J. pr. [2] 27, 428; 28, 504; G. 13, 493; Beckurts, Ar. Ph. [3] 14, 1041). Ptomaines are often produced in animal bodies which, after brief exposure, have been excluded from the air; e.g. corpses, sausages, and tinned meat. A crystalline ptomaine can be obtained from the body after arsenical poisoning (Husemann, Ar. Ph. [3] 16, 169; 19, 415). The alcoholic solution of hydrochlorides of ptomaines may be ppd. by an alcoholic solution of HgCl2; after twenty-four hours' standing, the pp. is boiled with a large quantity of hot water, which leaves the compounds of peptones and albuminates undissolved. The hot filtrate deposits the mercury double salt of choline (neurine) while the mother-liquor contains the remaining bases. Picric acid added to an aqueous solution of the hydrochlorides ppts. neuridine picrate, while the mother-liquor de-posits, on evaporation, broad needles of choline picrate C₅H₁₈NOC₆H₂(NO₂)₃OH. Cadaverine and putrescine can be separated by means of their aurochlorides. Saprine is isolated by means of its platinochloride. To distinguish a ptomaine from a vegetable alkaloid, Brouardel a. Boutmy (C. R. 92, 1056; cf. Tanret, C. R. 92, 1163) add a few drops of the solution of the sulphate of the alkaloid to potassium ferricyanide mixed with FeCl₃: a dark-blue pp. is formed if a ptomaine be present, while the vegetable alkaloids (except morphine, veratrine, eserine, aconitine, and ergotinine) have no action. The test is, of course, given by many other reducing agents. This ptomaine reaction is given (slowly) by a poisonous liquid alkaloid extracted by Stas's method from the intestines of persons who died of cholera (Villiers, C. R. 100, 91). Pouchet (C. R. 97, 1560; cf. Gautier, B. C. 1882, 710; Béchamp, C. R. 94, 973) obtained from urine and fæces some alkaloids closely resembling the ptomaines got by putrefaction of proteids out of contact of air. He separated them by decomposing their tannates with Pb(OH)₂ in presence of alcohol, the lead being finally removed by H2S, and the liquid dialysed. C3H3NO2 is liquid and dialyses with difficulty; it exhibits the alkaloidal reactions, is resinified by HCl, and reduces platinic chloride. The base C7H12N4O2 or C₁H₁₄N₄O₂ is crystalline, and passes through the membrane; it is insol. ether, nearly insol. alcohol, is feebly alkaline and yields crystalline salts. From the dialysate the platinochlorides of two crystalline unstable bases $C_5H_{12}N_2O_4$ and $C_7H_{18}N_2O_6$ can be got. Urine which had a tendency to deposit cystin and the fæces of the same patient contained cadaverine and putrescine (Baumann, H. 13, 562; cf. Dupré a. Bence-Jones, Pr. 15, 73; Stadthagen a. Brieger, Ar. pathol. Anat. 115, pt. 3). These diamines appear to be absent from normal urine, but present in cholera.

Cadaverine is Pentamethylene-diamine, vol.

iii. p. 305.

Putrescine is Tetramethylene-diamine.

Mydaleïne is extremely poisonous, while

gadinine is not poisonous.

Mydine C₈H₁₁NO. Occurs in putrid corpses (Brieger, *Ptomaine*, iii. 25). Has an ammoniacal odour and reduces gold chloride. Not poisonous. -B'C₆H₃N₃O₇. [195°]. Broad prisms. **Base** C₇H₁₇NO₂. Occurs in decaying horse-

flesh, four months old (Brieger). Acid in reaction. Poisonous. Does not form a picrate.— B'HAuCl₄. [176°]. Needles or plates. Typhotoxin C₂H₁₇NO₂. Formed by the action

of the typhus bacillus on muscle (Brieger, Ptomaine, iii. 86). Poisonous base.—B'HAuCl4. [176°]. Prisms.

Mydatoxin C₈H₁₉NO₂. Occurs in decaying horse-flesh and corpses (B.). Poisonous. Strongly alkaline.—B'₂H₂PtCl₆. [193°]. V. e. sol. water. Tetanine C₁₃H₂₀N₂O₄. Occurs in decaying

corpses (Brieger, B. 19, 3120), and is formed by the action of the tetanus bacillus on beef. Very poisonous base.—B'H₂PtCl₈. Plates.

Base C₃H₁₁N. (c. 100°). Accompanies te-

tanine, and also produces convulsions (B.). Volatile liquid. Its hydrochloride is crystalline [205°].—B'HAuCl₄. [130°].—B'₂H₂PtCl_e. Plates, decomposed at 240°.

Base C₃₂H₃₁N. Oily ptomaine, smelling like hawthorn (Delezinier, Bl. [3] 1, 178). In presence of air it acts chemically and physiologically like veratrine. Insol. water, sol. alcohol and ether. Its salts are deliquescent.

Base C₅H₁₁NO₂. [156°]. Occurs in putrid flesh and putrid fibrin (E. a. H. Salkowski, B. 16, 1192). Crystalline powder, v. e. sol. water. Not poisonous. -B'HCl: crystals, v. e. sol. water. -B'HAuCl, aq: yellow crystals.

Base C11H1,NOs. Erysipeline. Occurs in urine in erysipelas (Griffiths, C. R. 115, 667). Prisms, sol. water. Very poisonous.

Base C22H19NO2. Occurs in urine in puerperal fever (G.). Crystalline and poisonous.

Base C₁₀H₁₇N. A product of the decomposition of albumen by Bacterium allii, which is found in decaying onions (A. B. Griffiths, C. R. 110, 416). Minute deliquescent needles (from water), smelling like hawthorn.-B'2H2PtCls: crystalline, sol. hot water.

Base C_{1.}H₁₀N₂O₆. Occurs in urine of victims to glanders (A. B. Griffiths, C. R. 114, 1382). White crystals. Poisonous. Forms crystalline

salts.

Base $C_{20}H_{24}N_2O_3$. Occurs in urine in cases of pneumonia (G.). Minute needles, forming an alkaline aqueous solution $[a]_D = 23.5^{\circ}$.

Base C₈H₁₁N. (202°). S.G. 2 9865. Extracted, together with a base C10H15N from putrid cuttlefish (De Coninck, C. R. 106, 858, 1604; 108, 58, 809; 110, 1339; 112, 584). Mobile, stronglysmelling liquid, v. sol. water, alcohol, and ether. Turns brown in air. Appears to be a propylpyridine, as it yields nicotinic acid on oxidation by dilute KMnO₄ at 90°.—B'HCl: deliquescent radiating mass.—B'.H.PtCl₆. Converted by hot water into B'.PtCl₄, a nearly insoluble brown powder.—B'HAuCl₄. — B'HBr. — B'₂H₃HgCl₄.— B'₂H₂Hg₃Cl₈. — B'MeI. Needles, v. e. sol. alcohol.

Base C₁₀H₁₅N. (230°). Formed as above. Liquid, smelling like furze. Resinified by air.— B'HBr. White deliquescent needles.—B'HCl.-B'₂H₂PtCl₄. — B'₂PtCl₄. [206°]. — B'HAuCl₄: yellow pp., decomposed by hot water.

Base C₁₀H₁₃N. (200°). Extracted by Stas's method from fibrin that has putrefied for two months (Guareschi a. Mosso, J. pr. [2] 27, 428; J. Th. 1887, 487). Yields the alkaloidal reactions. Smells like pyridine. Acts physiologi-

cally like curare.— $B_2'H_2PtCl_6$: rose-red crystals. Base $C_{14}H_{20}N_2O_4$. [250°]. Formed in the putrefaction of fibrin, from which it can be extracted, together with the liquid alkaloid C10H18N, by chloroform and ether (Guareschi, G. 17, 509). Tables (from alcohol), sol. water and alcohol, v. sl. sol. chloroform. Its aqueous solution is neutral. Griffiths (Bl. [2] 7, 250; C. R. 113, 656) got, from urine of patients suffering from erysipelas, a crystalline ptomaine 'erysipeline C11H13NO3, sol. water, and very poisonous. It gives the alkaloidal reactions.

The term leucomaines (λεύκωμα, white of egg) is applied by Gautier (Sur les Alcaloïdes des Tissus Animaux, Paris, 1886) to alkaloids occurring in the tissues of living animals. From fresh beef and Liebig's extract of beef he obtained xanthocreatinine C5H10N4O, crysocreatinine C₃H₈N₄O, amphicreatinine C₆H₁₉N₇O₄, pseudoxanthine C₄H₁N₃O, and two feeble bases C₁₁H₂₄N₁₀O₅ and C₁₂H₂₅N₁₁O₅, both crystallising in tables.

Xanthocreatinine C, H, N, O. Thin sulphuryellow tables, v. e. sol. water, sol. boiling alcohol. Gives an odour of roast meat when heated. Resembles creatinine.

Chrysocreatinine C, H, N,O. Orange crystals, with feebly alkaline reaction. Sl. sol. water. Resembles creatinine.

Amphicreatinine C, H10 N, O4. Yellow prisms, sl. sol. water. A weak base, resembling crea-

Pseudoxanthine C4H5N3O. Yellow crystalline powder, sl. sol. cold water, sol. HClAq and NaOHAq. Reacts with HNO, and KOH like xanthine.

Carnine v. vol. i. p. 710.

PTYALIN v. vol. ii. p. 545.
PTYCHOTIS OIL. The essential oil got from the seeds of Ptychotis ajowan, an umbelliferous plant growing in Central India, contains thymol and a terpene C₁₀H₁₈ (172°) S.G. ¹² ·854 (Stenhouse, C. J. 9, 234; cf. Haines, C. J. 8, 289).

PULEGIUM OIL. The essential oil of Pule-

gium micranthum, growing on the Steppes of Southern Russia, contains oily C10H16O (227°) S.G. 11.932 (Butlerow, J. 1854, 594). It yields acetic and valeric acid when fused with potash. The oil of Mentha pulegium is described under

PENNYROYAL.

PULVIC ACID C18H12OA i.e.

CO.H.CPh: $C < \frac{C(OH):CPh}{CO}$. [215°]. Prepared

by the action of cold NaOHAq on a solution of the anhydride in acetone, and also by boiling vulpic acid with milk of lime (Spiegel, B. 13, 1630, 2219; 14, 1686; 15, 1550; A. 219, 1). Orange crystals or yellow plates (containing EtOH), sol. alcohol and ether, sl. sol. water. Decomposed by baryta-water into phenyl-acetic and oxalic acids. Alkaline KMnO, oxidises it to phenyl-glyoxylic and oxalic acids. On reduction by zinc-dust and NH, Aq it gives a mixture of earboxy-cornicularic acid (C18H14O5), cornicularic acid (C₁,H₁₆O₃), and the two dihydrides of cornicularic acid (C₁,H₁₈O₃).

Salts.—BaA" raq: golden plates.—AgHA": small prisms.—Ag₂A" aq: long needles.

Methyl ether v. Vulpic acid.

Ethyl ether Eth'. [128°]. Prepared by dissolving the anhydride in alcoholic potash.

Methyl derivative of the methyl ether $C_{18}H_{10}Me_2O_5$. [139°]. Colourless needles. An hydride $C_{18}H_{14}O_4$ i.e.

CO.O.C:CPh CO. [221°]. Formed, together with MeOH, by heating vulpic acid at 200°. Colourless needles, sl. sol. alcohol, insol. water. Converted by ammonia into C₁₈H₁₁(NH₂)O₄

[220°] crystallising in yellow prisms. PUNICIN. The purple of the ancients. Extracted from Purpura capillus and other shellfish which yield a colourless secretion which becomes purple on exposure to sunlight (Schunck, C. J. 35, 589; 37, 613). Obtained in Central America from Purpura patula. Yarn dyed by the Indians is warmed with HCl and then with ether. The colouring matter is then extracted with aniline, which on cooling deposits punicin as a crystalline powder, insol. boiling alcohol and ether, sl. sol. benzene and HOAc. Its solution in aniline shows an absorption band between C and D, with a sharp edge towards C. The band resembles that of indigo, but it disappears sooner on standing. In phenol it forms a sky-blue solution. Its solution in conc. H2SO4 shows an absorption band between D and E; this disappears on standing, the liquid becoming bright green. Water precipitates the purple from this solution, so that a sulphonic acid is not formed. Punicin begins to sublime at 190°. It is hardly attacked by HNO₂ (S.G. 1·2) or by aqueous CrO₂. With bromine it reacts, forming a body which crystallises from alcohol in yellow needles. Punicin dissolves in aqueous SnCl₂, but is deposited again on exposing the solution to air.

PURPLE OF CASSIUS. A purple-coloured solid, obtained by adding SnCl₂Aq to AuCl₃Aq in presence of a reducing agent; v. Tin, oxides or, in this volume.

PURPURIC ACID C.H.N.O. Not known in the free state.

Salts. - (NH4)HA" aq. Murexide. Formed by heating ammonium dialurate; by oxidising uranil with HgO; and by adding ammonia or ammonium carbonate to a mixture of

alloxan and alloxantin (Prout, A. Ch. 11, 48; Kodweiss, P. 19, 12; Liebig a. Wohler, A. 26, 319; Fritzsche, J. pr. 16, 380; 17, 47; Beilstein, A. 107, 176; Laurent, C. R. 35, 629; Gregory, A. 33, 334). Garnet-red four-sided prisms with golden-green lustre; when dried in vacuo it forms a brown powder. M. sol. hot water, forming a purple solution, insol. alcohol and ether. Decomposed by acids with formation of uranil and alloxan. The absorption spectrum has been studied by Hartley (C. J. 51, 199).-KHA". Crystals resembling murexide, forming in cold KOHAq a blue solution decolourised by heat.— NaHA".--BaA" 3aq; dark green powder. powder.---AgHA" 1 aq: brownish-red powder. Ag2A".

Isopurpuric acid v. TRI-NITRO-PHENOL, Re-

action 5.

Metapurpuric acid v. (a)-DI-NITRO-PHENOL, Reaction 2

PURPUREO-CHROMIUM SALTS v. CHROM-AMMONIUM SALTS, vol. ii. 159-60.

PURPUREO-COBALT SALTS v. COBALT-

AMINES, vol. ii. pp. 227-8.

PURPUREO-RHODIUM SALTS v. Rно-DIUM-AMMONIUM COMPOUNDS.

PURPURIN v. TRI-OXY-ANTHRAQUINONE.

PURPUROGALLIN C₂₀H₁₆O₉. quinone. [256°]. Formed by the slow oxidation of pyrogallol (Struve, A. 163, 162; Girard, Z. 1870, 86; Wichelhaus, B. 5, 848; De Clermont a. Chautard, C. R. 94, 1362; 102, 1072; Nietzki a. Steinmann, B. 20, 1277; Loew, J. pr. [2] 15, 322). Formed also by the action of aqueous KNO₂ on gallic acid (Hooker, B. 20, 3259). Obtained also from pyrogallol and quinone (Wichelhaus, B. 5, 847; Nietzki, B. 18, 1278). Prepared by adding a solution of K₃FeCy₆ (87 g.) in water (330 c.c.) to pyrogallol (20 g.) dissolved in cold water (330 c.c.), and filtering after half an hour; the yield is small (3 g.). Dark brown needles (from alcohol). May be sublimed. V. sl. sol. water, sl. sol. alcohol, m. sol. ether. Reduces AgNO, and Fehling's solutions. H₂SO₄ forms a crimson solution changed to violet by HNO₂. Ammonia forms a blue solution, changing through green to dark yellow. Yields naphthalene on distillation with zinc-dust. According to De Clermont and Chautard purpurogallin yields Na, A'v crystallising in deliquescent needles, Ba2A1 as crystalline plates, $C_{20}H_{12}Br_{4}O_{6}$ as red needles [204°] and $C_{20}H_{12}Ac_{4}O_{6}$ [186°]. By heating with $H_{2}SO_{4}$ these chemists obtained SO_{2} and $C_{20}H_{12}O_{16}$, which crystallised in brown needles forming a blue solution in KOHAG

PURPUROXANTHIN v. m-DI-OXY-ANTHRA-QUINONE

PURREIC ACID v. EUXANTHIC ACID.

PUTREFACTION v. FERMENTATION.

PUTREFACTIVE ALKALOIDS v. PTOMAÏNES. PUTRESCINE v. Tetra-methylene-diamine. PYKNOMETER. This name is sometimes

given to various forms of the specific gravity bottle, for determining the relative densities of liquids (cf. Densities, relative, vol. ii. p. 373).

PYRANIL-PYROIC ACID C10H10NO(CO2H). [165°]. Formed by heating phenyl-amido-pyrotartaric acid CO2H.CH2.CMe(NHPh).CO2H a little above its melting-point for some time (Reissert a. Tiemann, B. 19, 622; 21, 1942, 3257; 22, 2281; 24, 814). Crystalline powder, sl. sol. PYRENE. 349

water, alcohol, and ether. Converted by heating | strongly into the phenyl-imides of citraconic acid and of phenyl-amido-pyrotartaric acid. According to Anschütz (B. 21, 3252; 22, 731; 23, 895, 2979; A. 246, 115; 248, 269), pyranilpyroïc acid is the mono-anilide of mesaconic acid.

PYRAZINE. This name is used to denote the ring N<CH:CH>N.

Pyrazine hexahydride C₄H₁₀N₂ $\label{eq:nh} {\tt NH} < \stackrel{\rm CH_2.CH_2}{\tt CH_2.CH_2} {\tt NH}. \ \ {\it Piperazine}.$ Di-ethylenediamine. [104°-107°]. (137° uncor.). Formed from ammonia and ethylene bromide (Cloez, Instit. 1843, 213) or ethylene chloride (Natanson, A. 92, 48; 98, 291; Hofmann, Pr. 10, 224; 11, 278; B. 23, 3297). Formed also by boiling NO.C₆H₄.N CH₂·CH₂ N.C₆H₄.NO with alcoholic potash (Bischler, B. 24, 717). According to Majert a. Schmidt (B. 24, 241), it is not identical with spermine. Glittering tables (from water). May be sublimed. Strongly alkaline and absorbs CO, from the air.

Reactions .- 1. Aqueous NaOCl, saturated with Cl, forms ClN CH₂·CH₂·NCl [71°], crystallising in prisms, v. sl. sol. water, v. sol. alcohol (Schmidt a. Wichmann, B. 24, 3243). This body explodes at 80°-85°. Bromine water forms the corresponding di-bromo-piperazine, which is very unstable .- 2. Diazobenzene chloride and NaOHAq form Ph.N₂.N:C₁H₈:N.N₂Ph [129°] (S. a. W.).—3. Benzoic aldehyde yields the compound (N:C,H,:N):CHPh [247] (Schmidt a. Wichmann, B. 24, 3242).—4. Quinone forms an amorphous violet-brown body, almost insol. alcohol.—5. Hydroquinone in alcoholic solution forms C₄H₁₀N₂C₆H₆O₂, crystallising in needles [195°].—6. Phenol forms C₄H₁₀N₂2PhOH [101°], crystallising from spirit in prisms, v. sol. water.-7. Oxalic ether at 110° reacts forming the ether CO2Et.CO.N:C4H4:N.CO.CO2Et [124°] crystallising in broad needles (S. a. W.).—8. p-Chloro-nitro-benzene at 150° gives the compound NO₂.C₆H₄.N:C₁H₅:N.C₆H₄.NO₂ [248°] (Schmidt a. Wichmann, B. 24, 3240).

Salts.—B"H2Cl2 aq: needles, v. sol. water.-B"H_PtCl₆.—B"2HAuCl₄.—B"H_HgCl₄: stellate groups of needles (Sieber, B. 23, 326).—B"C₆H₂(NO₂)₈OH: yellow needles.—Carbonate: [162°-165°].—Urate: B"C₈H_N,O₃. The phosphate and the bismutho-iodide are

crystalline.

Di-acetylderivative C₄H₆N₂Ac₂.

[138·5°]. (above 310°). Formed from piperazine and Ac₂O. Needles, v. e. sol. water and alcohol. Nitrosamine NO.N:C₄H₈:N.NO. [158°] (Ladenburg, B. 24, 2640; Schmidt, B. 24, 3245; G. P. 12, 59222). Reduced by zinc-dust and HOAc to NH₂.N.C.H₂.N.NH₂, which crystallises in needles [c. 100°] (228°), and yields B'HCl, B'C₆H₂N₃O₇, an amorphous di-benzoyl derivative, and reacts with benzoic aldehyde, forming CHPh:N.N:C4H8:N.N:CHPh [205°], which crystallises in pearly plates.

PYRAZOLE C,H,N, i.e. CH:N NH. [70°]. 187°). Formed by heating the tri-carboxylic acid produced by saponification of the product slowly adding H₂SO₄ to aqueous KNO₂ under ap-

of direct addition of diazoacetic ether and acetylene dicarboxylic ether (Buchner, B. 22, Formed also by heating hydrazine hydrate (10.8 g.) with epichlorhydrin (10 g.) and ZnCl₂ (11 g.) (Balbiano, B. 23, 1103). Needles, v. sol. cold water, forming a neutral solution, sol. alcohol and ether. Gives a white pp. with HgCl₂, and also with ammoniacal AgNO₈.

Salts.—B'HCl. Hygroscopic prisms .-B'2H2PtCl6 2aq. At 210° it is split up into HCl and $(C_3H_3N_2)$ PtCl₂, a light-yellow powder.— AgC₃H₁N₂: sl. sol. water.—B'C₆H₃N₃O₇. [160°]. PYRAZOLE BLUE C₂₀H₁₆N₄O₂ *i.e.*

N.NPh.ÇO ÇO.NPh.N CMe. Formed by oxidation CMe - C = C of di-oxy-di-phenyl-di-methyl-dipyrazyl (Knorr, A. 238, 172; B. 25, 765). Sol. chloroform, conc. H₂SO₄ and HOAc. Its spectrum resembles that of bodies of the indigo group. Decomposes about 230°. Its sulphonic acid, got by the action of Br on di-oxy-di-phenyl-di-methyl-dipyrazyl disulphonic acid, is destroyed by excess of bromine (Möllenhoff, B. 25, 1949).

o-PYRAZYL-BENZOIC ACID C10H, N2O, i.e. CH:N čH:CH N.C₆H₄.CO₂H. [139° cor.]. Formed by oxidising o-tolyl-pyrazole with KMnO, (Balbiano, G. 19, 123). Yellow needles (from dilute

alcohol).-BaA'2.

Ethyl ether EtA'. (309° i.V.). p-Pyrazyl-benzoic acid. [265°]. Got in like manner from p-tolyl-pyrazole.—NaA'.—BaA'. Ethyl ether EtA'. [62°]. Tables.

PYRENE C16H10 i.e. CH:CH CH.C C:CH Mol. CH:CH

202. [149°]. (above 360°). S. (alcohol) 1.37 at 16°; 3.08 at 78°: S. (talvano) at 16°; 3.08 at 78°; S. (toluene) 16.5 at 18° (Bechi, B. 12, 1978). Occurs in coal tar (Graebe, A. 158, 285). Monoclinic tables (Fittig a. Hintz, B. 10, 2143), v. e. sol. CS, and ether. Yields pyrene-quinone and pyrenic acid on oxidation. SbCl, at 360° forms CCl, C15Cl10 [above 300°], C₁₄Cl₁₀ [above 300°], and other bodies (Merz a. Weith, B. 16, 2880). Bromine forms crystalline C₁₆H_aBr₄ and C₁₅H₃Br₅ (Graebe).

Picric acid compound C18H10C8H2N2O7. [222°]. Red needles (from alcohol), v. sl. sol. cold alcohol, m. sol. ether, v. sol. benzene.

Hexahydride C₁₆H₁₆. [127°]. Formed by heating pyrene with HIAq and P at 200°. Needles, v. e. sol. ether and boiling alcohol. alcoholic solution is not ppd. by picric acid.

Chloropyrenes. The following chloro-pyrenes are formed by passing Cl into a solution of pyrene in chloroform. They are separated by fractional crystallisation from alcohol, chloroform, and xylene (Goldschmiedt a. Wegscheider. M. 4, 237)

C₁₈H₂Cl. [119°]. Golden needles, forming in H₂SO₄ a solution with violet fluorescence. Yields C₁₆H₃Cl₂. Two di-chloro-pyrenes [156°] and [196°].

C₁₆H,Cl. [257°]. White needles.

350 PYRENE.

ethereal solution of pyrene (Goldschmiedt, M. 2, 580). Yellow needles (from alcohol), sl. sol. cold alcohol.

Di-nitro-pyrene $C_{1e}H_s(NO_2)_2$. Formed from pyrene and HNO_s (S.G. 145). Yellow needles (from HOAc). Reduced by tin and HCl to diamido-pyrene (Jahoda, M. 8, 449). Tetra-nitro-pyrene $C_{16}H_6(NO_2)_4$.

above Yellow needles (from HOAc), nearly 300°].

insol. alcohol.

Amido-pyrene C₁₆H₉.NH₂. [116°]. Got by reducing nitropyrene. Needles, sl. sol. water.—B'₂H₂SO₄ (dried at 100°).—B'HCl. Its solution dyes fir-wood red.

PYRENE CARBOXYLIC ACID C16H9.CO2H. [267°]. Formed by fusing its nitrile with potash (Goldschmiedt a. Wegscheider, M. 4, 256). Needles, sl. sol. water. Resolved by heat into pyrene and CO.—CaA'₂ aq.—BaA'₂2½aq.

Nitrile C₁₈H₉.CN. [150°]. Formed, together with the nitrile of the dicarboxylic acid,

by heating potassium pyrene disulphonate with KCy or K.FeCy. Needles (from alcohol). The solutions of this body show green fluorescence.— (C₁₇H₉N)₂C₈H₃N₃O₇. [134°]. Red needles, decomposed by alcohol.

Pyrene dicarboxylic acid C15H8(CO2H)2.

[above 300°].

Nitrile [above 300°]. $C_{16}H_6(CN)_2$. Formed as above. Yellow powder. Its solutions show green fluorescence.

PYRENE-KETONE C13H8O i.e.

[1:1'] $C_{10}H_6 < CH > CO$. [142°]. Formed by

distilling pyrenic acid with slaked lime (Bamberger a. Philip, B. 19, 1996, 3040; 20, 371; A. 240, 178). Golden tables (from dilute alcohol), volatile with steam. Reduced by zinc and HCl to C₁₂H₈(CH.OH). Oxidised by KMnO, to naphthalene (1,1')-dicarboxylic acid.

PYRENE-QUINONE C16H8O2 i.e.

(Bamberger a. Philip, B. 20, 369; A. 240, 166). [c. 282°]. Formed by oxidation of pyrene by chromic acid mixture (Graebe, A. 158, 295; Goldschmiedt, M. 4, 309). Red needles (from HOAc), sol. aqueous NaHSO,, v. sl. sol. alcohol and ether. Yields pyrene on distillation with and ether. Bromine forms C16H6Br2O2 and zinc-dust. C₁₆H₂Br₂O₂. Its alcoholic solution is turned claret-colour by a little NaOH. Reduced by zinc-dust and ammonia to hydropyrenequinone or pyrene-hydroquinone C₁₆H₁₆O₂, which forms golden crystals exhibiting a dark-blue fluorescence in alcoholic solution, and yielding $C_{16}H_8Ac_2O_2$ [167°].

PYRENE SULPHONIC ACID C16H9.SO3H. Got by fusing the disulphonic acid with potash (Goldschmiedt a. Wegscheider, M. 4, 242).-KA'aq: minute needles (from dilute alcohol).

PYRENE DISULPHONIC ACID C₁₆H₃(SO₃H)₂. Formed by heating pyrene with H₂SO₄. Sticky mass, v. sol. water, sl. sol. alcohol, insol. ether.—K₂A" 2½aq.—BaA" 3½aq.— CaA"2aq: yellow powder.

Formed by the oxidation of pyrene or pyrenequinone by chromic acid (Bamberger a. Philip, B. 19, 1427, 1995, 3036; 20, 369; A. 240, 168). Pale-yellow plates, blackened above 250°. V. sl. sol. alcohol. H.SO, forms an orange solution. Oxidised by KMnO, to naphthalene tetra-carboxylic acid. Reacts with hydroxylamine. Phenyl-hydrazine forms C₁₅H₈O₄(N₂HPh) 2aq, crystallising in yellow prisms. — BaA"aq.—Ag₂A": amber-yellow pp.

 $C_{13}H_6O < CO > 0.$ Anhydride

needles, formed by boiling the acid with HOAc. $Imide\ {\rm C_{1s}H_{e}O}{<}^{\rm CO}_{\rm CO}{>}{\rm NH}.$ Yellow plates,

got by dissolving the acid in NH,Aq.

PYRENOLINE C₁₉H₁₁N. [153°]. Formed by heating amido-pyrene hydrochloride with nitrobenzene, glycerin, and H.SO. (Jahoda, M. 8, 442). Golden scales (from alcohol), v. sol. water, forming solutions which exhibit green fluorrorming solutions which exhibit green hiddrescence.—B'HCl. [270°].—B'₂H₂PtCl₈. [above 290°].—B'H₂SQ₄aq. [246°].—B'C₆H₃N₃Q₇. Decomposes at 260°.—B'MeI. [212°].

PYRIDANTHRILIC ACID C₁₅H₁₆N₂Q₇ i.e. [2:1]CO₂H.C₆H₄.NH.CO.C₃H₂N(CO₂H)₂. [265°].

Formed by the action of alkaline KMnO₄ on explotivation of alkaline kMnO₄ on

cyclothraustic acid derived from a-diquinoline (Weidel a. Strache, M. 7, 289; 8, 197). Micaceous plates, sl. sol. hot water. Yields isocinchomeronic and anthranilic acids on oxidation.

PYRIDAZINE. The ring CH CH.CH CH (Ach, A. 253, 46; cf. Knorr, A. 236, 295).

PYRIDINE C_5H_5N i.e. $N \leqslant \stackrel{CH:CH}{CH:CH} > CH$. Mol. w. 79. (116°) (Schiff, B. 19, 566); (114°) (Ladenburg, A. 247, 4); (116·5°) (Perkin, C. J. 55, 701). S.G. 2 1·0033 (L.); 15·9855; 25·9778 (P.). M.M. 8·761. H.F.p. -19,370 (Thomsen); 7,117 (Ramsay, C. J. 35, 696). H.F.v. -20,530 (Thomsen, Th.). S.H. 418 (Colson, A. Ch. [6] 19, 408). S.V. 89·4 (Schiff); 91·6 (Ramsay). Occurs in bone oil, in coal-tar, and in target by distilling hitmingues shale and pret got by distilling bituminous shale and peat (Anderson, Tr. E. 16, 4; 20 [2] 247; P. M. [4] 2, 257; A. 80, 55; 96, 200; 105, 335; Greville Williams, P. M. [4] 8, 24; Church a. Owen, C. N. 2, 146; P. M. [4] 20, 110; Schulze, B. 20, 409). Occurs also in tobacco smoke, and, in small quantity, in fusel oil (Haitinger, M. 3, 688).

Formation.—1. By heating isoamyl nitrate with P2O5 (Chapman a. Smith, A. Suppl. 6, 329). 2. By distilling its carboxylic acids with lime. 3. By distilling oxytrialdine and oxytetraldine with soda-lime (Schiff, A. Suppl. 6, 21).—4. With other bases by heating glycerin with ammonium phosphate (Stöhr, J. pr. [2] 45, 23).—5. By passing ethyl-allyl-amine over PbO at 450° (Koenigs, B. 12, 2344).—6. By heating piperidine with H_2SO_4 at 300° (K.), with nitro-benzene ($4\frac{1}{2}$ pts.) at 260° (Lellmann a. Geller, B. 21, 1921), or with AgOAc and 10 p.c. HOAc at 180° (Tafel, B. 25, 1621).—7. Together with mono- and disubstitution products by the action of Br on

acetyl-piperidine (Hofmann, B. 16, 587).—8. By heating sodium-pyrrole with NaOMe and CH₂I₂ in sealed tubes at 200° (Dennstedt a. Zimmermann, B. 18, 3316).—9. From anhydroegonine, by treatment with conc. HCl and distillation of the product with zinc-dust (Einhorn, B. 22, 1365).

Purification.—By ppg. the crude base (20 g.) dissolved in (100 g. of) a 10 p.c. solution of HCl by adding HgCl₂ (135 g.) dissolved in hot water (1,000 g.). The crystalline double salt is distilled with NaOHAq (Ladenburg, A. 247, 4).

Properties.—Colourless liquid with powerful odour, miscible with water. Fumes with HCl. Strong poison. Should not be inhaled (Marcus a. Oechsner, Bl. [2] 38, 97; cf. Kendrick a. Dewar, Pr. 22, 432; 23, 290). Forms a hydrate B'3aq, (93°), S.G. 1·0219, V.D. 1·17, which is a colourless liquid (Goldschmidt a. Constam, B. 16, 2976). Not attacked by boiling HNO₃ or CrO₃. Pyridine ppts. the hydroxides of Fe, Al, and Cr from their salts. With CuSO₄ it gives a pale-blue pp. dissolving in excess, forming a deep-blue liquid. Pyridine passes unchanged into the urine (O. de Coninck, Compt. rend. Soc. Bool. 4, 755). Pyridine does not mix with liquid CO₂ (Ditte, C. R. 105, 612). It gives no reaction with hot alkaline NaOBr (Denigès, C. R. 107,

Reactions.—1. Dry chlorine forms di-chloropyridine [72°]. Chlorine passed into a solution of pyridine in CHCl₃ forms B'Cl₂, a white powder (Bally, B. 21, 1772). Chlorine-water ppts. a white powder. KOCl completely decomposes pyridine, forming N, CO₂, CHCl₃, and chloro-acetic acid (Keiser, Am. 8, 308).—2. Bromine added to a solution of the hydrochloride ppts. C₅H₅NBr₂, which is split up by heat into pyridine and bromine. Bromine acting in the cold on pyridine and water gives rise to (C_bH_sNBr₂)_zHBr [126°] (Grimaux, Bl. [2] 38, 127). Pyridine hydrochloride heated with Br at 200° forms mono- and di-bromo-pyridine.-HIAq at 300° yields n-pentane and NH, (Hofmann, B. 16, 590).—4. Sodium at 80° yields (γ)-dipyridyl C₁₀H₂N₂, dipyridine C₁₀H₁₀N₂, isonicotine C₁₀H₁₁N₂, and two nitrogenous oils (240°–260°) and (300°–310°) (Weidel a. Russo, M. 3, 884).—5. Its alkylo-iodides are converted to distillation with soils KOM into relations. by distillation with solid KOH into volatile bases .- 6. Excess of AcCl forms dehydracetic acid (Dennstedt a. Zimmermann, B. 19, 75).-7. Unites with CH,Cl.OH (the product of the union of formic aldehyde and HCl), and the product in aqueous solution gives with HgCl₂ a pp. of (C₂H₂N)(CH₂ClOH)HgCl₂ [162°], with platinic chloride a pp. of (C₃H₃N)₂(CH₂ClOH)₂PtCl₄ [216°], and with pieric acid the compound (C₃H₄N)CH₂(OH).C₄H₂(NO₂)₃O [c. 200°]. Treationally with ANO forms (CH)NCH (CH)NCH ment with AgNO, forms (C,H,N)CH2(OH)NO, crystallising from alcohol-ether in prisms (Hemmelmayr, M. 12, 533). -8. Alloxan and aqueous SO₂ form triclinic crystals of B'C₄H₂N₂O₄H₂SO₅ (Pellizzari, A. 248, 150).—9. Glycollic chlorhydrin forms the base C₇H₂NO (Coppola, G. 15, 331). Glycerin chlorhydrin reacts with formation of C₅H₅NCl.C₂H₅(OH)₂ [107°] separating from alcohol in hexagonal crystals and yielding (C₈H₁₈NO₂Cl)₂PtCl₄ [180°] and C₈H₁₂NO₂ClAuCl₅ [122°] (Krüger, J. pr. [2] 44, 130).—10. Chloroacetic acid at 100° forms C₈H₅NCl.CH₂CO₂H

[202°] which may be viewed as the hydrochloride $C_3H_5N < CH_2 > CO$ of 'pyridine-betaine' (von Gerichten, B.15, 1253; Krüger, B. 23, 2609; J. pr. [2] 43, 279, 297; 44, 136). The pyridinebetaine hydrochloride is also got by oxidising C,H,NCl.CH,CH(OH).CH.OH with chromic acid and is decomposed at 240° into CO2, MeCl, and pyridine. Pyridine betaine C,H,NO2, got by the action of moist Ag₂O on the hydrochloride, forms hygroscopic tables (containing and and yields the salts B'₂H₂PtCl₈ [211°], B'HCl [202°], B'₂HCl aq [159°], B'₂HHgCl₃ aq [134°], B'₂HBr aq, an auro-chloride B'HAuCl₄ [165°] (Jahns, B. 20, 2841), B'HBr, [200°], B'HNO₃, [145°], B'CrO₃ [c. 166°], B'C₄H₃N₃O₇ [143°], B'AgNO₈ [171·5°], and B'HBl 20q On reduction by soliton pages and support the same statements. B'HBiI, 2aq. On reduction by sodium-amalgam, pyridine-betaine yields C,H,NCl CH2.CH2OH. -11. Chloro-acetic ether yields the compound C₅H. NCl.CH₂.CO₂Et, a crystalline powder [100°] (Krüger, J. pr. [2] 43, 274). This ether, C₈H₁₀NO₂Cl, forms the salts B'₂PtCl, [213°], B'AuCl, [117°], B'HgCl₂ [123°], B'CdCl₂ [141°], and B'₂CdCl₂ [107°].

Salts. - B'HCl. Deliquescent mass. -B'2H2PtCl8. [242°]. Orange triclinic prisms. The aqueous solution is converted by long boiling into crystalline B', H2Cl2PtCl, and B'2PtCl, a yellow insoluble powder.—B'₂PtCl₂. Yellow tables, almost insol. cold water (Jorgensen, J. pr. [2] 33, 504). An isomeride crystallises in needles. 55, 044). An isomeriae crystatuses in necures, B',PtCl₂, aq. —B',PtCl₂, —B'₂(NH₃),PtCl₂, —B'(NH₃),PtCl₂, —B'(NH₃),PtCl₂, —B'(NH₃),PtCl₂, B'₂(NH₃),PtCl₂, aq. Minute colourless prisms. —B',HHg,Cl₃, [178°]. Needles (from water).—B',2HgCl₂.—B'HAuCl₄, [285°]. Prisms, sl. sol. water.—B'C,H₃N₃O₃, [162°] Needles, sl. sol. water.—B'2ZnCl2: short prisms, very soluble in cold water (Lang, B. 21, 1578).—B'2ZnCl2 2aq (Lachovitch, M. 9, 516).- B'_2ZnBr_2 : prisms, v. sol. pyridine.— $B'_2H_2ZnCl_4$.— B'_2CuCl_2 [180°-190°].— $B'CuSO_4$ 3aq.— — B'₂CuCl₂ [180°-190°]. — B'CuSO₄ Saq. — B'₄Cu₂Cl₂. — B'₆CuBr₂. — B'₄Cu₂Cl₂. — B'₄Cu₂Cl₂. — B'₄Cu₂Cl₂. — B'₄Cu₂Cl₂. = B'₄CuCl₄. — B'₂CdCl₂: needles, almost insol. alcohol, sol. water. — B'₆CdBr₂. — B'₂CdI₂. — B'CdI₂ (Monari, J. 1884, 629). — B'HgCl₂. — B'₂HgCy₂. — B'₂CaCl₂: white powder. — B'₄NiBr₂. — B'AgBr: white needles, sol. cold pyridine. — B'AgI. — B'AgCy: prisms (Varet, Bl. 1815, 843). — B'CuCy: vellow lamelles. [3] 5, 843).—B'₂CuCy: yellow lamellæ, v. sol. warm pyridine (Varet, C. R. 112, 391). Loses pyridine on warming.—B'H2SO4: deliquescent mass.—B'HI: tables, v. sol. water, but not deliquescent. — B'HI, [89°]. Green crystalline powder (Dafert, M. 4, 508). — B'HBr. Delipowder (Dafert, M. 4, 508). — B'HBr. Deliquescent needles. — B'₂HBr. [126°]. Red tables.—B'₂Cl. Waxy mass (Keiser, Am. 8, 312). — B'HNO₃.—B'₂AgNO₃. [87°] (Jörgensen, J. pr. [2] 33, 502).—B'₃AgNO₃.—B'₄CuSO₄.—B'CuSO₄ 3aq. — B'₄CuS₂O₄.—B'₂SiCl₄. Amorphous (Harden, C. J. 51, 47).—B'₂SiF₄ (Corney a. Smith, Am. 10, 294).—B'₃2SiF₄.—B'₂H₄FeCy₅ 2aq: monoclinio prisms (Mohler, B. 21, 1015).—B'ICl. [132°]. Needles (Pictet a. Krafft, Bl. [3] 7, 72).—B'IClHCl. [180°]. Long yellow needles (from water).—B'.RhCl.HCl. 2aq (Jörgensen, J. pr. [2] water).—B',RhCl₂HCl 2aq (Jörgensen, J. pr. [2] 27,478).—B',RhCl₂.—B',Rh₂PtCl₁₀.—B',Rh₂Cl₂Br. —B',RhCl₂(NO₂) (dried at 100°).—B',Rh₂Cl₄SO₄ (dried at 100°

Acetate B'23HOAc. (140°). Miscible with

852 PYRIDINE.

water, forming an acid solution (Gardner, B. 23, 1588).—Formate B'3CH₂O₂. (149°). Liquid, v. sol. water.—Propionate B'3C₃H₅O₂. (149°). Liquid. — Cupric oxalate compound B'2CuC2O4: minute prisms (Seubert a. Rauter,

B. 25, 2825).

B'MeCl. Methylo-chloride needles (Ostermeyer, B. 18, 591). Converted by hot aqueous solution of picric acid into B'MeOC₆H₂(NO₂)₈ ½aq, which crystallises in greenish-yellow explosive needles [34°]. - $B'_2Me_2PtCl_8$. [188°] (O.); [207°] (B.). -(B.). [188°] (O.); [207°] [253°].—B'MeClICl. [8 B'MeÂuCl. [253°].—B'MeClICl. [82°] (U.); [90°] (B.). Yellow plates.—B'MeClICl. [180°]. [82°] (O.); Unstable yellow crystals (Bally, B. 21, 1774).

Methylo-perbromide B'MeBr. [48°].

Methylo-iodide B'MeI. Converted at 290° into methyl-pyridine hydriodides. On heating with alcoholic potash at 45° it yields a brown resin, forming a deep-red solution in alcohol, turned bright-red by HCl, orange-red by acetic acid, and ruby-red by ammonia (O. de Coninck, C. R. 102, 1479). Chlorine forms B'MeICl, [90°] (v. supra). Oxidised by alkaline K₂FeCy, to oxy-methyl-pyridine or v-methyl-pyridone $CH \stackrel{CH.CO}{< CH:CH} > NMe$ (250°), a liquid miscible with water (Decker, J. pr. [2] 47, 28; cf. Pechmann, B. 24, 3144).

Silvery Ethylo-iodide B'EtI. (Anderson, A. 94, 364). At 300° it yields pyridine, NH_3 , (a) and (γ)- ethyl-pyridine, and diethyl-pyridine (Ladenburg, B. 16, 2059; 18, Oxidised by alkaline 2961). Yields B'2Et2PtCl6.

K₃FeCy₆ to CH CH:CH NEt (250°).

Ethyleno-iodide B'C2H4I2. Prisms (Coppola, G. 15, 332). Ag₂O yields a base C,H, NO. By heating pyridine with ethylene bromide and some alcohol at 100° there is found $\mathbf{B'_2C_2H_4Br_..}, \text{which yields } \mathbf{B'_2C_2H_4PtCl_6}$ (Davidson, A. 121, 254).

Benzylo-chloride B'PhCH2Cl. Reduced by sodium-amalgam to the unstable C21H21N2 (Hofmann, B. 14, 1503).—(B'PhCH₂Cl)₂PtCl₄.

Nitro-benzylo-chlorides $B'C_0H_4(NO_2).CH_2CI.$ o [c. 76°], m [70°-100°], p [90°-100°]. These bodies are reduced by tin and HClAq to B'C,H4(NH2Cl).CH2Cl, which are split up by heat into pyridine hydrochloride and C.H. NH₂Cl (Lellmann a. Pekrun, A. 259, 54).

Phenacylo-bromide B'BzCH2Br. Prisms (Bamberger, B. 20, 3344).—B'₂(BzCH₂)₂Cr₂O₇.
References.—Bromo-, Chloro-, Oxy-Amido-,

and Oxy- PYRIDINE. Dipyridine C₁₀H₁₀N₂ v. DIPYRIDYL DIHYDRIDE.

Dipyridine $C_{10}H_{10}N_{2}$. (275°). S.G. ¹³ 1·124. Is probably a dipyridyl dihydride. Formed by heating nicotine with KOH and K₂FeCy₆, and also by heating the product of the action of S on nicotine at 150° with finely-divided copper (Cahours a. Etard, Bl. [2] 34, 452). Inactive liquid.—B'HHgCl,—B'H-PtCl₆ 2aq.— B'₂H₂FeCy₂2aq: brownish-green tables.
(a)-PYRIDINE CARBOXYLIC ACID

C_eH_bNO₂ i.e. CH:CH.C.CO₂H CH:CH.N Picolinic acid. [136°]. Formed by oxidation of (a)-methylpyridine with KMnO₄ (Weidel, B. 12, 1994), and by oxidation of (a)-phenyl-pyridine (Skraup, M. 4, 477). Obtained also, together with its hexahydride, from comenamic acid by successive treatment with PCl₅ and H₂SO₄, the resulting di-chloro-picolinic acid being reduced by heating for three days at 155° with HI dissolved in

HOAc (Ost, J. pr. [2] 27, 285).

Preparation.—The three acids got by oxidation of crude methyl-pyridine from animal oil are converted into copper salts. Cupric pyridine (a)-carboxylate is extracted by hot water. The residue is treated with H.S., and the difficultlysoluble (γ) - acid separated from the (β) - acid (Ost, J. pr. [2] 27, 286).

Properties.—Needles, v. sol. water and alcohol, almost insol. ether, benzene, CHCl3, and CS₂. May be sublimed. FeSO, gives a red colouration with picolinic acid and with all the carboxylic acids of pyridine that contain CO.H in the (a)-position (Skraup, M. 7, 210). The absorption of the ultra-violet spectrum has been

studied by Hartley (C. J. 41, 45).

Reactions.—1. Yields pyridine on distillation with lime or with alcoholic potash at 240°. The Cu salt on distillation gives pyridine and (a)dipyridyl [70°] (Blau, M. 10, 375; B. 21, 1077). -2. Sodium-amalgam forms δ-oxy-adipic acid (Weidel, M. 11, 522).-3. Furning HI at 170° forms (a)-methyl-pyridine and piperidine (Seyfferth, J. pr. [2] 34, 241).-4. Zinc-dust and HOAc reduce it to (a)-methyl-pyridine.

Constitution.—This may be deduced from its formation from (β) -naphthoquinoline vid (β) phenyl-pyridine carboxylic acid (Skraup a. Co-

benzl, M. 4, 436).

Salts. — HA'HCl: unstable crystals. -H₂A'₂H₂PtCl₆2aq: orange-red crystals.—NH₄A': triclinic tables.—KA'.—BaA'₂½aq.—CaA'₂aq.—

MgA'22aq.

Hexahydride C.H. N.CO.H. Piperidine (a) carboxylic acid. The chief product of the action of HI at 160° on mono- or di-chloro-picolinic acid (Ost, J. pr. [2] 27, 287). Got also by reducing picolinic acid (Ladenburg, B. 24, 640). Syrup, v. sol. water. Salts.—B'HCl. B'2H2PtCl₂ 2aq.—B'MeCl. [191°]. Needles.

Pyridine (R) carbornia.

Pyridine (\$)-carboxylic acid ÇH:CH.C.CO₂H Nicotinic acid. [230°]. CH: N.CH

Formation.—1. By oxidising nicotine with HNO, (Weidel, A. 165, 330), CrO, (Huber, A. 141, 271; B. 3, 849), or KMnO, (Laiblin, A. 196, 129).—2. A product of oxidation of coal-tar bases (Weidel; Mohler, B. 21, 1009).—3. By heating quinolinic acid at 160° or with HCl at 180° (Ost, J. pr. [2] 27, 286; Lippmann a. Fleissner, M. 8, 315).—4. By oxidation of (β) methyl-pyridine (Weidel, B. 12, 2004), (\$\beta\$)-ethyl-pyridine (Stoehr, J. pr. [2] 43, 155) or (\$\beta\$)-phenyl-pyridine (Skraup, M. 4, 453).—5. By saponification of its nitrile, which is got by distilling sodium-pyridine sulphonate with KCy (Fischer, B. 15, 63). -6. By heating three of the pyridine dicarboxylic acids (Hoogewerff a. Van Dorp, R. T. C. 1, 1, 107; A. 204, 117; 207, 226; Weidel a. Herzig, M. 1, 16).—7. By heating berberonic acid at 215° (Fürth, M. 2, 420).—8. By the action of Zn and HClAq on chloro-nico-

tinic acid (Pechmann a. Welsh, C. J. 47, 145).

Properties.—Needles, sl. sol. cold water, sol. alcohol, nearly insol. ether. May be sublimed.

Reactions.—1. Yields pyridine on distillation with lime. - 2. Sodium-amalgam yields a mixture of δ-oxy-α-methyl-glutaric acid and its lactone, which, on treatment with alcohol and HCl, yields a mixture of two ethers. One of these, C10H17ClO4, is converted by sodium-amalgam into a-methyl-glutaric acid, and the other, C₈H₁₄O₄, is also converted by successive treatment with PI, and with Zn and dilute H,SO, into a-methyl-glutaric acid (Weidel, M. 11, 502).— 3. Bromine and water at 120° yield CO₂, pyridine, and bromoform .- 4. The K salt heated with MeI at 150° forms C, H, NMeI.CO, Me, which on saponification yields C_bH₄NMe(OH).CO₂H [130°], converted at 100° into trigonellin C,H,NMe<0 [218°], which occurs in the seeds of Trigonella fanum gracum (Jahns, B. 18, 2521; Hantzsch, This anhydride forms the salts B. 19, 31). C.H., NO, HCl, B', H, PtCl, aq, B'HAuCl, [198°],

and B'43HAuCl [186°].

Constitution.—This may be deduced from its formation from (a)-naphthaquinoline vid (a)-

phenyl-pyridine carboxylic acid.

Salts. — HA'HCl. Colourless prisms.— H_A'_2H_2PtCl_2 2aq.—B'_2H_AuCl_3.—B'HNO_1 aq.— NH_A': needles.—KA'.—MgA'_2: needles.— CaA'₂ 5aq: monoclinic crystals; a:b:c =1.537:1:629; $\beta = 62^{\circ} 50'$. — Cu(OH)A' (De Coninck, Bl. [2] 42, 100).—AgA': needles (from

hot water).

Nitrile C₈H₄N.Cy. [49]. Formed by distilling sodium pyridine sulphonate with KCy (Fischer, B. 15, 63). Needles or prisms, sol. water. — B'HCl. — B',H,PtCl,: yellow soluble needles. Tables (by sublimation). Converted by means of hydroxylamine into the amidoxim $C_bH_1N.C(NH_2):NOH$ [128°], which yields an acetyl derivative [148°] and a benzoyl derivative [190°], and is converted by phenyl cyanate into C₅H₄N.C(NOH).NH.CO.NHPh [167°], by phenyl into C,H,N.O N,S O.NHPh thiocarbimide [241°], and by succinic anhydride at 100° into $C_bH_4N \stackrel{N.O}{\leqslant} C.CH_2.CH_2.CO_2H$ [178°]. acetyl and benzoyl derivatives are converted by heating into $C_bH_4N \stackrel{N.O}{\swarrow} CMe$ [109°] and $C_bH_4N \stackrel{N.O}{\leqslant} OPh$ [189°] respectively (Mi-

chaelis, B. 24, 3489).

Hexahydride C₂H₁₀N(CO₂H). Nipecotinic acid. [250°]. Got by reducing nicotinic scid in alcoholic solution by Na (Ladenburg, B. 25, 2768). Crystals, v. e. sol. water, insol. alcohol and ether.—HA'HCl. [240°].—H,A',H,PtC [213° cor.]. — HA'HAuCl. [197° cor.]. HA'HCl5HgCl. [231°].—MeA'HCl [208°]. [240°].—H₂A'₂H₂PtCl₂. AuCl₄. [197° cor.].— Me₂A',H₂PtCl_a. — Nitrosamine [112°]. O.H.N.O.

Pyridine (γ) -carboxylic acid

N CH.CH CO.CO.H. Isonicotinic acid. [8059]

(S.); [306°] (B. a. H.); [309·5°] (W. a. H.).

Formation.—1. By heating pyridine s-tricarboxylic acid (Skraup, B. 12, 2331) and three

of the pyridine di-carboxylic acids (Hoogewerff a. Van Dorp, A. 204, 112; Weidel a. Herzig, M. 1, 28; Böttinger, B. 14, 68).—2. By the action of $KMnO_4$ on (γ) -methyl-pyridine (Behrmann a. Vor. IV.

Hofmann, B. 17, 2696; Ladenburg, B. 21, 287). 3. By heating di-chloro-pyridine carboxylic acid [210°] with HI (B. a. H.).

Properties.—Needles, sl. sol. cold water, insol. alcohol. Yields pyridine on distilling with lime and δ-oxy-ethyl-succinic acid on reduction

with sodium-amalgam (Weidel, M. 11, 517).
Salts.—NH,A': needles.—CaA', 4aq: silky
needles, m. sol. water.—HA'HCl: monoclinic

prisms.—H_A'_H_PtCl₀ 2sq: monoclinic crystals. Hexahydride C₅H₁₀N(CO₂H). Got by reducing the acid in alcoholic solution by Na (Ladenburg, B. 25, 2778). Branching groups of needles, v. e. sol. water, insol. alcohol. Blackens at 300°, but is not melted at 320°.—HA'HCl. [228°]. Trimetric crystals; a:b:c = 922:1: 979. [228]. Trimetric crystals; a:b:c = 922:1:979.-H.A'₂H₂PtCl₈. [239°]. — Aurochloride: [197°]. Nitrosamine C₂H₁₀N₂O₃. [101°].

Pyridine (aa)-dicarboxylic acid

CH.C(CO₂H) N. Dipicolinic acid. [226°]
CH:C(CO₂H) N. Dipicolinic acid. [226°] (L. a. R.; S.); [237°] (E.); [236° cor.] (Collie, C. J. 59, 179). Formed by oxidation of (aa)-dimethyl-pyridine [145°] (derived from acetoacetic ether or from coal-tar) by KMnO, (Epstein, A. 231, 26; Ladenburg a. Roth, B. 18, 52; 19, 790; 20, 130; A. 247, 32; Lange a. Rosenberg, B. 20, 132; cf. Dewar, C. N. 23, 18). Got also by oxidation of (a)-methyl-(a)-ethyl-pyridine by

dilute (2 p.c.) KMnO, (Schultz, B. 20, 2724).

Properties.—Hair-like needles (containing $1\frac{1}{2}$ aq) or anhydrous scales; sl. sol. cold alcohol, water, and ether. On heating at 245° in a current of H it yields pyridine and pyridine (a)carboxylic acid. PCl, forms a chloride [610],

(284°). FeSO, gives a reddish-yellow colour. Salts. — CaA" 2aq: minute prisms. — CuA" 2aq: dark-blue prisms.

Pyridine (aB)-dicarboxylic acid

C(CO,H).C(CO,H) N. Quinolinic acid. [281°]. •55 at 6.5°. Formed by the oxidising action of KMnO, on quinoline (Hoogewerff a. van Dorp, B. 12, 747; R. T. C. 1, 107; A. 204, 117), on cinchonine (H. a. D.), on c- and p- methylquinoline (Skraup, M. 2, 157), on c-oxy-quinoline, on quinoline o-sulphonic acid (O. Fischer a. Renouf, B. 17, 755), and on (a)-oxy-quinoline carboxylic acid (La Coste a. Valeur, B. 20, 103).

Properties. — Monoclinic needles, a:b:c= $^{\circ}542:1:^{\circ}607$; $\beta = 64°54'$; sl. sol. water and alcohol, insol. ether. Begins to decompose at 140°, and forms nicotinic acid. Yields pyridine when distilled with lime. Reduced by sodium-amalgam to the δ-lactone of butane aβy-tricarboxylic acid CH₂CH₂CH₂CO₂H (Perlmutter, M. 13, 840).

Salts.-KHA" 2aq: triclinic plates (Lippmann a. Fleissner, M. 8, 811).-K2A" 2aq. BaA" aq.—Ag₂A": orystalline.—AgHA" aq.

Anhydride C,H,N<00>0. [135°]. Got from the acid and Ac.O (Bernthsen a. Mettegang, B. 20, 1208). Prisms. Converted by benzene and AlCl, into C.H.BzN.CO.H [147° converted by heat into the ketone C.H.BzN (807° uncor.), which yields a crystalline phenylhydrazide [143.5°]. Pyridine $(\alpha \gamma)$ -dicarboxylic acid

CO,H.O CH: C(CO,H) N. Lutidinio acid. [240°] (V.); [235°] (B.; L. a. R.). Formed by the oxidising action of KMnO₄ on $(\alpha\gamma)$ -dimethyl-pyridine (Ramsay, P. M. [5] 4, 241; 6, 19; Weidel a. Herzig, M. 1, 20; Ladenburg a. Roth, B. 18, 915; A. 247, 37), on (a)-methyl-(γ) ethyl-pyridine (Schultz, B. 20, 2726), on diethyl-pyridine, on (aa)-di-methyl-dipyridyl (Heuser a. Stochr, J. pr. [2] 44, 409), and on the methyl-pyridine carboxylic acid obtained from uvitonic acid (Böttinger, B. 14, 68; 17, 93; Voigt, A. 228, 54).

Properties.—Needles (containing aq), m. sol. cold water, sol. alcohol, insol. ether. Gives a blood-red colour with FeSO₄. Yields pyridine on distillation with lime. Converted by heat into CO2 and isonicotinic acid. PCla yields a

chloride [203°].

Salts.—KHA" ½aq. Crystals.—(NH₄)₂A".—
(NH₄)HA" aq: hygroscopic needles.—CaA" 3aq.
CaA" aq.—CaA" ½aq.—CaH₂A", 2aq.—BaA" aq.
—BaA" 1½aq (B.).—BaA" 3aq.—CdA" 4aq (Waage, M. 4, 727).—MgA" 5aq.—CuA" 3aq. CuA" 4aq (B.).—Ag₂A" 2aq: white pp.

Pyridine $(a\beta')$ -dicarboxylie acid $CH \stackrel{CH:C(CO_2H)}{\sim} N$. Isocinchomeronic acid.

[236°].

Formation.—1. By oxidation of lutidine (150°-170°) with KMnO, (Ramsay, P. M. [5] 4, 246; Weidel a. Herzig, M. 1, 1; Lange a. Rosenberg, B. 20, 135).—2. By oxidising (\$\beta\$) methyl-(a)-ethyl-pyridine (Ladenburg, A. 247, 44).—3. By oxidation of quinine (Ramsay a. Dobbie, C.J. 33, 102; B. 11, 324).—4. By heating the dihydride of potassium pyridine tricarboxylate (Weiss, B. 19, 1311).—5. By the action of alkaline KMnO, on cyclothraustic acid and on pyridanthrilic acid (Weidel a. Strache, M. 7, 290). - 6. By oxidising (β) -ethyl- (α) -stilbazole (Plath, B. 22, 1062).

Properties.—Small prisms (containing aq), almost insol. cold water, alcohol, and benzene, sol. hot HClAq. On heating with HOAc at 220° it is split up into CO₂ and nicotinic acid [230°]. On heating with lime it gives pyridine. PCl, yields a chloride [61°] converted by NH₃ into an amide [297°]. FeSO₄ gives a reddish

Salts.—(NH₄)HA" aq. sms, sl. sol. cold [253°]. Triclinic prisms, sl. sol. cold water.—(NH₄)₂A".

KHA" laq: needles.—K₂A" aq.—CaA" 2aq.Ca(HA")₂ 3aq.—MgA" 5aq.—CuA" aq.—Ag₂A".

Methyl ether MgA". [117.5°] (R.).

Pyridine $(\beta\beta')$ -dicarboxylic acid

CH ⟨C(CO,H).CH⟩N. Dinicotinic acid. [322°]. Formed by heating pyridine (2,3,5)-tri-carboxylic acid and pyridine (2,3,5,6)-tetra-carboxylic acid (Riedel, B. 16, 1613; Hantzsch a. Weiss, B. 19, 286; Weber, A. 241, 12). Formed also by heating di-chloro-pyridine di-carboxylic acid with conc. HIAq at 180° (Guthzeit, A. 262, 130)

Properties.—Small prisms (from HOAo), split up by heat into CO₂ and nicotinic acid.

Salts.—PbA" 2aq.—Ag2A" aq.—Ag2A" 12aq. - HA'HCl 2aq: needles, decomposed by water.

HA',H.PtCl; : orange-red needles.
Pyridine (βγ)-dicarboxylic acid
CO,H.C CH CH CH CH Cinchomeronic aqid. [259°].

Formation.—1. By oxidation of einchonine or einchonidine by HNO, of S.G. 1.4 (Weidel, A. 173, 76).—2. By oxidation of quinine by HNO, the yield being 28 p.c. (Weidel a. Schmidt, B. 12, 1146).—3. By heating apophyllenic acid with conc. HClAq at 240° (Von Gerichten, B. 13, 1635).—4. By heating pyridine tricarboxylic acid (formed from cinchonic acid) at 190° (Hoogewerff a. van Dorp, B. 13, 61; Skraup, M. 1, 184; Weidel a. Brix, M. 3, 604).—5. By the action of KMnO4 on methyl-pyridine carboxylic acid and on isoquinoline (Hoogewerff a. van Dorp, R. T. C. 2, 23; 4, 285).—6. By heating pyridine pentacarboxylic acid (Weber, A. 241, 16) .- 7. By oxidation of methyl-nicotinic acid, derived from (\$\beta\$)-collidine (Oechsner de Coninck, Bl. [2] 43, 106).—8. By boiling berberonic acid with HOAc (2 pts.) and Ac₂O (1 pt.) for six hours (Mayer, M. 13, 344; cf. Fürth, M. 2, 426).

Properties.—Prisms (from HClAq), v. sl. sol. water and ether, sl. sol. alcohol. nicotinic and some nicotinic acid on heating (Hoogewerff a. van Dorp, A. 207, 217). Gives pyridine on distillation with lime. Sodiumamalgam yields NH, and cinchonic acid C,H,O,, i.e. $CO_2H.CH < CH_2.CO_2$ (Weidel a. Hoff, M.

CO₂H

13, 578). Cinchonic acid [169°] forms monoclinic crystals, v. sol. hot water and alcohol, yields $BaC_7H_6O_6$ Saq, $Ba_3(C_7H_7O_7)_2$ Saq, CaA'' 2aq, $Ca_3(C_2H_2O_2)_2$ (dried at 190), and oily Et₂A", whence PCl₃, followed by alcohol, yields CO.Et.CH(CH.Cl).OH(CO.Et).CH..CO.Et, which is a heavy oil. Cinchonic acid is reduced by HI to butane tricarboxylic acid C, H10O6 [184°], whence Ca, A'', 8aq, accompanied by an isomeric butane tricarboxylic acid [133°]. FeSO, gives no colour.

Salts.—Na₂A" 2aq: tables.—NaHA".—BaA" 1½aq: needles, sl. sol. water.—CaA" 3½aq: prisms. — CaA"3aq. — CuA''33aq: small blue crystals.—Ag₂A": white pp.—AgHA".—HA'HCl: monoclinic prisms, decomposed by

water.—H₂A'₂H₂PtCl_e: golden prisms.

Anhydride (C_bH₃N),C₂O₃. [77°]. Formed by boiling the acid with Ac₂O (Goldschmiedt a. Strache, M. 10, 156). Plates. May be sublimed. Converted by NH, gas into C,H,N(CO,NH,).CONH, [229°], which is converted by heat into a yellow powder [130°], and which yields the amic acid C₈H₂N(CO₂H).CONH₂ [237°], orystallising in needles.

Mono-ethyl ether HEtA". [133°]. Formed from the anhydride and EtOH. Plates (from benzene).—AgEtA": long needles.

Mono-methyl ether HMeA". [154°]. Anhydride of the Methylo-hydroxide $C_8H_7NO_4$ i.e. $C_5H_8NMe(CO_2H) < O$. Apophyllenic acid [242°]. Formed by oxidation of cotarnine by HNO₃ (Wohler, A. 50, 24; Anderson, Tr. E. 23, 347; C. J. 5, 257; Gerichten, B. 18, 1685). Formed also by heating cinchomeronic acid with MeI and MeOH at 100° (Roser, A. 234, 116). Needles (anhydrous) or octahedra (containing aq), sol. hot water, insol. alcohol and ether. HClAq at 250° decomposes it, forming cinchomeronic acid and McCl. — BaA'₂. — AgA'.—Ag₂A'(NO₂).—H₂A'₂H₂PtCl₄ aq.

Bromo-apophyllenic acid C₈H₆BrNO, 2aq. [205°]. Formed by oxidation of bromo-tarconine. Yields BaA'₂3aq and $H_2A'_2H_2PtCl_8$ (Gerichten, A. 210, 91).

Pyridine $(\alpha \alpha' \beta)$ -tricarboxylic acid

CH CCO,H).C(CO,H) N. Formed by oxidising di-methyl-nicotinic acid with KMnO, (Weiss, B. 19, 1309). Crystallises from alcohol in plates (containing 2aq), v. e. sol. water. Decomposes at 160° into CO₂ and isocinchomeronic acid. FeSO, colours its neutral solution red. — RH_RA''', 5aq: needles.—Ca₂A''', 4aq.—Pb₂A''', 5aq. Pyridine (αα'γ)-tricarboxylic acid

 $CO_2H.C \stackrel{CH.C(CO_2H)}{<} N.$ Trimesitic acid. Carbolutidinic acid. [244°]. Formed by oxidation of uvitonic acid (Böttinger, B. 13, 2048; 14, 69), and by oxidation of the tri-methylpyridine obtained from acetoacetic ether and aldehyde-ammonia (Voigt, A. 228, 31). Tables or spheroidal groups of needles (containing 2aq), sol. hot water, sl. sol. alcohol and ether. Yields isonicotinic acid on sublimation. FeSO, gives a violet-red colour.

Salts.— $K_3A'''_5aq$: needles.— $Ca_3A'''_24aq$.— $Ba_3A'''_26aq$.— $BaH_4A'''_24aq$.— $Mg_3A'''_212aq$.— $Cu_3A'''_212aq$.— $Ag_3A'''1_2aq$.

Ethyl ether Et_3A''' . [127.5°].

Amide. [above 280°]. Pyridine $(\alpha\beta\beta')$ -tri-carboxylic acid

CH ⟨C(CO,H):C(CO,H)⟩N. Carbodinicotinic acid. [323°]. Formed by oxidation of quinoline (Py. 2)-carboxylic acid (Riedel, B. 16, 1615), methyl-pyridine (\(\beta\beta'\)-dicarboxylic (Weber, A. 241, 11), and of the parvoline got by heating propionic aldehyde-ammonia with propionic aldehyde at 200° (Dürkopf, B. 21, 832, 2707; 23, 689). Spherical aggregates (containing $1\frac{1}{2}$ aq), v. sol. hot water. At 155° it is split up into CO_2 and dinicotinic acid.—Ba₃A'''₂5aq.—Ag₂HA''' $1\frac{1}{2}$ aq: rosettes of plates.

Pyridine $(a\beta\gamma)$ -tri-carboxylic acid At 155° it is

 $CO_2H.C \leqslant C(CO_2H):C(CO_2H) \geqslant N.$ [250°] (H. ...

D.; S.); [257°] (D. a. R.). S. 1.2 at 15°

Formation .- 1. By the action of KMnO, on quinine, cinchonine, cinchonidine, quinoïdine, and cinchonic acid (Dobbie a. Ramsay, C. J. 35, 189; Hoogewerff a. van Dorp, B. 12, 158; 13, 152; A. 204, 84; Skraup, A. 201, 312; Strache, M. 10, 642).—2. By oxidation of cinchonine by HNO₃ (Weidel, A. 173, 101; B. 12, 415).—3. By oxidation of methyl-pyridine (αβ)-dicarboxylic acid (Hoogewerff a. van Dorp, R. T. C. 2, 18), of di-methyl-pyridine carboxylic acid (Michael, B. 18, 2027), of (α)-oxy-cinchonic acid (Weidel a. Cobenzl, M. 1, 865).—4. By oxidation of papaverine (Goldschmiedt, M. 6, 397). Trimetric plates (containing 1½aq), v. sol. hot water, m. sol. alcohol, nearly insol. ether. Blackens at 200°. FeSO gives a reddish colour. H₂S forms a red amorphous body (D. a. R.). Decomposed by long heating at 180°, or by boiling with HOAc into CO₂ and cinchomeronic acid. Yields MeI and pyridine on distilling with lime. MeOH at 100° form CO, and apophyllenic acid. Sodium-amalgam gives NH, and cinchonic acid. PCl, forms a chloride (206° at 40 mm.). Salts. — K,A" 3aq. — Ba,A", 16aq.

 $Ba_2A'''_212aq.$ — $Ca_2A'''_214aq.$ — $Ca_2A'''_213aq.$ — $Ca_3A'''_22aq.$ — $Cu_3A'''_29aq:$ light-blue pp.— CuHA'''3 2aq: hexagonal prisms.—CuH,A''', 2aq. —Cd₂A" 26aq.—Ag₂HA" aq.—Ag₃A" 2aq; amorphous pp.—AgH₄A" 2½ aq.—H₄A"'HCl: crystalline powder (Roser, A. 284, 125).

Pyridine (aB'7)-tricarboxylic acid
CO₂H.C CH.C(CO₂H)-N. Berberon Berberonic acid. [243°]. Formed by oxidising berberine with nitric acid (Weidel, B. 12, 410; Fürth, M. 2, 416). Triclinic prisms (containing 2aq), sol. hot water, v. sl. sol. hot alcohol, insol. ether. Yields pyridine on distillation with lime. FeSO, gives a red colour. At 215° it is split up into CO2 and nicotinic acid; above 243° it yields isonicotinic acid.

Salts. — K₃A"'4½aq. — K₂HA"'3aq. — KH₂A"'1½aq. — Ca₃A"'₂8aq. — Cd₃A"'₂4aq. — Ag, A": white pp., insol. water.

Pyridine $(\beta\beta'\gamma)$ -tricarboxylic acid $CO_2H.C < \stackrel{C(CO_2H).CH}{\subset (CO_2H):CH} N.$ (β) - Carbocinchomeronic acid. [261°]. Formed by heating dipotassium pyridine pentacarboxylate at 220° (Weber, A. 241, 17). Plates (containing 3aq), v. sol. hot water. Yields cinchomeronic acid when heated. Gives no colour with FeSO4. Cu,H,A''', 24aq.—Ag,A''' 2aq: crystalline pp. Pyridine $(\alpha\beta\beta'\gamma)$ -tetra-carboxylic acid

 $C_{\mathfrak{s}}H_{\mathfrak{s}}NO_{\mathfrak{s}}$ i.e. $CO_{\mathfrak{s}}H.C \leqslant \stackrel{C(CO_{\mathfrak{s}}H).C(CO_{\mathfrak{s}}H)}{C(CO_{\mathfrak{s}}H)} \stackrel{N.}{=} OH$ Formed by oxidising $(\alpha \gamma)$ -di-methyl-pyridine $(\beta \beta')$ -dicarboxylic acid (Weber, A. 241, 23). Prisms (containing 2aq or 3aq). At 120° it loses CO2, forming (β)-carbocinchomeronic acid. FeSO, gives a dark-red colour.—Ba2A1v 4aq.— Ag, HAiv, aq: crystalline.

Pyridine $(\alpha\alpha'\beta\beta')$ -tetracarboxylic acid

ing di-methyl-pyridine dicarboxylic acid (Hantzsch, B. 19, 286; Weber, A. 241, 4). Needles (containing 2aq), v. sol. water. Decomposes at 150° into CO, and dinicotinic acid.-CaH₂A^{iv} 2aq: needles, v. sol. water.—Cu₂A^{iv} 5aq.

 $\begin{array}{lll} -\text{Ag}_{A}\text{A}^{\text{iv}} & \text{2aq: bulky pp.} \\ \text{Pyridine } & (a\alpha'\beta'\gamma)\text{-tetra-carboxylic acid} \\ \text{CO}_{2}\text{H.C} < & \overset{\text{CH}}{\subset} & \overset{\text{C}(\text{CO}_{2}\text{H})}{\subset} & \text{CO}_{2}\text{H} \\ \end{array} \right) \text{N.} \quad \text{Formed} \end{array}$ Formed by oxidation of tri-methyl-pyridine carboxylic acid and of di-methyl-pyridine di-carboxylic acid Michael, A. 225, 142). Formed also by oxidation of flavenol by alkaline KMnO, (Fischer a. Täuber, B. 17, 2927). Slender needles (containing 2aq), v. sol. water, v. sl. sol. alcohol and ether. Not decomposed at 150°. FeSO, gives a brownish-red colour. - Ba, Aiv 2 aq. - Ba, Aiv aq. -Cu₂A" 2½aq.—Ag₄Aiv aq.

Pyridine penta-carboxylic acid NC,(CO2H)8. Formed from potassium tri-methyl-pyridine dicarboxylate and KMnO, (Hantzsch, A. 215, 62; Weber, A. 241, 15). Crystalline mass of minute needles (containing 2aq), extremely sol. water, v. sl. sol. ether. Acid to litmus and to taste. Loses 2aq at 120° and decomposes, without having melted, at 220°. The neutral alkaline salts are very soluble, but the acid alkaline salts are sl. sol. water. The acid does not combine with HCl. Distilled with lime it gives pyridine. FeSQ4 gives a dark-red colour.

Salts.—KH,A' 2 or 8aq. When heated swells up like Pharaoh's serpent.—K2H3AV 312aq. K₃Av. — Ba₃Av₂ 11aq. — Ca₃Av₂ 12aq. CaH, A v 2 aq. - Mg, A v 2 12 aq. - Ca 2 (NH 4) A v 5 aq. -Ag,HA 2aq.

Double salt with oxalic acid

KH4AVKHC2O4 5aq.

References.—Bromo- and Oxy-pyridine carb-OXYLIC ACID.

PYRIDINE TETRAHYDRIDE C.H.N i.e. CH₂ CH : CH NH. Piperidein. Formed by heating amido-valeric aldehyde with solid KOH (Wolffenstein, B. 25, 2782).—B'HAuCl₄. [141°]. —B₂'H₂PtCl₆.—B'HCl. [230°].—B'HBr. [178°].

Pyridine hexahydride v. Piperidine.

PYRIDINE (B)-SULPHONIC ACID C₈H₄N.SO₂H. Formed by heating pyridine (1 pt.) with H₂SO₄ (3 pts.) at 320° (0. Fischer, B. 15, 62; 16, 1183). Small needles or plates, sol. water, sl. sol. alcohol, insol. ether. Br added to its boiling aqueous solution forms dibromopyridine. Potash-fusion gives oxy-pyridine [123°]. On distillation with KCy it yields the nitrile of nicotinic acid. The K salt, heated with

MeI at 150° forms crystalline $C_5H_4NMe < ^{>}_0$

(Hantzsch, B. 19, 36).—BaA', 4aq: needles. C,H,NSO, Pyridine sulphonic acid? [155°]. Formed from pyridine and ClSO₂H (Wagner, B. 19, 1157). Crystalline, decomposed by water into pyridine and H2SO4.

Pyridine disulphonic acid C,H,N(SO,H)2. Formed by heating piperidine (1 pt.) with H₂SO₄ (10 pts.) (Königs, B. 16, 735; 17, 592). Needles (from HOAc), v. sol. water, nearly insol. alcohol and ether. PCl_s at 200° forms tri-chloropyridine [48°]. — Na₂A" 4aq. — K₂A" 3aq. — PbA" 4 aq.

PYRIDONE v. OXY-PYRIDINE.

(aa)-DIPYRIDYL $C_{10}H_{8}N_{2}$ i.e. $C_{5}H_{4}N.C_{5}H_{4}N.$ [70°]. (272.5°). V.D. 5.6. Formed by distilling cupric picolinate (Blau, B. 21, 1077; M. 10, 875). Crystals (from water), m. sol. water, v. sol. alcohol. Not hygroscopic. Strong base. FeSO, colours its aqueous solution red. Oxidised by KMnO₄ to picolinic acid.—B"H₂PtCl₆.— B"H₂FeCy₆.—B"C₆H₂N₅O₇. [155·5°]. Needles. Hexahydride. Formed by reducing the

 $\begin{array}{ll} \textbf{base with zinc and HClAq.} & \textbf{Alkaline oil.} \\ \textbf{Dodecahydride} & \textbf{C}_{10}\textbf{H}_{20}\textbf{N}_{2}. & \textbf{Dipiperidyl.} \end{array}$ (259° cor.). Formed by reducing the base, dissolved in isoamyl alcohol, by Na. Powerful base. Very deliquescent. Not poisonous. Forms with CS₂ a compound [93°], and yields a nitrosamine [159°].—B"H,PtCl, 21 aq.

(ββ)-Dipyridyl C₁H₂N₂. [68°]. (287°) (L. a. O.); (292° at 736 mm.) (S. a. V.). Formed by distilling its dicarboxylic acid with KOH (Skraup a. Vortmann, M. 4, 591), and by the dry distillation of pyridine disulphonic acid (Leone a. Oliveri, G. 15, 276). Extremely deliquescent needles, miscible with water and alcohol, sl. sol. ether. Yields nicotinic acid on oxidation.—B",H,PtCl,.—B"2C,H,N,O,. [232°].

Hexahydride C₁₀H₁₄N₂. Nicotidine.

(288°). Got by warming the base with tin and conc. HClAq. Poisonous oil, v. e. sol. water and alcohol, m. sol. ether.—B",H,PtOl,: orange-red pp.—Picrate: [202°].

(77)-Dipyridyl $C_{10}H_8N_2$. [114°]. (305° cor.). V.D. 5'9 (calc. 5'5). Formed by boiling pyridine with sodium (Anderson, A. 154, 274; Weidel a. Russo, M. 3, 854). Formed also by heating its (aa)-dicarboxylic acid with HOAc at 180° (Heuser a. Stochr, J. pr. [2] 44, 407). Tables, sl. sol. cold water, v. sol. alcohol. Crystallises from water in tables (containing 2aq). [73°]. Tastes bitter. Yields isonicotinic acid on oxidation. Br forms $C_{10}H_{\phi}Br_2N_2$ crystallising from alcohol in needles. MeI and EtI form crystalline B"2MeI and B"2EtI respectively.

Salts. — B"H₂Cl₂. Monoclinic crystals; a:b:c=1.064:1:.595. $\beta=112^{\circ}$ 33'.—B"H₂ZnCl₄. - B"H₂HgCl₄. Monoclinic tables; β=91° 3'. — B"H,PtCl_s. -= '673:1: '341: B"2HNO_s. [256°]. Trimetric prisms; a:b:c

= 841:1: 397. - B"HNO₃AgNO₃. -B"H₂SO₄2aq. Dihydride C₁₀H₁₀N₂. Dipyridine. (290°) at 735 mm. V.D. 5.0 (calc. 5.5). Formed by the action of Na on pyridine (Anderson, C. J. 22, 406; Weidel, M. 3, 879). Liquid, sol. water and alcohol.—B"H2PtCl8. — B"2MeI. — B"Me2PtCl4 (Ramsay, C. J. 36, 264).

Hexahydride $C_{10}H_{14}N_2$. Isonicotine. [78°]. (above 260°). Formed by reducing $(\gamma\gamma)$ -dipyridyl with tin and HClAq (W. a. R.). Deliquescent needles, sol. water, alcohol, and benzene. Has hardly any smell. Strongly alkaline and caustic. Poisonous, acting somewhat like curari. Its salts are much less poisonous. Oxidised by KMnO, to isonicotinic acid.—B"2HNO₄. Deliquescent needles. — B"H₂PtCl₆ aq. — B"₂H₄Hg₅Cl₁₆. — B"2MeI: triclinic prisms (from MeOH).

Dodecahydride $C_{10}H_{20}N_2$. [122°]. Formed by reducing $(\gamma\gamma)$ -dipyridyl in alcohol by Na (Ahrens, B. 21, 2929). Needles, insol. water, v. sol. alcohol and ether.—B"H2PtCl8.—B"HAuCl4. Picrate: needles, blackening when heated.

(281°). Formed by Dipyridyl C₁₀H₈N₂. passing pyridine vapour through a red-hot tube (Roth, B. 19, 360). Oil.—B"2HCl: hygroscopic needles.—B"H₂PtCl₉.—Picrate: [208°].

 $(\alpha\beta)$ -Dipyridyl $C_{10}H_8N_2$. (296° cor.). Formed by heating its carboxylic acid with lime (Skraup a. Vortmann, M. 3, 599; Blau, B. 24, 326). Oil, sol. alcohol and ether. — B"H₂PtCl₂ aq. — B"C₂H₃N₃O₇. [149·5°]. Yellow needles. Dodecahydride C₁₀H₂₀N₂. [69°]. (269°)

cor.). Formed by reducing the base with isoamyl alcohol and Na (Blau, M. 13, 332). Hygroscopic crystalline mass. Not identical with nicotine hexahydride. Strongly alkaline, absorbing CO₂ from the air. V. e. sol. water, but much water gives a turbidity. V. sol. alcohol, m. sol. ether. CS₂ forms a compound [205°].—B"H₂Cl₂. V. e. sol. [238°]. — B"2HAuCl₄. [212°]. — B"2C₆H₂N₂O₇. [215°].

Bensene sulphonyl derivative. [157°].

Nitrosamine C₁₀H₁₈N₂(NO)₂. [88°]. Dipyridyl dodecahydride? C₁₀I C10H20N2? (251°). S.G. 4 ·956. Is perhaps (αβ)-dipyridyl hexa-hydride (Blau). Formed by the action of Na on nicotine in alcoholic solution (Liebrecht, B. 18, 2970; 19, 2590). Lævorotatory liquid, v. sol. water, alcohol, and ether. Alkaline in reaction. Smells like piperidine. Readily unites with CS2. Forms an oily nitrosamine and an oily di-acetyl derivative (c. 405°).—B"2HCl. Sol. alcohol.—B"H₂PtCl_{*}. [202°].—B"H₂I₉: brown needles.—

reducing nicotine obtained a mixture of bases (235°-265°) yielding a sparingly soluble platinochloride [218°], which gave a base (244°), possibly methyl-nicotine hexahydride $C_{11}H_{22}N_2$

(α)-PYRIDYL-ACRYLIC ACID C, H, NO, i.e. C, H, N.CH:CH.CO, H. [203°]. Formed by heating α-oxy-tri-ω-chloro-propyl-pyridine with alcoholic potash (Einhorn, B. 20, 1593; 23, 220; A. 265, 215). Got also by heating α-oxy-pyridylpropionic acid. Small transparent needles, v. sl.

sol. cold water, v. sol. alcohol.

Reactions. — 1. \mathbf{Br} in HOAc forms CaH4N.CHBr.CHBr.CO2H [127°], which melts at 146.5° when containing HOAc of crystallisation.-2. Hydrogen bromide in HOAc forms (C₅H₄N.CHBr.CH₂.CO₂H)HBr [164°], crystallising in needles.—3. MeI yields HA'MeI [220°] converted by AgBr into HA'MeBr [242°].

Salts. HA'HCl. [220°]. H2A'2H2PtCl. [210°]. Red prisms. — HA'HAuCl,. [195°].

HA'HBr. [223°].—CaA'2.—AgA': needles. Methyl ether MeA'.—MeA'HCl. [186°]. MeA'HBr. [242°]. MeA'HI. [220°]. Ethyl ether EtA'. Needles.

(a)-PYRIDYL-BUTYLENE C₀H₁₁N C₅H₄N.CH:CHEt. (148° at 75 mm.). By distilling oxybutyl-pyridine C₅H₄N.CH₂.CH(OH).C₂H₅ with KOH in vacuo or by heating it with HClAq at 165° (Matzdorff, B. 23, 2711). Colourless oil, smelling like conyrin. - Platinochloride:

[140°].—B'HAuCl₄. [130°]. Small needles.
(αβ)-DIPYRIDYL (β)-CARBOXYLIC ACID
C₁₁H_{**}N₂O₂ i.e. C₅H₄N.C₅H₄N.CO₂H. [183°].
Formed by heating the dicarboxylic acid at 200° (Skraup, B. 15, 896; M. 3, 597). Needles (containing 1½aq), sl. sol. cold water and alcohol. Gives a yellow colour with FeCl₂.—CaA'₂ 2aq.- $AgA' \frac{1}{2}aq$: prismatic needles.

 $(\alpha \bar{\beta})$ -Dipyridyl $(\beta \gamma)$ -dicarboxylic acid

[215°]. Formed by oxidation of phenanthroline by KMnO, (Skraup a. Vortmann, B. 15, 896; M. 3, 587). Triclinic prisms (containing 2aq), sl. sol. cold water, v. sol. alcohol. FeSO, colours its aqueous solution red. Salts.-KHA" aq.-CaA" 3aq.—CuA" 3aq.—BaA" 13aq. Crystals.— AgHA" 4aq.—H₂A" 2HCl.—H₂A"H₂PtCl₆ 3aq.— (H₂A")₂H₂PtCl₆ 6aq : golden prisms.

(ββ)-Dipyridyl (αα)-dicarboxylic acid C₁₂H₈N₂O₄. [213°]. Formed by oxidation of pseudo-phenanthroline by KMnO₄ (Skraup a. Vortmann, M. 4, 583). Thick prisms (containing ½aq), sl. sol. cold water, alcohol, and ether. FeSÖ, gives an orange-yellow colour.—K,A "5aq. — KHA"2aq. — CaA"5aq. — CuA" $\frac{3}{2}$ aq. — CaA"5aq. — CuA" $\frac{3}{2}$ aq. — Ag₂A" $\frac{1}{2}$ aq.—Ag₂A"AgNO₂.—H₂A''HCl aq. Monoclinic prisms; a:b:c=1.27:1:2.29; β =110° 16'.— H2A 'H2PtCl, 8aq: orange crystalline pp

 $(\gamma\gamma)$ -Dipyridyl (aa)-dicarboxylic acid. [247.5°]. Formed by oxidising (aa)-di-methyl-dipyridyl by KMnO₄ (Heuser a. Stoehr, J. pr. [2] 44, 405). Needles, v. sl. sol. water and alcohol. gives a reddish-yellow colour. AcOH at 180° gives $(\gamma \gamma)$ -dipyridyl.

Dipyridyl tetracarboxylic acid? [96°]. Got by oxidising diquinolyl (Claus, B. 14, 1942). Needles (from hot water). Pb2Aiv. Ag4Aiv.

PYRIDYLENE-PHENYLENE-KETONE

PHENYLENE PYRIDYL RETONE.

Pyridylene-phenylene-ketone sulphonic acid $C_eH_s(SO_sH) < CO.U:N-CH.CH.$ Formed by oxidising (B)-naphthoquinoline sulphonate with alkaline KMnO₄ (Immerheiser, B. 22, 408). Yellow plates (from water), v. sl. sol. alcohol and ether. Yields an oxim crystallising in yellow flakes and a phenyl-hydrazide crystallising in minute orange needles.—KA' aq.—BaA', 2aq.—PbA', 3aq.—AgA' aq.

PYRIDYL-ETHYLENE C,H,N.CH:CH, (160°). Formed by the action of NaOHAq on the hydrochloride of \$\beta\$-bromo-\$\beta\$-pyridyl-propionic acid (Einhorn, \$B. 23, 221; \$A. 265, 229). Liquid.—B'HAuCl₄. [144°]. Yellow needles.

(a)-PYRIDYL ETHYL KETONE C.H.NO i.e. C₅H₄N.CO.C₂H₅. (205°). Formed by distilling calcium picolinate with calcium propionate (Engler a. Bauer, B. 24, 2530). Oil, sol. alcohol. Sodium-amalgam forms a pinacone [136°] Phenyl-hydrazine sulphonic acid yields a crystalline compound [268°].-B'HgCl2: crystalline.-B'EtI. [160°].

C,H,N.C(NOH).C,H,. OximNecdles. Yields an acetyl derivative [46°] and

a benzoyl derivative [69°].

(β)-Pyridyl ethyl ketone C₂H₄N.CO.C₂H₃ Formed by distilling calcium nicotinate with calcium propionate (Engler, B. 24, 2539). Yields a phenyl hydrazide [145°] and a phenyl-hydrazide sulphonate [235°] which forms B'2H2PtCla. B'C₆H₈N₈O₇, and B'HgCl₂ [130°]. Oxim C₅H₄N.C(NOH).C₂H₅.

(α)-PYRĬDÝL METHÝL KĚTONE

C_sH₄N.CO.CH_s. (192°). Formed by distilling calcium picolinate with calcium acetate (Engler a. Rosumoff, B. 24, 2527). V. sol. alcohol and ether. Readily volatile with steam. Yields an oxim [120°], a phenyl-hydrazide [155°], and a phenyl-hydrazide sulphonate which is not melted at 300°.—B'C,H,N,O,. [131°].—B'HgCl. [150°].—B'MeI. [161°].—B'EtI. [205°].
(β)-Pyridyl methyl ketone C₃H,N.CO.CH,

(220°). Formed by distilling calcium nicotinate with calcium acetate (Engler a. Kiby, B. 22, 597). Oil, v. sol. acids. Yields an oxim [112°], which yields B'HCl [204°]. The phenyl-hydrazide [137°] crystallises from alcohol in yellow needles. -B'HgCl₂. [158°]. White needles.

DI-(a)-PYRIDYL-PROPANE C₁₃H₁₄N₂ i.e. 2 1.0281 S.G. $CH_{2}(CH_{2}.C_{5}H_{4}N)_{2}.$ (323°). Formed by heating picoline with methylal and ZnCl₂ for 10 hours at 290° (Ladenburg, B. 21, 8100). Yellow oil, v. sol. alcohol and ether .-[215°].--B"2HAuCl, 1 aq. Salts: B"H,PtCl,

Balts: B'H₂FCl₁₀. [210].—B 2HAUO, 15aq.

—B"H₂Hq₄Cl₁₀. [161°]. Large plates.

Dodecahydride C₁₃H₂₆N₂. Dipipecolylmethane. [54°]. (195° at 26 mm.). Formed by reducing the base with Na and alcohol. Crystalline mass, sl. sol. water.—B"2HCl: very hygroscopic needles. — B"2MeCl. [171°]. -C; H24Me2N22MeI. Crystals, v. e. sol. water.

C.H.N.CO.Pr. (216°-220°). Formed by distilling calcium picolinate with calcium butyrate (Engler a. Majmon, B. 24, 2536). Oil. Yields an oxim [48°] which forms a benzoyl derivative [57°]. Forms a phenyl-hydrazide [82°] and a

(a)-PYRIDYL PROPYL KETONE

phenyl-hydrazide sulphonic acid [251°]. ketone is reduced, in dilute alcoholic solution, by sodium-amalgam to a pinacone C₁₈H₂₄N₂O₂ [146°]. The chloro-iodide melts at 85°.

Salts.—B'₂H₂PtCl_s.—B'HgCl₂. [c. 78°].-

B'MeI. [79°].

(8)-Pyridyl propyl ketone C,H,N.CO.Pr. (246°-252°). Formed by distilling calcium nicotinate with calcium butyrate (Engler, B. 24, 2541). Yellow needles, sol. alcohol. Yields a phenyl-hydrazide [182°], a phenyl-hydrazide sulphonate [283°], a crystalline oxim, and an ethylo-iodide [192°]. — BHgCl₂. [178°]. — B'C.H.N.O.

(B.3)-(a)-PYRIDYL-QUINOLINE C,4H,0N2 i.e. ÇH.CH.C.CH:CH [104°]. Formed by CH. N .C.CH:C.C,H,N' heating the Ag salt of its carboxylic acid (O. Fischer a. H. van Loo, B. 19, 2475). Prisms.

B'2H2PtCla

Carboxylic acid C15H10N2O2 i.e.

 $C_{\bullet}H_{\bullet}N.C \leqslant_{N-CH}^{CH:CH} > C.CO_{2}H.$ [273°]. Formed by oxidising (β)-diquinolyl with CrO₃. Needles, v. sl. sol. water.—AgA': pale yellow pp.

PYRO.. Use of this prefix applied to inorganic compounds; for pyro-compounds v. the compounds to the names of which pyro- is prefixed. Thus pyro-phosphoric acid will be found under Phosphoric Acid, and pyro-phosphates under Phosphates.

PYROCATECHIN $C_0H_0O_2$ i.e. $C_0H_4(OH)_2[1:2]$. Catechol. o-Di-oxy-benzene. Oxyphenic acid. [104°] (F. a. M.); [111°] (Mortinon). (240°–245°). H.C.p. 685,200. H.C.v. 684,900. H.F. 85,800 (Stohmann, J. pr. [2] 45, 334). Occurs in urine aspecially after administration. in urine, especially after administration of benzene or phenol (Baumann, H. 1, 244; 3, 157; Nencki a. Giacosa, H. 4, 335; Schmiedeberg, H. Occurs in the green leaves of the Virginia creeper (Ampelopsis hederacea) (Gorup-Besanez, B. 4, 905) and in the sap of the plants from which kino is prepared (Flückiger, B. 5, 1). Occurs sometimes in raw beet sugar (Lippmann, B. 20, 3298). Occurs in wood-tar (Béhal a.

Desvignes, Bl. [3] 9, 144).

Formation. - 1. By dry distillation of catechin, moritannic acid, and all varieties of tannin that turn green with FeCl, (Zwenger, A. 37, 327; Wagner, J. pr. 52, 450; 55, 65; Eissfeldt a. Uloth, A. 92, 101; 111, 215).—2. By the dry distillation of wood (Buchner, A. 96, 188).-3. By heating cellulose, starch, or cane sugar with water at 200°-280° (Hoppe-Seyler, B. 4, 15).—4. By potash-fusion from o-iodo-phenol (Körner, Bull. Acad. Belg. [2] 24, 166; Lautemann, A. 120, 815).—5. By the action of HI on guaiacol (Gorup-Besanez, J. 1867, 688; Baeyer, B. 8, 155).-6. By the dry distillation of protocatechnic acid and of quinic acid (Strecker, A. 118, 285; Hasiwetz a. Barth, J. 1864, 405; Tiemann a. Haarmann, B. 7, 617).—7. By potash-fusion from o-phenol sulphonic acid (Kekulé, Z. 1867, 643), benzoic acid, gum guaiacum (Hlasiwetz a. Barth, A. 130, 352; 134, 282), and, together with resorcin, from o- and m-bromo-phenol (Fittig, B. 8, 364).—8. By soda-fusion from phenol (Barth a Schreder, B. 12, 419).—9. From phenol and H₂O₂ (Martinon, Bl. [2] 43, 157).— 10. By passing a rapidly alternating electric surrent through a solution of phenol.—11. A

product of the action of water at 200° on benzene hexachloride (Meunier, C. R. 100, 1591).

Preparation.-1. From HIAq and guaiacol at 200° or by heating guaiacol with conc. HClAq for 4 hours at 175° (Perkin, jun., C. J. 57, 587). By fusing o-phenol sulphonic acid with potash at 350° (Degener, J. pr. [2] 20, 308).
 Properties.—Large plates (from benzene) or

needles (from water), v. sol. water, alcohol, and ether, m. sol. benzene and chloroform, insol. ligroin. Gives an acid reaction in presence of borax (Lambert, C. R. 108, 1017). FeSO, gives no colour. FeCl, colours the aqueous solution green, turned violet-red by alkalis (Ebstein a. Müller, Fr. 15, 465). The alkaline solution absorbs oxygen, becoming brown. It reduces AgNO₃, AuCl₃, and platinic chloride. Ppts. a conc. solution of egg-albumen. Does not ppt. gelatin. Lead acetate gives a white pp. Quinone in ethereal solutions forms C.H.O.C.H.O. crystallising in deep-green needles with violet lustre [153°] (Clermont a. Chautard, C. R. 102, 1072). Ppts. a solution of quinine sulphate, forming $C_{20}H_{24}N_2O_2$ H_2SO_4 $C_6H_6O_2$ aq, which separates from alcohol in yellow crystals [167°], v. sl. sol. cold water.

Reactions.—1. Nitric acid acts violently, forming oxalic acid .- 2. Phthalic anhydride and ZnCl₂ at 150° form 'pyrocatechin phthaleïn'

C.H. < CO.Ö -C(C₈H₃(OH)₂)₂, a yellow mass forming a blue solution in alkalis and yielding a tetrabenzoyl derivative [202°] (Baeyer a. Kochendörfer, B. 22, 2196).—3. Phenyl cyanate at 100° forms C₆H₄(O.CO.NHPh)₂ [165°] crystallising in needles, v. sol. alcohol (Snape, C. J. 47, 772). 4. The disodium compound $C_0H_1(ONa)$ treated with CO_2 in the cold forms $C_0H_4(O.CO_2Na)_2$, which at 100° changes to the compound $C_0H_2(O.CO_2Na)(OH).CO_2Na$ and at 210° to $C_6H_2(OH)_2(CO_2Na)_2$ (Schmitt a. Hähle, J. pr. [2] 44, 2).-5. Ammonium carbonate and water at 140° react forming protocatechuic acid and C_oH_a(OH)(CO₂H)₂[1:2:3].—6. K₂S₂O, acting on K salt forms crystalline C_oH_a(O.SO₃K)₂ and C₈H₄(OH)(O.SO₃K) (Baumann, B. 11, 1913).—7. Cl.CONH₂ forms C₆H₄(O.CONH₂)₂ [178°] crystallising from alcohol in needles.-8. Chlorine passed into its solution in acetic acid forms

CCl CCl₂CO crystallising (with 2aq) from ether-ligroin, and from ligroin (with aq). melting at 94° (Zincke a. Klein, B. 21, 2719).— 9. KOH and ClCO₂Et form C₆H₄CO₃ [118°] (Bender, B. 13, 697), (225°-230°) (M. Wallach, A. 226, 84).

Estimation. — By extracting its acidified aqueous solution with ether, evaporating the ether, dissolving the residue in water, and pre-cipitating with lead acetate. The pp. is dried at 100° and weighed (Degener, J. pr. [2] 20, 303).

Salts.—C₈H₄O₂Pb. White pp.—C₆H₅O₃Sb or C_eH₄ SbOH. Formed by adding SbCl_s to a

solution of pyrocatechin saturated with NaCl (Causse, Bl. [3] 7, 245). Prisms, insol. water, alcohol, and ether, sol. alkalis and mineral acids. Ac₂O at 125° forms C_eH₄(OAc), and Sb(OH)(OAc),. Di-acety l derivative C_eH₄(OAc),

Needles (Nachbauer, A. 107, 243).

Di-bensoyl derivative $C_6H_4(OBz)_2$. [84°]. Plates (Doebner, A. 210, 261; Hinsberg,

A. 254, 254).

Mono-methyl ether C_cH₄(OH)(OMe). Guaiacol. Mol. w. 124. [28·5°]. (20·5°) (Tiemann a. Koppe, B. 14, 20·16; Béhal a. Choay, Bl. [3] 9, 142). S.G. № 1·125 (V.); 2·1·153; ½ 1·143 (B. a. C.). A product of distillation of gum guaiacum (Sobrero, A. 48, 19; Deville a. Pelletier, A. 52, 403; Völckel, A. 89, 349). Occurs among the products of the distillation of wood (Hlasiwetz, A. 106, 362; Gorup-Besanez, A. 143, 151). Prepared by heating calcium vanillate with slaked lime (Tiemann, B. 8, 1123) and by heating pyrowith MeI and KMeSO, (Gorup-Besanez, A. 147, 248, or with NaOMe, MeOH, and MeI). Liquid, with peculiar odour, sol. alcohol and ether, sl. sol. water, sol. dry glycerin and ligroïn. FeCl, gives a green colour in its alcoholic solution. Dissolves in alkalis. Yields C₆H₅.OMe on heating with zinc-dust (Marasse, A. 152,64). PCl, forms C₆H,Cl.OMe (Fischli, B. 11, 1463). I and KOHAq give a coffee-brown pp. [125°-130°] (Messinger a. Vortmann, B. 22, 2320). The K salt is converted by acetochloration of the converted by acet hydrose into $C_6H_4(OMe)(O.C_6H_{11}O_5)$ [157°] (Michael, Am. 6, 339). Phthalic anhydride and SnCl, at 115° form 'guaiacol-phthalein,' which yields a crystalline benzoyl derivative (Baeyer, B. 22, 2199). H₂SO, forms two sulphonic acids (Tikmann a. Koppe, B. 14, 2019).— C_sH₄(OK)(OMe) 2aq.—KHA'₂ aq: prisms (from alcohol), decomposed by water.—Pb(OH)A': flocculent pp. — $C_6H_4(O.SO_3K)(OMe)$: white needles.— $C_6H_4(OAo)(OMe)$. (235°-240°). V.D. 82·7 (obs.). Colourless liquid.

Di-methyl ether C₆H₄(OMe)₂. Veratrole. (205°). V.D. 68·6 (obs.; H=1). S.G. 15 1·086. Formed by heating veratric acid with baryta (Merk, A. 108, 60; Koelle, A. 159, 243; Tiemann, B. 14, 2016). It is obtained also from $C_8H_4(OK)(OMe)$ and MeI (Marasse). Solidified at 15°

Methyl ethyl ether C₆H₄(OMe)(OEt). (213°). V.D. (H=1) 75·6 (obs.). Liquid. Methyl propyl ether C₆H₄(OMe)(OPr). (240°-245°). Liquid (Cahours, Bl. [2] 29, 270). Di-ethyl ether C₆H₄(OEt)₂. [44°]. Formed from pyrocatechin, Etl, and alcoholic potash

(Herzog a. Zeisel, M. 10, 152).

derivative CoH4(OC,H7)2. Di-benzyl[61°]. Yellowish needles (from alcohol). Forms a nitro-derivative crystallising in needles [98°].

The mono-benzyl derivative is liquid, but its nitro-derivative forms yellow needles (from alcohol) [129°] (Schiff a. Pellizzari, A. 221, 378; G. 13, 507).

Sulphonic acid C₆H₃(OH)₂(SO₃H)[4:3:1]. Formed by fusing phenol (a)-disulphonic acid with KOH at 300° (Barth a. Schmidt, B. 12, 1260). Deliquescent needles, v. sol. water and alcohol, insol. ether.—KA'.—NaA'aq.—BaA'2.

References .- AMIDO-, BROMO-, CHLORO-, and NITRO- PYROCATECHIN, and TRI-BROMO-GUAIACOL.

PYROCINCHONIC ACID v. DI-METHYL-

MALBÏO ACID.

PYROCOLL C₁₆H₆N₂O₂ (Magnanini, B. 22, 2502). [269°]. A product of the distillation of gelatin when free from fat but containing albumen, casein, or gluten (Weidel a. Ciamician, M. 1, 279; 2, 29). Formed also, together with

HOAc, by heating the acetyl derivative of pyrrole (a)-carboxylic acid (Ciamician a. Silber, B. 17, 103; G. 14, 162, 563). Colourless plates, insol. water and cold alcohol, sl. sol. ether. Sublimes before fusion.

Reactions .- 1. Boiling KOHAq converts it into pyrrole carboxylic acid.—2. Alcoholic NH, forms the amide of pyrrole carboxylic acid.—3. PCl₃ forms C₁₀Cl₂N₂O₂ [above 320°] insol. ether and C₁₀Cl₁₀N₂O [197°] sol. ether (Ciamician a. Danesi, G. 13, 28). The perchloro-pyrocoll The perchloro-pyrocoll C₁₀Cl₈N₂O₂ is converted by boiling KOHAq into tri-chloro-pyrrole carboxylic acid, and by PCl, into C₁₀Cl₁₄N₂O₂ [144°].—4. Bromine forms C₁₀H₃BrN₂O₂ [192°], C₁₀H₄Br₂N₂O₂ [290°], and C₁₀H₂Br₄N₂O₂, which is converted by KOHAq into di-bromo-pyrrole carboxylic acid (Ciamician, G. 11, 330; 12, 29; B. 16, 2388).

Reference.--CHLORO-, BROMO-, and NITRO-

PYROCOLL.

PYROCRESOL. C₁₅H₁₄O? An inappropriate name given by Schwarz (B. 15, 2201; 16, 2141; M. 3, 726; cf. Armstrong, C. J. Proc. 3, 114) to some neutral substances found in coal-tar.

(a)-Pyrocresol [195°]. Thin silvery plates, yielding crystalline C₂₈H₂₂Br₅O₂? and oxidised by Chlorine acting on a solution of (a)-pyrocresol in chloroform gives C₁₈H₁₁Cl₈O [225°]? HI reduces (a) pyrocresol to a hydrocarbon $C_{1a}H_{22}$? (Bott, C. J. Proc. 3, 114).

(β)-Pyrocresol [c. 124°]. Vields, on oxidation, '(β)-Pyrocresol oxide 'C₁₃H₁₂O₂ [95°].
 (γ)-Pyrocresol [165°]. Vields '(γ)-pyrocresol oxide ' [77°] on oxidation. Bromine forms C₂₂H₁₂Br₂O₂ oxystallising in trimetric plates.

PYROGALLIC ACID v. PYROGALLOL. PYROGALLOL C_eH_eO_s i.e. C_eH_e(OH)₈[1:2:3]. c-Tri-oxy-benzene. Pyrogallic acid. Mol. w. 126. [131°] (Etti, B. 11, 1882; cf. Stenhouse, A. 179, 236); [134°] (Stohmann). (210°). S. 40 at 12°. H.C.p. 633,300 (Berthelot a. Louguinine, A. Ch. [6] 13, 839; C. R. 104, 1577). H.F. (from diamond) 137,700 (B. a. L.); 132,000 (Stohmann, J. pr. [2] 45, 336). Occurs in woodtar as dimethyl ether.

Formation.-1. By heating gallic acid (alone or mixed) with pumice stone (2 pts.) (Braconnot, A. 1, 26; Pelouze, A. 10, 159; Liebig, A. 101, 47).—2. By heating di-iodo-o-oxy-benzoic acid with KOHAq (Lautemann, A. 120, 299).—3. By heating (α) - or (β) - chloro-phenol sulphonic acid with KOH at 190° (Petersen a. Baehr, A. 157, 136) .- 4. By heating gallic acid (10 g.) with glycerin (30 c.c.) at 200° as long as CO, comes off (Thorpe, Ph. [3] 11, 990).—5. By heating gallic acid (1 pt.) with aniline (2 pts.) at 120° (Cazeneuve, Bl. [8] 7, 549). The product is aniline pyrogallate C.H.O.2NPhH, [56°], which gives off aniline when exposed to air, or when shaken with benzene.

Properties.—Prisms, v. sol. water, m. sol. alcohol and ether. Tastes bitter. Poisonous (Personne, Z. [2] 5, 728). Its alkaline solution rapidly absorbs oxygen, turning brown, and giving off a little CO in bulk about 10th of the oxygen absorbed (Calvert a. Cloez, A. 130, 248). Reduces KMnO₄ (Monier, C. R. 46, 577) and salts of mercury, Ag, Au, and Pt. FeSO₄ gives

a white milkiness, but if a ferric salt is present, or the pyrogallol solution has become slightly oxidised by keeping, FeSO₄ gives an indigo-blue colour, changing to brownish-red on standing and on adding HOI (Jacquemin, C. R. 77, 593; Cazeneuve, Bl. [2] 44, 114; C. R. 101, 56). FeCl₂, in absence of air, gives a transient blue colour, restored by cautious addition of alkali. A solution of pyrogallol renders borax slightly acid (Lambert, C. R. 108, 1017). Nitrous acid colours the aqueous solution brown (Schönbein, Fr. 1, 319). K₂CrO, and H₂SO, give a dark colour. Reduces CuSO, and, on addition of alkalis, gives a black colour changed by NH, to red. Cu(OAc)2 gives at once a black colour. Pyrogallol fused with ammonium oxalate yields ammonium rufigallate, which dissolves in water with red colour and gives with K_*FeCy_* and $K_2Cr_2O_*$ a dark-brown pp. insol. alcohol (Kliebahn, Fr. 26, 641). A solution of iodine in presence of Na2SO4 gives a purple colour (Nasse, B. 17, 1186). A solution of HgCl₂ and pyrogallol in alcohol gives a black pp. with alkaloids but not with glucosides (Schlagdenhaufen, Ph. [3] 4, 772). An alkaline solution of pyrogallol absorbs about 50 vols. of NO in 12 hours, but no N₂O (Russell a. Lapraik, C. J. 32, 35). Pyrogallol does not react with hydroxylamine (Baeyer, B. 19, 163). A solution of potassium pyrogallate which is absorbing oxygen in contact with alumina gives out a feeble light, especially in presence of Na,S (Lenard a. Wolf, P. [2] 34, 918).

Reactions.—1. Fuming HNO, forms oxalic acid .- 2. Bromine forms tri-bromo-pyrogallol .-3. Chlorine in presence of HOAc forms mairogallol (vol. iii. p. 165) and crystalline leucogallol C₁₈H₈Cl₁₂O₁₂ 2aq, which yields tri-chloro-pyrogallol when boiled with water and zinc-dust (Stenhouse a. Groves, C. J. 28, 706; Webster, C. J. 45, 208; Hantzsch a. Schniter, B. 20, 2033).-4. Ozone passed through a solution of pyrogallol (1 mol.) and KOH (3 mols.) in water forms a syrupy acid C₆H₆O₇, which yields BaC₆H₄O₇ (Boeke, B. 6, 486).—5. Purpurogallin or pyrogalloquinone is formed by oxidation by alcoholic AgNO₃, by KMnO₄ and H₂SO₄, or by excess of FeCl₂ (Girard, C. R. 69, 865). It is also got from pyrogallol and quinone (Wichelhaus, B. 5, 847; Nietzki, B. 20, 1278). It forms garnet-red needles (by sublimation), sl. sol. water, m. sol. alcohol and ether, forming yellow solutions. Alkalis impart a transient blue colour. Purpurogallin dyes cotton mordanted with alumina violet-blue. When an aqueous solution of pyrogallol is mixed with gum arabic and exposed to the air, purpurogallin C20H16O, separates, the yield in the course of two months being 67 p.c. of the pyrogallol used (Struve, A. 163, 160; De Clermont a. Chautard, C. R. 94, 1189, 1254). A solution of pyrogallol and Na₂HPO₄ also yields pyrogalloquinone on exposure to air (Loew, *J. pr.* [2] 15,322). Purpurogallin forms C₂₉H₁₂Ac₄O₄ and C₂₉H₁₂Br₄O₉.—6. An ammoniacal solution exposed to the air forms brown pyrogallein C₁₈H₂₀N₆O₁₀ (Rösing, J. 1858, 259).-7. Ammonium carbonate at 130° forms pyrogallol carboxylic acid C,H,O, (v. TRI-OXY-BENZOIC ACID) and pyrogallol dicarboxylic acid C_sH_sO , [270°] (Senhofer a. Brünner, M. 1, 468). 8. Distillation over zinc-dust yields benz-

ene.—9. ClCO₂Et acting on the K salt forms C₉H_sO₄ [105°] converted by aniline into diphenyl-urea and the mono-ethyl ether of pyrogallol (Bender, B. 13, 698).—10. A few drops of POOl, added to a mixture of pyrogallol (2 pts.) and acetone (1 pt.) react violently, forming gallacetonin C₉H₁₀O₃, which crystallises from 15 p.c. alcohol in whetstone-shaped crystals, insol. water, decomposing about 250°, and yielding C₉H₈AcO₈ (Wittenberg, J. pr. [2] 26, 76). Its asolutions are turned purple by FeCl₃ and reduce AgNO₃.—11. A few drops of H₂SO₄ added to a mixture of pyrogallol (12 g.) and acetoacetic ether (8 g.) form di-oxy-methyl-coumarin C₁₀H₂O₄, which crystallises from water in needles [235°] and yields O₁₀H₆Ac₂O₄ [176°] (Wittenberg, J. pr. [2] 26, 68; Pechmann, B. 16, 2127; 17, 2188).—12. Phthalic anhydride forms, on heating, $O_{20}H_{10}O$, (Baeyer, B. 4, 457, 663; A. 209, 261).—13. Phenyl cyanate at 100° forms $O_{6}H_{10}(O.CO.NHPh)_{6}[178^{\circ}]$ crystallising in minute needles (Snape, C. J. 47, 774).-14. Cyanogen passed into an aqueous solution forms C, H, O, Cy, or a polymeride thereof, as an unstable crystalline pp. (Loew, J. pr. [2] 15, 826).—15. Chloro-acetic acid ($2\frac{1}{2}$ pts.) followed by NaOHAq forms $C_0H_3(O.CH_2.CO_2H)_3$ crystallising in needles [198°], S. 1.3 at 14.5° and yielding K_sA''' and KH_A^M aq (Giacosa, J. pr. [2] 19, 398).—16. Benzotrichloride at 160° yields 'pyrogallol-benzeïn' C₃₈H₂₄O₁₁, crystallising in minute red plates with green lustre forming a blue solution in alkalis and a bluish-violet solution in alcohol. Zinc and HOAc reduce it to the anhydride of hexa-oxy-diphenyl-methane. Pyrogallol-benzein yields $C_{ss}H_{20}Ac_4O_{11}$ [208°], $C_{ss}H_{20}Bz_4O_{11}$ [251°], $C_{ss}H_{20}O_{11}(C_5H_9O)_4$ [228°] (Döbner a. Förster, A. 257, 60).—17. Benzoic acid and $ZnCl_2$ at 145° form tri-oxy-benzophenone (Alizarin yellow A) $C_8H_2(OH)_3.OO.C_8H_5[141^\circ]$, which crystallises with aq. The same body is got by heating pyrogallol with benzotrichloride (G. P. 50,451 [1889] and 54,661 [1890]; Graebe a. Eichengrün, A. 269, 297).—18. On heating pyrogallol with salicylic acid and zinc chloride tetra-oxy-benzophenone $C_0H_4(OH).CO.C_0H_2(OH)_8[149^\circ]$ is formed (G. s. E.). 19. Ac_2O forms $C_{10}H_{10}O_4$ [280°] crystallising in white prisms (Causse, Bl.[3]3,867). Fuming HClAq at 170° forms $C_2H_{14}O_4$ as a black powder, sol. NaOHAq (Böttinger, A. 202, 280).— 20. Benzene sulphonic chloride added to a solution of pyrogallol kept slightly alkaline forms $C_0H_3(0.SO_2Ph)_3$ [142°], sol. alcohol, sl. sol. ether (Georgesen, B. 24, 418).—21. Formic aldehyde and dilute HClAq form CH2(C6H2(OH)3)20 a microcrystalline powder, insol. water, sol. alcohol (Caro, B. 25, 947).—22. Aldehyde and dilute H₂SO₄ mixed with Na₂SO₄ added slowly at 60° gives colourless crystals of C₈H₈O₂2aq, which when dried at 30° over H₂SO₄ leaves violet C, H,O, aq (Causse, Bl. [3] 3, 865; cf. Michael a. Ryder, Am. 9, 133) .- 23. Benzoic alldehyde and a large quantity of HClAq form amorphous $C_{2a}H_{22}O_7$ (Baeyer, B. 5, 280). Benzoic aldehyde, alcohol, and a little conc. HClAq form a pp. of $C_{2s}H_{2s}O_{s}$, which yields $C_{2s}H_{14}Ac_{s}O_{s}$ (Michael a. Ryder, Am. 9, 130). On heating with benzoic acid alone pyrogallol forms resinous $C_{24}H_{22}O$, and red $C_{24}H_{16}O$, which may be reduced to colourless $C_{24}H_{22}O$, (Baeyer, B. 5, 25). 24. KOH (83 pts.) and K_2S_2O , (70 pts.) heated

with water (33 pts.) and pyrogallol (25 pts.) at 70° form C₆H₃(OH)₂(O.SO₃K), crystallising in needles, v. e. sol. water (Baumann, B. 11, 1913).—25. ClCONH₂ forms C₆H₃(O.CO.NH₂)₈. [1786], crystallising from alcohol in plates.

Salts.—C₀H₃(OH)₂(ONH₄). Crystals, got by passing NH₃ into an ethereal solution of pyrogallol (De Luynes, A. Surpl. 6, 252).— C_sH_s(OH)(O₂Pb) aq. Crystalline pp. got by adding lead nitrate to ammonium pyrogallol. Converted by conc. NH₃Aq into C₀H₄O₃3PbO.— (C₆H₅O₂)Pb.OAc. Ppd. by adding lead acetate to a solution of pyrogallol (Deering, C. J. 26, 702). —C_sH_s(SbO)Ô_s. Plates, got by ppg. pyrogallol solution with tartar-emetic (Rösing). SbCl_s (100 g.) dissolved in a saturated solution (250 c.c.) of NaCl, filtered, and slowly added to pyrogallol (50 g.) dissolved in saturated NaClAq (250 g.), forms a white pp. which changes to crystals of $C_6H_3(OH) < 0 > SbOH 2aq$, insol.water and alcohol, sol. mineral acids (Causse a. Bayard, Bl. [3] 7, 794). At 100° the above liquids on mixing yield $C_8H_3O_3Sb$, which is also crystalline, whence Ac_2O at 100° forms $C_8H_3O_3Ac_3$. AcCl in Ac2O converts both antimonites into $C_6H_5(OAc)_3$ $C_6H_6O_32NPhH_2$ [56°] (Mylius, B. 19, 1003).

Acetyl derivative CaH3(OAc)3. Got from pyrogallol and AcCl (Nachbauer, A. 107, 244).

Benzoyl derivative C,H,(OBz)s. Got, together with a mono- or di- benzoyl derivative [131.5°], from pyrogallol and BzCl (Skraup, M. 10, 391).

Di-methyl ether C,H,(OH)(OMe)2. (253°). Occurs in beech-wood creosote (Hof-mann, B. 11, 333; 12, 1373). Formed from pyrogallol (1 mol.), KOH, and MeI (2 mols.) at 160°. Prisms (from water). Its alkaline soluticn does not turn brown in air. Oxidised by means of $K_2Cr_2O_7$ and HOAc to corrulignone $C_{12}H_4O_2(OMe)_4$. Yields $C_4H_3(OAc)(OMe)_2$ as a sticky mass, and C₆H₃(OBz)(OMe)₂ [118°].

Tri-methyl ether C.H. (OMe) s. (235°). Formed by the action of MeI and KOH on pyrogallol dissolved in MeOH (Will, B. 21, 607). Needles, insol. water and alkalis, v. sol. alcohol and ether. Yields C₆Br₈(OMe), [81°]. Conc. HNO₃ forms C₆H₂(NO₂)(OMe), [100°] and the di-methyl derivative of dioxyquinone.

Mono-ethylether C₆H₃(OH)₂(OEt). [95°] Formed, together with the di- and tri-ethyl ethers, by heating pyrogallol with KOH and KEtSO, at 100° (Benedikt, B. 9, 125; M. 2, 212; Hofmann, B. 11, 797). Needles, m. sol. water, v. e. sol. alcohol and ether. FeSO, gives a

bluish-violet colour. Volatile with steam.

Di-ethyl ether C_eH₃(OH)(OEt)₂. [79°].

(262°). Crystals (from dilute alcohol). Oxidised by K₂Cr₂O₇ and HOAc to ethyl-corulignone. Nitrous acid passed into an ethereal solution forms C₂₀H₂₁O₅HNO₅, which forms a brown solution in water.

Tri-ethyl ether C₈H₃(OEt)₂. [39°]. Got from pyrogallol, alcoholic potash, and EtI (Herzig a. Zeisel, M. 10, 151). Needles, insol. KOHAq.

Ethylene ether $C_6H_3(OH):O_2:C_2H_4$. (267°). Formed from pyrogallol, alcoholic potash, and ethylene bromide (Magatti, B. 12, 1860). Yields C₆H₂(OBz):O₂C₂H₄ [109°].

References.—AMIDO-, BROMO-, CHLORO-, and NITRO- PYROGALLOL.

PYROGALLOL CARBOXYLIC ACID v. GALLIC ACID.

Pyrogallol dicarboxylic acid

C_aH(OH)_s(CO₂H)₂[1:2:3:4:5]. Gallocarboxylic acid. [270°]. S. 05 at 0°. H.C. 633,700. H.F. **Gallocarboxylic** 231,300 (Stohmann, J. pr. [2] 40, 128). Formed by heating pyrogallol or gallic acid with ammonium carbonate at 130° (Senhofer a. Brunner, M. 1, 468). Needles (containing 3aq), sl. sol. cold water. Gives a violet colour with FeCl_s.— K2A" 2aq: needles. — BaA"aq. — CaA" 6aq. — Ag₂A''

PYROGALLOL SULPHONIC ACID C.H.SO. i.e. $C_8H_2(OH)_3$, SO_8H_2 aq. Formed by dissolving pyrogallol in H_2SO_4 (Personne, BL. [2] 12, 169; 20, 531; Schiff, A. 178, 179). Hygroscopic crystals.—KA' 2aq: prisms, v. sol. water. Converted by heating with POCl, into C, H10S2O11, a flocculent mass, v. sol. alcohol, which is converted by warming with HOAc and Ac₂O into crystalline C₁₂H₃Ac₅S₂O₁₁ and C₁₂H₃Ac₄S₂O₁₁.

PYROGALLOQUINONE v. PYROGALLOL, Re-

action 5, and Purpurogallin.

PYROGENTISIC ACID is HYDROQUINONE. PYROGLUTAMIC ACID is OXY-TETRA-METHENYL DIHYDRIDE CARBOXYLIC ACID.

PYROGLYCERIN v. DIGLYCERIN.

PYROGUAIACIN C₁₈H₁₉O₃ i.e. C₁₈H₁₉O(OH)₃. [180·5°]. (258°) at 80-90 mm. V.D. 9·53 (calc. 9·76). A product of the distillation of gum guaiacum (Pelletier a. Deville, C. R. 17, 1143; Ebermaier, J. pr. 62, 291; Nachbauer, A. 106, 382; Hlasiwetz, A. 106, 381; 119, 277; Wieser, M. 1, 594). Plates, v. sl. sol. hot water, sl. sol. alcohol and ether. Forms a blue solution in H₂SO₄. On distillation with zinc-dust it yields guaiene $C_{12}H_{12}$ [100°], which is converted by oxidation into a lemon-yellow quinone $C_{12}H_{10}O_{2}$ [121°]. Potash-fusion forms $C_{12}H_{12}O_2$ [202°], which is reduced by zinc-dust to guaiene. Pyroguaiscin yields C₁₈H₁₈Ac₂O₈ [122°], C₁₈H₁₆Bz₂O₈ [179°], C₁₈H₁₅Br₂O₈ [172°], and the salt C₁₈H₁₆K₂O₃.

PYROLIC ACID v. SEBACIC ACID.

PYRO-MECAZONIC ACID C5H5NOs. Formed by heating oxycomenic acid, with NH3Aq and also, in small quantity, along with the isomeric amido-pyro-meconic acid, by the reduction of nitro-pyromeconic acid. Prepared from oxypyro-mecazonic acid by reducing with HI (Ost, J. pr. [2] 19, 203; 28, 441; 27, 258).

Properties.—Streaked trimetric tablets. Combines with mineral acids, but not with acetic acid. Gives rise to a di-acetyl derivative [155°] and a bromo-derivative C, H, BrNO. Is stable in acid solutions, but soon turns brown in alkaline solution. Gives a blue colour with FeCl, BaCl₂ and NH₃ give a pp. which turns bright

blue in air.

Reactions.—1. Suspended in ether and mixed with HNO, forms 'pyromecazone' C,H,NO, Pyromecazone is a brick-red powder, insol. ether, v. sol. water, gives no colour with FeCl, and turns the skin violet. With BaCl, and NH, it gives, on exposure to air, a crimson pp. Crystallises from alcohol as C_sH_sNO_s,EtOH. Aqueous SO₂ converts pyromecazone back into pyrome-cazonic acid. These properties resemble those of a quinone.-2. Suspended in HOAc and

treated with HNO, it forms yellowish prisms of C,H2(NO2)NO, aq. nitro-pyromecazone body is decomposed by water at 30° into nitropyro-mecazonic acid C₅H₄(NO₂)NO₃, CO₂ coming off through oxidation of some of the substance. SO₂ rapidly changes nitro-pyromecazone into nitro-pyromecazonic acid, so that the former resembles quinone in its ready reducibility. Nitro-pyromecazonic acid gives a blood-red colour with FeCl, nitro-pyromecazone gives no colour with FeCls. It yields the salt CsH3NaN2O5. Salt.—HA'HClaq: needles.

Oxypyromecazonic acid C5H5NO4. Formed by passing SO₂ into an ethereal solution of nitrosopyromeconic acid, and boiling the product (v. Nitroso-PYROMECONIC ACID) with CHCl, (Ost, J. pr. [2] 19, 177). Needles (containing aq or 2aq). Coloured dirty-violet by FeCl. - NaHA'2. --KHA'2-TlHA'2-CaA'2-HA'HCl.

PYROMECONIC ACID C,H,O,. [117°]. (225°

Preparation. - By distillation of meconic acid in a current of CO2, some comenic acid being also formed (Ost, J. pr. [2] 19, 182; 23, 441; cf. Robiquet, A. 5, 90; A. Ch. [2] 5, 282; 51, 236; Stenhouse, P. M. [3] 24, 128; A. 49, 18; F. D. Brown, P. M. [4] 4, 161; 8, 201; A. 84, 32; 92, 321; Ihlée, A. 188, 31). The yield is 14 p.c.

Properties.—Glittering prisms from water or alcohol. V. sol. chloroform, much less sol. ether. Gives a blood-red colour with FeCl. Feeble acid. Does not react with hydroxylamine (Oderheimer, B. 17, 2081). Bromine forms C.H.BrO, crystallising in prisms and yielding PbA', aq. Excess of Br yields C.H.BrO, aq PbA', aq. Excess of Br yields C,H,BrO, aq [109°]. ICl forms C,H,IO, yielding BaA', aq and PbA'.

Salts.—Forms two series of unstable salts.

alkaline to litmus: KA'.—HNaA'₃.—BaA'₂3aq.

—BaH₂A'₄.—CaA'₂aq.—CaH₂A'₄.—MgA'₂.—
SrA'₂aq.—PbA'₂.—FeA'₃.—CuA'₂.—C₅H₄O₃,HCl.
Formed by passing HCl into ethereal solution of the acid. Decomposed into its constituents by water or alcohol. Sulphates C, H,O,.H,SO, and (C,H,O₃),H,SO₄. Got by adding H,SO₄ to an ethereal solution (Ost, J. pr. [2] 19, 189).

Acetyl derivative C,H,AcO₂. [91°]. Ob-

tained by warming with AcCl

Mitroso-pyro-meconic acid $C_sH_s(NO)O_s$. An unstable compound of this body with an equivalent of pyro-meconic acid is formed by passing N2O2 into an ethereal solution of that acid (Ost, J. pr. [2] 19, 195). An aqueous solution of SO₂ converts this compound by addition of H₂ into C₅H₅(NO)O₅,C₅H₄O₅, a compound which is split up by boiling chloroform, leaving so-called oxy-mecaronic acid, C,H,NO4. This acid gives a violet colour with FeCl, and a blue pp. with BaCl, and NH, By Sn and HCl it is reduced to pyro-mecaronic acid C.H.NO.

References .- AMIDO- and NITEO-PYROMECONIC ACID

PYROMELLIC ACID C10HaOa. [238°]. tained in the electrolysis of aqueous KOH with carbon electrodes, and got also by oxidation of mellogen by KOCl (Bartoli a. Papasogli, G. 12, 113; 13, 51). Crystalline (containing 2aq), sol. water and alcohol. Yields the crystalline salt C, H, Na, O, and an ethyl ether crystallising in colourless needles [below 100°], insol. water.

40, 140; 43, 540). H.C.v. 778,900. H.F. 369,600. Formed by gently distilling mellitic acid (Erdmann, A. 80, 281) and by oxidising s-durene (Jacobsen, B. 17, 2517). Crystallises from water in triclinic tables (containing 2aq), v. sol. hot water. Forms various compounds on heating with (a)-naphthol (Grabowski, B. 4, 726; 6, 1065).—Ca₂A'' 6aq.—Pb₂A''aq.—Ag,A'': pp.

Methyl ether Me₄A''. [138°]. Plates, sl.

sol. hot alcohol (Baeyer, A. 166, 339).

Ethyl ether Et, A'v. [53°]. Formed from the Ag salt and Etl (Baeyer, A. Suppl. 7, 36). Needles (from alcohol).

Chloride C.H2(COCl)4. Got by heating the acid with PCl.. Crystalline mass, v. sol. ether.

Anhydride C₀H₂(C₂O₃)₂. [268°]. Got by distilling the acid. Needles (by sublimation), v. sol. hot water, being re-converted into the acid.

Tetrahydride $C_6H_6(CO_2H)_4$. Formed by the action of sodium-amalgam on the ammonium salt of pyromellitic acid (Baeyer). Amorphous hygroscopic mass, v. sol. water. When heated with H2SO4 it yields pyromellitic, trimellitic, and isophthalic acids.

Iso-tetrahydride C,H,(CO,H),. 200°]. Accompanies the preceding acid. Needles (containing 2aq). Behaves like the preceding hydride when heated with H₂SO₄. Yields a methyl ether [156°] which may be distilled.

References .- AMIDO- and NITRO- PYROMEL-

LITIC ACID.

PYROMUCIC ACID C,H,O, i.e.

CH.CH C.CO₂H. Furfurane (a)-carboxylic acid. Mol. w. 112. [132°]. S. 3.6 at 15°; 25 R_{∞} 40.18 (in a 2.19 p.c. aqueous at 100°. solution).

Formation.—1. By the distillation of mucic acid (Houton-Labillardière, A. Ch. [2] 9, 365; Pelouze, A. 9, 273; Boussingault, A. 15, 184; Liès-Bodart, A. 100, 327).—2. By distilling dehydromucic acid (Heinzelmann, A. 193, 184).— 3. By oxidation of furfuraldehyde by Ag₂O (Schwanert, A. 114, 63; 116, 257).—4. Together with furfuryl-carbinol by boiling furfuraldehyde with alcoholic potash (Ulrich, Z. 1861, 186; Schmelz a. Beilstein, A. Suppl. 3, 275; Limprioht, A. 165, 279).—5. By distilling iso-saccharic acid in a current of CO₂ (Tiemann a. Haarmann, B. 19, 1271).—6. Occurs, together with pyromycuric acid C₇H,NO₄ [165°] which yields BaA', 11aq, in the urine of dogs dosed with furfuraldehyde (Jaffé a. Cohn, B. 20, 2311).

Preparation.—By oxidising furfuraldehyde with alkaline KMnO, below 20° (Volhard, A. 261,

Properties.—Long white needles (by sublimation), v. sol. alcohol and ether. FeCl, gives no colour.

Reactions.—1. Bromine in excess forms cobromic acid. Dry Br gives C₅H₄Br₄O₅ mucobromic acid. [160°], oxidised by dilute CrO_s to di-bromopyromucic acid (Tönnies, B. 11, 1086).—2. Distillation with soda-lime yields furfurane.—3. On heating with lime and ammonia-zinc-chloride it yields pyrrole.
Salts.—KA': needles.—NaA'.—BaA'₂: orys-

tals, sol. water and alcohol.—CaA'2: crystalline

powder.—CuA'₂ Saq.—PbA'₂ 2aq.—AgA': scales. Ethyl ether EtA'. [34°]. (209°). Laminæ, v. sol. alcohol and ether, insol. water (Malaguti, A. 25, 276). Combines with dry chlorine, forming C.H. Cl.O. S.G. 195 1.496. Dry Br forms C.H. Br.O. [480] (Tönnies) and also the ethers of $(\beta\delta)$ -di-bromo-pyromucic acid [167°], of $(\beta\gamma)$ di-bromo-pyromucic acid [192°] and of (δ)-bromopyromucic acid [183°] (Hill a. Sanger, A. 232, 65).

Chloride C₅H₃O₂Cl. (170°).

[143°]. Amide C₄H₃O.CONH₂. Formed from the ether or chloride and NH, Aq (Wallach, B. 14, 751; 214, 227; Ciamician a. Dennstedt, B. 14, 1058; G. 11, 294). Large plates, sol. alcohol, sl. sol. ether. Converted by PCl₅ or P₂O₅ into the nitrile.

Ethylamide C₄H₃O.CONHEt. [258° cor.). Formed from pyromucic ether and ethylamine solution at 100°. Converted by PCl, into crystalline C4H2O.CCl2.NHEt. PCl3 in smaller quantity (1/3 pt.) forms the amidine C4H3O.C(NEt).NHEt, a liquid (240°) which gives B', H, PtCl_a.

Anilide C, H, O.CO.NHPh. [123.5°]. Long

needles (Schiff, B. 19, 849).

Furfuryl cyanide. Nitrile C.H.O.CN. (148°). Liquid with sweet taste, sl. sol. water, smelling like benzoic aldehyde. On reduction it gives CH2O.CH2.NH2.

(B)-Pyromucic acid, got by oxidation of fucusol, is a mixture of pyromucic acid with methyl-pyromucic acid [108°] (Bieler a. Tollens,

A. 258, 126).

Isopyromucic acid. [82°]. Formed, together with pyromucic acid, in the distillation of mucic acid. It decomposes BaCO₃ very slowly, and can be separated by shaking the product with BaCO, and extracting with ether (Limpricht, A. 165, 256). Plates (by sublimation), v. e. sol. water, alcohol, and ether. FeCl₃ gives a green colour. Br yields mucobromic acid.

References. — Bromo-, Bromo-sulpho-, NITEO-, and SULPHO- PYROMUCIC CHLORO-, ACIDS

PYROMUCIC ALDEHYDE v. FURFURALDE-HYDE.

PYROMYKURIC ACID C,H,NO. Occurs in the urine of rabbits dosed with furfuraldehyde (Jaffé a. Cohn, B. 20, 2311). Prisms (from water). Split up by boiling baryta-water into glycocoll and pyromucic acid.—BaA'2 12 aq: plates.—A urea compound (C,H,NO,)CO(NH2)2 [120°] occurs in the urine of dogs fed with meat. It crystallises from benzene in needles, v. sol. water and alcohol, sl. sol. ether, and is decomposed by heating with BaCO_s into urea and pyromykuric acid.

PYRONE. A name given to the ring co < CN:CH > 0.

PYROPHORUS. Any finely-divided substance which takes fire very easily when exposed to air is called a pyrophorus.

PYRORACEMIC ACID is PYRUVIC ACID. PYROTARTARIC ACID C, H,O, i.e.

CO₂H.CHMe.CH₂.CO₂H.

Mol. w. 132. [115°]. Methyl-succinic acid. S. 66 at 20°. S.H. 3098 (from 0° to 50°) (Hess. 4. Ch. [2] 35, 410). H.C.v. and p. 515,200

(Stohmann, J. pr. [2] 40, 209); 511,672 (Louguinine, C. R. 107, 597). H.F. 230,800 (S.).

Formation.—1. By dry distillation of tartar (Valentin Rose, Gehlen's Journ. 3, 598) and of tartaric and racemic acids (Gruver, N. J. T. 24, 2, 55; Pelouze, A. Ch. [2] 56, 297; Weniselos, A. 15, 148; Fourcroy a. Vauquelin, A. Ch. 35, 161; 64, 42; Arppe, A. 66, 73).—2. By the reduction of ita-, citra-, and mesa-conic acids and of dibromopyrotartaric acid with sodiumamalgam (Kekulé, A. Suppl. 1, 342; 2, 95).—3. By saponifying its nitrile with HClAq (Maxwell Simpson, A. 121, 161).—4. By heating powdered tartaric acid with HClAq at 180° (Geuther a. Riemann, Z. [2] 5, 318).-5. By potash-fusion from gamboge (Hlasiwetz a. Barth, A. 138, 73).— 6. By heating pyruvic acid alone at 170° or with HClAq at 100°, or by boiling it with baryta. 7. By boiling β-acetyl-α- or β- methyl-succinic ether with potash (Conrad, A. 188, 217; Kressner, A. 192, 135).—8. By heating allyl iodide (1 mol.) with KCy (2 mols.) and a little alcohol, and boiling the resulting nitrile with KOHAq (Claus, B. 5, 612; 8, 100; A. 191, 38).—9. By heating propane tricarboxylic acid (Bischoff a. Guthzeit, B. 20, 614).—10. By oxidation of β acetyl-n- and iso- butyric acid (Bischoff, A. 206,

Properties.—Triclinic prisms grouped like frog's feet (from ether), v. sol. water, alcohol, and ether. Begins to boil at 200°, yielding an anhydride together with some butyric acid (Claus, A. 191, 48). Its solution is not ppd. by lime-water or lead acctate, but gives a pp. with

lead subacetate.

Reactions.—1. Bromine acts very slowly in the cold, but when the acid (10 pts.) is heated with Br (24 pts.) and water (10 pts.) at 120° for 6 hours bromocitraconic anhydride is formed (Lagermark, Z. 6, 299). When the acid (1 mol.) is heated with Br (4 mols.) at 132° acetylene tetrabromide is formed (Bourgoin, A. Ch. [5] 12, 419).-2. The K salt yields, on electrolysis, O, CO, and CO₂ at the positive pole (Reboul a. Bourgoin, C. R. 84, 1231).—3. Distillation with P₂S₃ yields methyl-thiophene.—4. On heating the acid (5 pts.) with resorcin (9 pts.) and H2SO. (18 pts.) at 150° there is formed pyrotartrylfluorescein C₁₇H₁₆O_e, a brownish-red powder, sl. sol. water, v. sol. dilute acids and alkalis. Its dilute alkaline solutions fluoresce yellowishgreen. It yields C₁₇H₁₂Br₄O₆ (Hjelt, B. 17, 1280)

Salts.—K₂A" aq: deliquescent. — KHA": monoclinic prisms.--Na_A" 6aq: efflorescent laminæ.-NaHA": small prisms.-(NH4)2A": deliquescent; gives off NH3 on evaporation, leaving (NH₄)HA", which crystallises from water in prisms, permanent in the air.—BeA".—BeH₂A"₂.—CaA" 2aq: powder. S. 1 at 100°.—CaH₁₀A"₆ 2aq. BaH₂A''₂ 2aq. — SrH₂A''₂ 2aq. — BaH₂A''₂ 4aq. — MgA'' 3aq. — MgA'' 6aq. — Al(OH)A''. — Bi₄A''₄(HO)₁O. — Al(OH)A''. — Bi₄A''₄(HO)₁O. — Al(OH)A''. — Bi₄A''₄(HO)₁O. — Al(OH)A''. CdA" 3aq.—CdA" 2aq.—MnA" 3aq.—CuA" 2aq.—Cu₂(OH)₂A".—Fe(OH)A" 2aq.—PbA" 2aq.— Pb,A"O₂.—NiA"2aq.—NiH,A",2aq.—ZnA"3aq. —Ag,A": white curdy pp., blackened by light.

Ethyl ether Et.A". (218° oor.). S.G. 15 1.0189; 25 1.0113. M.M. 9.347 at 17.4° (Perkin, C. J. 45, 516; cf. Malaguti, A. 25, 274).

Chloride C, H,O2Cl2. (190°-193°) (Hjelt, B. 16, 2624). Reduced in ethereal solution by sodium-amalgam and HOAc to liquid CaHeOs (204°), S. 20, which is the anhydride of an acid C,H,,O,.

Anhydride C, H,O,. (245° cor.) (Lebedeff. A. 182, 327). Got by heating the acid at 200° alone or with P₂O₅. Heavy oil, slowly converted

by water into the acid.

CH₂·CH(CO.NH₂).CH₂·CONH₃ Amide[175°]. S. 7 at 10°. Large plates (Henry, C. R. 100, 943).

Imide CH₃.CH<CO.NH_{CH₂.CO}. [66°]. Formed by heating acid ammonium pyrotartrate (Arppe, A. 87, 228). Six-sided trimetric plates, sol. water, alcohol, and ether.

C,H,(CO.NHMe), Di-methylamide

[115°] (Henry, Bl. [2] 43, 619).

Mono-anilide CO₂H.CH₂.CHMe.CONHPh. [147°]. Formed from aniline and pyrotartaric anhydride in chloroform (Arppe, A. 90, 141; Anschütz, A. 246, 122). Got also by reducing the anilide of mesaconic acid with sodiumamalgam. Is identical with Reissert's 'pyranilpyroic lactone dihydride.' Bulky needles. m. sol. water.—PbA'2: white pp. —AgA'

Phenyl-imide CH, CH CO NPh [989] (Arppe, A. 90, 139); [104°] (Biffi, A. 91, 105). Formed by heating the acid with aniline at 100° for ten minutes. Minute needles.

p-Nit**r**o-anilide $C_2H_6(CO_2H)(CO_1NH,C_6H_4NO_2)$. [above 150°]. Formed by boiling the p-nitro-phenyl-imide with Na₂CO₂Aq. Minute tables, v. sl. sol. hot water. - AgA': white pp.

water.—AgA': wnite pp.

p-Nitro-phenyl-imide
C₃H₆:C₂O₂:NO₆H₄NO₂. [155°]. Formed by nihe phenyl-imide. Needles (from al-

cohol), almost insol. water. $p extbf{-}Bromo extbf{-}anilide$ C_sH_s(CO₂H).CO.NHC_sH_sBr. O.H).CO.NHC, H.Br. [158°]. Formed the anhydride and p-bromo-aniline from

(Anschütz a. Hensel, A. 248, 269).

Di-bromo-anilide

C₃H₆(CO₂H).CO.NHC₆H₃Br₂[1:2:4]. [139°]. Nitrile CH₃.CH(CN).CH₂CN. [c. [c. 12°]. (254°). Formed, amongst other products, by the action of KCN on allyl chloride (Pinner, B. 12, 2058). Prisms. Prepared from propylene bromide and alcoholic KCy at 100° (Maxwell Simpson, A. 121, 160). Liquid, sol. water, alcohol, and ether.

Ethyl ether of the seminitrile CO₂Et.CH₂.CHMe.CN. (198°). S.G. 24.5 1.0275. V.D. 4.6. Formed from a-bromo-propionic ether and alcoholic KCy (Zelinsky, B. 21, 8162). Oil. -NaA': very hygroscopic.

References.—Bromo-, Chloro-, Iodo-, Oxx-AMIDO-, and OXY- PYROTARTARIC ACID.

n-Pyrotartaric acid v. GLUTARIC ACID.

Isopyrotartaric acid v. DI-METHYL-MALONIC ACID

PYROTEREBIC ACID v. HEXENOIC ACID. PYROTRITARIC ACID C,H,O, i.e.

O CMe:CH Uvic acid. Di-methyl-furfurane carboxylic acid. Mol. w. 140. [135°]. S. 25 at 100°. A product of the distillation of tartaric acid (Wislicenus a. Stadnicki, A. 146,

306). Prepared by heating pyruvic acid with NaOAc and Ac_2O at 140° , the yield being 20 p.c. (Böttinger, B. 13, 1969; A. 172, 241; 208, 122; 247, 255), by heating pyruvic acid (80 g.) with dry sodium succinate (55 g.) at 110° (Fittig a. Parker, A. 267, 212), and by heating pyruvic acid (17.5 g.) with dry potassium propionate (20 g.) at 140° (Bischoff, B. 24, 2021). Formed also by heating methronic acid as long as CO2 is given off (Fittig, A. 250, 190; cf. Harrow, C. J. 33, 425). Needles (from hot water), nearly insol. cold water, v. sol. alcohol and ether. Not affected by potash-fusion. PCl, gives a chloride, reconverted by water into the acid. Does not react with hydroxylamine or phenyl-hydrazine. Water at 160° forms acetonyl-acetone. bromine it is converted into a tetra-bromoderivative and its tetra-bromide, which are both reduced back to pyrotritaric acid by sodiumamalgam. An excess of bromine at 100° forms penta-bromo-pyrotritaric acid. By dry distillation it gives di-methyl-furfurane, a substance which is also obtained by distilling acetonylacetone with ZnCl₂ (Paal, B. 20, 1074; Dietrich a. Paal, B. 20, 1077).

Salts.-NaA' 2aq.-CaA', 2aq: prisms.-CaA', 4aq. — CaA', 6aq. — BaA, 4aq: plates.— BaA', 2aq.—BaA', 5aq.—ZnA', 8aq.—AgA'. Methyl ether MeA'. (192°). Got by dis-

tilling the Ag salt of carbopyrotritaric monomethyl ether (Knorr a. Cavallo, B. 22, 156). Ethyl ether EtA'. (208°

Pyrotritaric carboxylic acid v. METHRONIC

Carbopyrotritaric acid v. DI-METHYL-FUR-FURANE CARBOXYLIC ACID.

PYROXANTHIN C₁₅H₁₂O₈. [162°]. Extracted by NaOHAq from the product of the dry distillation of wood (Scanlan, J. pr. 7, 94; Gregory, A. 21, 143; Hill, Am. 3, 332; B. 11, 456). Orange needles (from alcohol), sol. benzene and HOAc. Forms a purple solution in H₂SO₄. Reduced by zinc-dust and HOAc to a colourless body. Bromine forms C₁₅H₁₀Br₆O₃ crystallising in small colourless triclinic needles, converted by alcohol and powdered Sb into C₁₅H₁₀Br₂O₂, crystallising in yellow monoclinic needles, v. sol. hot alcohol.

PYROXYLIN v. CELLULOSE.

PYRRODIAZOLE C₂H₃N₅ i.e. N \leqslant CH:N CH.NH. [121°]. Formed by heating its carboxylic acid at

120° (Andreocci, B. 25, 229). Needles, v. e. sol. water and alcohol.

Pyrrodiazole carboxylic acid

 $CO_2H.C \stackrel{N.NH}{\leqslant_{N:CH}}$. Formed by oxidising methylpyrrodiazole with alkaline KMnO. crystalline powder, v. sl. sol. water, alcohol, and

PYRROLE C,H,N i.e. CH:CH>NH. Pyrrhol. Pyrroline. Mol. w. 67. (130° i.V.). S.G. 12'5 ·9752. S.V. 92·1 (Ramsay). H.F.p. 4,056 (Ramsay). Occurs among the products of distillation of coal (Runge, P. 31, 67) and of bones (Anderson, Tr. E. 20 [2] 247; 21 [4] 571; A. 80, 63; 105, 349).

Formation.-1. By distilling ammonium mucate alone or with glycerin at 190° (Schwanert, A. 116, 279; Goldschmidt, Z. [2] 8, 280).—2. By heating its carboxylic acid.—3. By distilling PYRROLE. 365

succinimide with zinc-dust (Chichester Bell, B. 13, 877).—4. By distilling ammonium saccharate (Bell a. Lapper, B. 10, 1962).—5. By heating di-chloro-maleic imide with PCl₃ at 200°, and reducing the resulting perchloride C₄Cl₄N with zinc-dust and HOAc (Ciamician a. Silber, B. 17, 554; 19, 3027). The dichloro-maleic imide may be got from succinimide and PCl₃.—6. By reducing tetra-iodo-pyrrole with zinc-dust and KOHAq (C. a. S.).—7. By heating albumen with Ba(OH)₂ at 150° (Schützenberger, Bl. [2] 25, 289).—8. By distilling pyromucic acid with ammonia-zinc chloride and CaO (Canzoneri a. Oliveri, G. 16, 487).—9. By passing di-ethylamine through a red-hot tube (Bell, B. 10, 1868). 10. By passing ethyl-allyl-amine over PbO at 450° (Koenigs, B. 12, 2344).—11. By distilling glutamic acid (Haitinger, M. 3, 228).

Preparation.—Bone oil is shaken with acids and distilled. The fraction 98°-150° is heated with KOHAq and distilled, and the portion 125°-140° heated with solid KOH with inverted condenser. The solid C.H.NK thus obtained is washed with ether and decomposed by water, and the pyrrole distilled with steam (Ciamician, B.

13, 70; 19, 173; G. 16, 336).

Properties.—Oil with fragrant odour, resembling chloroform. Tastes hot and pungent. V. sol. alcohol and ether, insol. dilute alkalis. Turns brown in air. Turns pine-wood soaked in HClAq red. Dissolves in cold dilute acids, but on warming the solution pyrrole-red is formed as a bulky amorphous pp. FeCl, added to its solution in HClAq gives a green colour changing to black. Alcoholic HgCl₂ gives a white pp. Isatin and HOAc or dilute H₂SO₄ give a deep-blue pp. which forms a deep-blue solution in HOAc and H₂SO₄ (V. Meyer, B. 16, 2974; Ciamician, B. 17, 142).

Reactions .- 1. HNO, forms oxalic acid. 2. Potassium acts very strongly, forming a colourless liquid, which solidifies on cooling to C.H.NK. Boiling with solid KOH gives the same body (Anderson, A. 105, 352). Sodium acts but slightly, forming C.H.NNa only at a very high temperature. NaOHAq has no action on pyrrole.—3. HCl passed into an ethereal solution ppts. crystalline (C₄H₅N)₅HCl, whence ammonia liberates an unstable base (Dennstedt a. Zimmermann, B. 21, 1478).—4. Hydroxylamine forms NH, and crystalline C₄H₈N₂O₂ [175°]. Converted by adding Na to its boiling alcoholic solution into tetramethylene-diamine (Ciamician, G. 14, 156; B. 22, 1968). The compound C4H8N2O2, when heated with phenyl-hydrazine, yields C₁₆H₁₈N₄, which crystallises from alcohol in plates [125°].—5. By a dilute solution of sodium hypochlorite (15 pts. of active chlorine in 1,000 pts. of water to 10 pts. of pyrrol) it is converted into di-chloro-maleic acid and chlorinated pyrroles. On the other hand, a concentrated solution of sodium hypochlorite (45 pts. of active chlorine in 500 pts. of water to 10 pts. of pyrrole) yields dichloro-acetic acid. An alkaline solution of bromine yields dibromomaleimide. Iodine in presence of alkalis yields tetra-iodopyrrol (Ciamician a. Silber, B. 17, 1743; 18, 1763).-6. Potassium pyrrole heated with ethyl iodide yields p-ethyl-pyrrole, di-ethyl-pyrrole C,H,Et:NEt, and probably also C,H,Et:NH. Potassium pyrrole and benzyl-chloride yield

ν-benzyl-pyrrole (247°) (Ciamician, B. 22, 659; G. 17, 185).—7. Pyrrole (50 g.), boiled with paraldehyde (50 g.) and ZnCl₂ (12 g.), forms oily ethyl-pyrrole C₂H₂N (164°) (Dennstedt a. Zimperson B. 2000). ethyl-pyrrole C₆H₅N (104⁻) (LPEILISECUE & ZAILI-mermann, B. 19, 2189).—8. Pyrrole (1 pt.), acctone (10 pts.) and a drop of HCl give C₁₄H₁₈N₂ [291⁵], whence alcoholic AgNO₃ forms (C₁₄H₁₈N₂)₂AgNO₃, crystallising in needles (Baeyer, B. 19, 2184). The compound C₁₄H₁₈N₂ yields, on distillation, C₁₀H₁₂N (275°-285°), which is probably mesityl-pyrrole, and whence Ac2O and NaOAc form ν-acetyl-pyrrole. The compound C₁₄H₁₅N₂, examined by Raoult's method, appears to have the formula C₂₅H₃₆N₄. On heating pyrrole with acetone and ZnCl₂ isopropyl-pyrrole (174°) is formed. Among the products of the action of acetone on pyrrole there is also a yellow oil $C_{18}H_{17}N$ (?) (300°-305°), which soon blackens in air. This oil forms an acetyl derivative [above 360°], and is reduced by tin and alcoholic HCl to C₁₃H₁₈N (274°), which yields B'HCl [228°] and B'HSnCl₃ [170°] (Dennstedt a. Zimmermann, B. 20, 850, 2449; 23, 1370).— 9. Di-ethyl ketone, MeOH, and HCl give C₁₈H₂₈N₂ [210°], which yields B'₂AgNO₃, crystallising in needles (Dennstedt a. Zimmermann, B. 20, 2455).—10. Chloroform converts potassium pyrrole into (β) -chloro-pyridine. Bromofo**rm** acts in like manner, giving bromo-pyridine. CH₂Cl₂ has no action, but CCl₄ forms chloropyridine, while benzylidene chloride yields (β)phenyl-pyridine (Ciamician, B. 14, 1160; 15, 1172; 20, 191; G. 16, 140).—11. Readily combines with diazo- compounds, forming azo- and disazobodies, but no compounds analogous to the diazoamides. Their constitution is probably $NH \stackrel{C(N_2X):CH}{\subset CH} = CH$ and $NH \stackrel{C(N_2X):CH}{\subset (N_2X):CH}$ (O. Fischer a. Hepp, B. 19, 2251).—12. Pyrrole (5 g.) added to a lukewarm solution of alloxan (11 g.) in water (300 c.c.) forms colourless crystals of C_sH₁N₂O₄ or NH₂·CO.NH.CO.CO.CO.C₄H₂NH, sl. sol. hot alcohol, nearly insol. water and ether, and yielding Ag_2A'' . Pyrrole-alloxan is converted by caustic potash solution into $C_1H_4N_2O_2$ or $NH_2CO.CO.CO.C_4H_3NH$, which crystallises in white plates, v. sol. warm alcohol, and yields AgA' and crystalline MeA' (Ciamician, B. 17, 106, 1711; 19, 1708; G. 16, 198, 857).—

13. Pyrrole and MeOH distilled over zinc-dust yield (a)- and (β)- methyl-pyrrole, di-methylpyrrole, and di-methyl-dipyrrole (Dennstedt, B. 24, 2559).—14. Acetyl chloride, acting on potassium-pyrrole, forms v-acetyl-pyrrole and pyrryl methyl ketone (pseudo-acetyl-pyrrole), which may be separated by steam-distillation. Ac O at 800° forms pyrrylene di-methyl diketone (Ciamician a. Silber, G. 15, 193).—15. Propionic anhydride and sodium propionate form, on boiling, v-propionylpyrrole, pyrryl ethyl ketone, and pyrrylene diethyl diketone (Dennstedt a. Zimmermann, B. 20, 1760).—16. Bz₂O and NaOBz at 220° form pyrryl phenyl ketone (pseudo-benzoyl-pyrrole). 17. COCl, in benzene, added to C,H,NK in ether, forms CO(NC4H4)2, separating from ligroin in monoclinic crystals [63°] (238°) (Ciamician a. Magnaghi, B. 18, 415).—18. Phthalic anhydride and HOAc at 185° form $C_{12}H_1NO_2$ [241°], converted by Br into $C_{12}H_2Br_2NO_2$ [199°], and by boiling KOHAq into $C_{12}H_2NO_3$ [174°–184°], which yields MeA' [105°] (Ciamician, B. 17, 2957;

Anderlini, B. 21, 2869). The compound C₁₂H₇NO₂ is reduced by water and Na to $C_4H_4N:C<_{C_6H_4}^0$ CH.OH [118°] (Ciamician, B.

19, 2206; 21, 1554).

Salts.—C,H,NK. Solid, absorbing CO₂ at 200°, forming pyrrole (β)-carboxylic acid.— B'2HgCl₂: crystalline powder, insol. water, sl. sol. cold alcohol.—B'43CdCl2.—B'ICl: black pp. (Dittmar, B. 18, 1612).—Picrate: [c. 71°]; unstable (Hooker, B. 21, 3299).

Acetyl derivative C.H.NAc. (182° i.V.). Formed by the action of AcCl on potassiumpyrrole suspended in ether, and also as a byproduct in the preparation of pyrryl methyl ketone by heating pyrrole with Ac,N and NaOAc (Ciamician, B. 16, 2348; 18, 881). Oil, volatile with steam, saponified by hot KOHAq. Reddens acidified pine-wood. Gives a pp. with aqueous $\mathbf{HgCl_2}$.

PropionylderivativeC₄H₄N.COEt. (193°) (Dennstedt a. Zimmermann, \vec{B} . 20, 1760).

Dihydride C,H,N. Pyrroline. (90° i.V.). Formed by reducing pyrrole with zinc-dust and HOAc (Ciamician a. Dennstedt, B. 16, 1536; 22, 2512; G. 15, 481). Hygroscopic liquid, v. sol. water. Yields a nitrosamine C₄H₆N(NO) Absorbs CO₂ from the air.—B'HCl. [174°]. Flat prisms, v. sol. hot alcohol.—
B'₂H₂PtCl₆: triclinic crystals.—B'HAuCl₄ [152°].
—B'C₆H₃N₃O₇. [156°]. Yellow crystals (from water).
Benzoyl derivative C₄H₆NBz. (160° at 2 mm.). Syrup, formed from the hy-

drochloride and BzCl at 110°. V. sol. alcohol.

Tetrahydride C.H.N. Pyrrolidine.
(88.5°). S.G. 22.5 8520. Formed by reducing the dihydride by HIAq at P at 250° (Ciamician a. Magnaghi, G. 15, 483; B. 18, 2079). Formed also, together with tetra-methylene-diamine, by the action of Na on ethylene cyanide or succinimide in alcohol, and by distilling tetramethylenediamine hydrochloride (Ladenburg, B. 19, 782; 20,442,2215; Petersen, B.21,290). Formed from chlorobutylamine hydrochloride and KOHAq (Gabriel, B. 24, 3234). Liquid, with pungent ammoniacal odour. Yields a liquid nitrosamine Prisms. — B'HAuCl₄. [206°]. — B'C₄H₂N₄O₇. [112°].—B'₄H₃Bi₄I₉.—B'H₂CdI₄. [219°]. Needles. Pyrrole red C₁₂H₄N₂O? Formed by boiling provide on its carbon size and solid side of the control of

pyrrole or its carboxylic acid with dilute H2SO4 or HCl (Anderson, A. 105, 357; Schwanert, A. 116, 280). Reddish-brown flakes, insol. water, ether, acids, and alkalis, sl. sol. alcohol. Yields

pyrrole on distillation.

Homopyrrole v. METHYL-PYRROLE.

References .- Bromo-, Chloro-, Iodo, Nitro-, and Oxy-amido- Pyrrole.

PYRROLE - AZO- v. Azo- compounds and DISAZO- COMPOUNDS.

PYRROL-CARBO-KETONIC ACID v. PYRRYL-GLYOXYLIC ACID.

PYRROLE (a)-CARBOXYLIC ACID C.H.NO. i.e. C, H3 (CO2H)NH. Carbopyrrolic acid. [191°].

Formation.—1. By the action of boiling baryta on its amide, which is got by distilling ammonium mucate (Malaguti, A. 15, 179; Schwanert, A.116, 270; Ciamician, B.17, 1(1).—
2. By boiling pyrocoll with KOHAq (Weidel a. Ciamician, M. 1, 285).—3. By fusing potassium (a)-methyl-pyrrole with potash (Ciamician, B.

14, 1054; G. 11, 228).—4. By heating pyrrole with ammonium carbonate and water at 140° (Ciamician a. Silber, B. 17, 1150; G. 14, 162).-5. By heating pyrrole with alcoholic potash and CCl₄(Ciamician a. Silber, B. 17, 1437).—6. By the action of CO₂ on potassium pyrrole at a high temperature (C. a S.).

Properties.-Prisms (from water), sol. water and alcohol. Decomposed by heat into CO2 and pyrrole. HClAq forms, on warming, CO2, pyrrole

red, and NH,Aq.

Salts.-NH,A': m. sol. water.-CaA'2: scales.—BaA'₂ (dried at 100°): plates, sol. water and alcohol.—PbA'₂: nacreous scales, v. sol. water.—AgA': small needles, sl. sol. water.

Methyl ether Med'. [173°]. Prisms.

Ethyl ether Etd'. [39°] (231°).

Amide C.H.(CONH.)N. [178°]. Lamiv. sol. alcohol and ether, m. sol. water.

Acetyl derivative C.H.N.CO.OAc. [75°].

Formed from the Ag salt and AcCl. Scales, readily decomposed by water into acetic acid and pyrrole carboxylic acid. At 75° it decom-

poses into HOAc and pyrocoll.

Pyrrole (β)-carboxylic acid C₄H₈(CO₂H):NH. [162°]. Formed by fusing potassium (β)-methylpyrrole with potash (Ciamician, B. 14, 1054). Needles, partially decomposed by boiling water into CO₂ and pyrrole. The lead salt is sl. sol. water.—BaA'₂: needles. A pyrrole carboxylic acid [166°], got by potash-fusion from isopropylpyrrole, and yielding a methyl ether [129°] (Dennstedt a. Zimmermann, B. 20, 855), is probably identical with the (β) - acid.

Pyrrole v-carboxylic acid

ether C₄H₄N.CO₂m₄. 80°). Formed from potassium-(Ciaminian a. Ethylurethane. (180°). pyrrole and ClCO₂Et in ether (Ciamician a. Dennstedt, G. 12, 84). Oil.

Amide C,H,N.CONH2. [167°]. Formed

from the ether and NH,

Nitrile C,H,N.CN. Tetrol cyanuramide. [210°]. Got by passing CyCl into potassiumpyrrole in ether (Ciamician a. Dennstedt, G. 13, 102). Needles, insol. water, sol. hot alcohol. Boiling alcoholic potash forms pyrrole, CO2, and NHg.

Pyrrole dicarboxylic acid C.H.NO. i.e. C₄H₂(CO₂H)₂NH. Formed by oxidising pyrrylene dimethyl diketone, and fusing the product with potash (Ciamician a. Silber, G. 16, 377; B. 19, 1958; 20, 2601). Needles (from dilute alcohol), sol. ether and hot water. Blackens at 260°, giving pyrrole and CO₂. FeCl₂ gives a brown pp. The Ba salt crystallises in needles. The Ag salt is a curdy pp.

Mono-methyl ether MeHA". [243°].

Di-methyl ether Me₂A". [132°]. Di-ethyl ether Et₂A". [82°].

References .- Bromo-, CHLORO, and NITRO-PYRROLE CARBOXYLIC ACIDS.

PYRROLINE. A name used both for Pyrrole and Pyrrole DIHYDRIDE.

PYRROLYLENE v. BUTINENE.

PYRROLYLENE TETRABROMIDE v. Tetra-BROMO-BUTANE.

PYRRONE v. DI-PYRRYL-KETONE.

PYRROYL-FORMIC ACID v. PYRRYL-GLY-OXYLIC ACID.

PYRROYL - PYRROL C,H,N.CO.C,H,NH [68°]. Formed, together with di-pyrryl-ketone

CO(C₄H₂NH), by intramolecular transformation of carbonyl-pyrrole by heating it to 250° for a few hours (Ciamician a. Magnaghi, B. 18, 1829). White silky plates. Not volatile with steam. By heating with aqueous KOH it is split up'into

pyrrole and pyrrole-(a)-carboxylic acid.

PYRROYL-PYRUVIC ACID. Ethyl ether C4H3NH.CO.CH2.CO.CO2Et. [123°]. Formed by the action of NaOEt and oxalic ether on pyrryl methyl ketone (Angeli, B. 23, 1794, 2155). Yellowish plates, v. sol. alcohol, sl. sol. water. Decomposed by alkalis, even in the cold, into oxalic acid and pyrryl methyl ketone. Hydroxylamine forms C₁₀H₁₀N₂O₃ [124°], which on saponification yields the acid CH CH.N.N.C.CO.2H [179°], crystallising in white needles. Aniline (2 pts.) in HOAc forms C4H3NH.CO.CH2.C(NPh).CO2Et [115°]. Phenyl-hydrazine forms C₁₆H₁₅N₃O₂ [168°] which is the ether of an acid [215°].

Anhydride $CH \leqslant_{CH.N.CO.CO}^{CH:C.CO.CH_2}$. Formed by adding HCl to the mother-liquor from which the ether has separated (Angeli, B. 23, 1795). Yellow needles (from benzene), v. sol. alcohol. o-Phenylene-diamine forms $C_{14}H_pN_sO$, a reddish-yellow crystalline powder forming a bluish-green solution in H.SO, and yielding a benzoyl derivative [c. 166°]. Aniline forms C_4H_3N < $CO.CH_2$ [218°], whence cold KOHAq

forms an acid C₁₄H₁₂N₂O₃ [179°].

PYRRYLENE DIETHYL DIKETONE

C₄H₂NH(COEt)₂. [117°]. Formed by heating pyrrole with propionic anhydride at 260° (Dennstedt a. Zimmermann, B. 20, 1761). Plates.

PYRRYLENE DI-METHYL DIKETONE

C₈H₉NO₂ i.e. NH CAc:CH. $[162^{\circ}].$ Formed by heating pyrrole, pyrryl methyl ketone, or acetyl-pyrrole with Ac2O at 250° (Ciamician, B. 17, 432, 2953; 18, 881, 1466; 19, 1957; 20, 2595). Needles, sol. hot water and hot KOHAq. May be sublimed. Yields a nitro-derivative [149°] and a di-bromo-derivative [172°].—AgA'.

PYRRYLENE DISTYRYL DIKETONE (C_eH₃,CH:CH.CO)₂C₄H₂NH. [240°]. Formed by heating pyrrylene dimethyl diketone with benzoic aldehyde and KOHAq (Ciamician a. Dennstedt, B. 17, 2953). Crystals (from HOAc), sl. sol. alcohol. Forms a violet solution in H.SO.

PYRRYL ETHYL KETONE C, H, NO i.e. C,H,.CO.C,H,NH. [52°]. (224°). Formed by boiling pyrrole with propionic anhydride and sodium propionate (Dennstedt a. Zimmermann, B. 20, 1761). Colourless needles.—AgA': pp.

PYRRYL-GLYOXYLIC ACID C.H.NO. aq i.e C,H,NH.CO.CO,H aq or C,H,NH.C(OH),CO,H. Formed by oxidation of pyrryl methyl ketone by KMnO, (Ciamician a. Dennstedt, B. 16, 2350; 17, 2949). Crystallises from benzene in yellow needles (of CoH, NO4) melting at 76°, v. sol. hot water. After drying over sulphuric acid the acid (C,H,NO,) decomposes at 114°. The aqueous solution gives a red colour with FeCl. On heating with HClAq it gives a crimson colour turned yellowish-green by alkalis.-AgA': colourless needles, sol. hot water.

Methyl ether MeA'. [72°]. (285°). Mono-

clinic crystals, sl. sol. water.

Carboxy-pyrryl-glyoxylic acid

CO2H.C4H2NH.CO.CO2H. Formed by oxidation of pyrrylene dimethyl diketone (Ciamician a. Silber, B. 19, 1412, 1961). Crystalline, v. sol. hot water.—Ag₂A": canary-yellow pp.

Methyl ether Me₂A". [145°]. DI-PYRRYL KETONE CO(C,H,NH)₂. [160°]. Formed, together with (C₄H₄N)₂CO, by the action of COCl₂ on C₄H₄NK. Formed also, together with pyrroyl-pyrrole, by heating (C₄H₄N)₂CO at 250° for a few hours (Ciamician a. Magnaghi, B. 18, 414, 1829). Trimetric crystals; a:b:c = 2.531:1:2.901. V.sol. alcohol, ether, and benzene, nearly insol. water. - Ag₂A": yellow pp.

PYRRYL METHYL KETONE C₆H,NO i.e.

CH₃.CO.C₄H₃NH or NH<CAc:CH CII :CH. Pscudoacetyl-pyrrole. [90°]. (220° uncor.). V.D. 8.8 (calc. 3.8). Formed by boiling pyrrole with Ac2O and NaOAc (R. Schiff, B. 10, 1501; Ciamician a. Dennstedt, B. 16, 2348; 17, 432, 2944; 18, 1456; 20, 2605; G. 15, 175). Formed also by heating pyrrole with Ac₂O and ZnCl₂ (Dennstedt a. Zimmermann, B. 19, 2204), and by heating CH₃.CO.C.H₃N.CO.K with K₂CO₃ at 290° (Ciamician a. Silber, B. 19, 1963). Long monoclinic needles, sol. hot water, v. sol. aqueous alkalis, sparingly volatile with steam. Not converted

into pyrrole and KOAc by potash.

Reactions.—1. Yields pyrryl-glyoxylic acid when oxidised by KMnO₄.—2. Benzoic aldehyde and dilute KOH form pyrryl styryl ketone.—3. Water and sodium-amalgam reduce it to crystalline pyrryl methyl pinacone $C_{12}H_{16}N_2O_2$ [120°] and pyrryl-methyl-carbinol CH₃.CH(OH).C₄H₂NH Pyrryl-methyl-pinacone an oil (290°-300°). C₄H₄N.CMe(OH).CMe(OH).C₄H₄N crystallises in prisms (containing 2aq) melting at 98° when hydrated.—4. Oxalic ether and NaOEt form

pyrroyl-pyruvic ether (Angel, B. 23, 1357, 1794). 5. Cold fuming H₂SO₄ yields an unstable sulphonic acid, which forms a crystalline K salt. Salt.—C4H3AcNAg. Crystalline pp

Oxim CH_s.C(NOH).C,H_sNH. [146°]. Needles. Phenyl-hydrazide

CH₂.C(N₂HPh).Č₄H₂NH. [147°]. White powder. References.—Amido-, Bromo-, Bromo-nitro-,

and Nitro- Pyrryl METHYL KETONE PYRRYL METHYL KETONE CARBOXYLIC **ACID** C,H,NO, i.e. CH, CO.C,H,N.CO,H. [186°]. Formed by saponification of its methyl ether, obtained by heating methyl pyrrole (a)-carboxylate with Ac₂O at 260° (Ciamician a. Silber, G. 14, 169; B. 17, 1155). Leaflets, sol. water, alcohol, and ether. FeCl, gives a brown pp.-CaA'₂7aq: triclinic prisms.—*PbA'₂: needles.—AgA': powder, sl. sol. water.

Methyl ether MeA'. [118°]. Gammoniacal AgNO, a pp. C.H.AgNO. Gives with

PYRRYL PHENYL KETONE C, H, NO i.e. C.H. CO.C.H.NH. [78°]. Formed by heating pyrrole with benzoic anhydride and dry NaOBz at 200°-240° (Ciamician a. Dennstedt, B. 17, 432, 2955). Needles, v. sol. alcohol, sl. sol. hot water.—C, H, AgNO.

PYRRYL STYRYL KETONE C₁₉H₁₁NO i.e. C_cH_a.CH:CH.CO.C,H_aNH. [142°]. Formed by boiling pyrryl methyl ketone with C_cH_aCHO and KOHAq (Ciamician a. Dennstedt, B. 17, 2947). Yellow needles, sl. sol. alcohol, insol. water.-C1. H10 ONAg: yellow needles, insol. NH.Aq.

PYRUVIC ACID C₂H₄O₃ i.e. CH₈:CO.CO₂H. Pyroracemic acid. Di-oxy-propionic acid. Mol. w. 88. [9°]. (c. 165°). S.G. $\frac{15}{25}$ 1·2752; $\frac{25}{25}$ 1·2700. M.M. 3·557 (Perkin, C. J. 61, 836).

Formation.—1. By distilling racemic or tartaric acid (Berzelius, A. 13, 61; Völckel, A. 89, 65; Wislicenus, A. 126, 225.—2. By gradually heating tartaric acid with HClAq in sealed tubes to 180° (Geuther a. Riemann, Z. [2] 5, 318).—3. By heating tartaric acid with conc. H₂SO₄ at 40°–50° (Bouchardat, C. R. 89, 99).—4. By heating silver di-a-chloro-propionate with water (Beckurts a. Otto, B. 10, 265; 18, 227).—5. By distilling glyceric acid (Moldenhauer, A. 131, 338; Böttinger, A. 196, 92).—6. By boiling acetyl cyanide with dilute HCl (Claisen a. Shadwell, B. 11, 1563).—7. By oxidation of a cold aqueous solution of calcium lactate by KMnO₄ (Beilstein a. Wiegand, B. 17, 840).

Preparation.—1. Tartaric acid (3 lbs.) is distilled from a glass retort (30 lbs. capacity). The distillation takes three hours. A fresh quantity of tartaric acid is then added and the operation repeated. The distillate is fractionally distilled (Clewing, J. pr. [2] 17, 243).—2. Dried, and finely powdered, tartaric acid is mixed with an equal weight of sand and distilled (Seissl, A. 249, 297).
3. Tartaric acid is distilled with KHSO₄, the yield being 50 to 60 p.c. (Erlenmeyer, B. 14, 321).

Properties.—Liquid, smelling like acetic acid, v. e. sol. water, alcohol, and ether. Colourless crystals below 9° (Simon, Bl. [3] 9, 111). Partially decomposed on distillation. Reduces ammoniacal AgNO₂, forming a mirror, and yielding CO₂ and acetic acid. Coloured cherry-red by alkaline sodium-nitroprusside, the colour being discharged by HOAc (Von Bitto, A. 267, 377). Readily ppd. by phenyl-hydrazine. Forms an oxim.

Redctions.-1. Dilute H2SO4 at 150° splits it up into aldehyde and CO₂ (B. a. W.).—2. By long heating at 170° it is converted into CO. and acetic, pyrotartaric, uvic, and citraconic acids (Böttinger, B. 9, 670, 837, 1823).—3. HClAq at 100° forms CO₂ and pyrotartaric acids (De Clermont, B. 6, 72).—4. Boiling with Ag₂O yields CO₂ and acetic acid.—5. Reduced to lactic acid by sodium-amalgam, and by Zn and H2SO4. 6. HI reduces it to propionic acid. -7. Zinc when added to its alcoholic solution forms CO₂H.CMe(OH).CMe(OH).CO₂H (Böttinger, B. 9, 1064).-8. PCl, forms di-a-chloro-propionic acid (Klimenko, B. 3, 465; Beckurts a. Otto, B. 11, 386).—9. Dry bromine at 0° forms crystalline di-bromo-lactic acid (?). Br and water give di-bromo-pyruvic acid (Wislicenus, A. 148, 208; Wichelhaus, A. 152, 265).—10. Oxidised by chromic acid mixture to OO, and acetic acid.— 11. HNO forms oxalic acid and CO .- 12. Boiling baryta-water forms uvitic, uvic, pyrotartaric, oxalic, and acetic acids and CO₂ (Finckh, A. 122, 182; Böttinger, B. 8, 957; A. 172, 241, 253; 188, 313; 208, 129). Baryta-water added to a cold solution of pyruvic acid ppts. barium hydruvate BaC₂H₂O₂, or the basic salt 18. On heating with excess of quicklime it yields a little aldehyde (Hanriot, C. R. 101, 1156; Bl. [2] 45, 81).—14. HON followed by HCl forms lactic acid. KCN followed by HCl yields

CH₂.C(OH)(CN).CO₂H v. CYANO-OXY-PROPIONIC On warming pyruvic acid with HCN ACID. under pressure the product is CMe(NH2)(CO2H)2, crystallising in unstable prisms (Körner a. Menozzi, G. 17, 104).—15. Alcoholic ammonia forms methyl - pyridine dicarboxylic acid.-16. Aniline added to an ethereal solution forms CoH,NO, i.e. CH, C(NPh).CO,H [122°] (Böttinger, B. 10, 818). Aniline alone forms, on heating, C₁₄H₁₈N₂O, crystallising from dilute alcohol in needles [195°], while p-toluidine gives $C_{16}H_{20}N_2O$ [238°] (Lazarus, B. 17, 998). Aniline and fur- $C_6H_4 < N \xrightarrow{C(CO_2H):CH} C.C_4H_5O_4$ **furaldeh**yde form crystallising in greenish-yellow needles [210°-215°] (Doebner, A. 242, 285). Tetra-amido-anisole forms the quinoxaline C'H(OMe)(N2C3H4O)2 as orange-yellow flakes, v. sol. alkalis (Nietzki a. Kurtenacker, B. 25, 284).—17. H.S passed into water containing the Agsalt forms thiolactic acid (Böttinger, B. 9, 404; Lovèn, J. pr. [2] 29, 376).—18. PH_3 and HCl passed into an ethereal solution form 'phosphortrianhydropyruvic' acid $\mathbf{C}_{\scriptscriptstyle{0}}\mathbf{H}_{\scriptscriptstyle{9}}\mathbf{PO}_{\scriptscriptstyle{6}}$, crystallising in needles, insol. alcohol and ether, and forming with phenyl-hydrazine the compounds $C_9H_9PO_8^3PhN_2H_3$ [132°] and $C_{45}H_{45}N_{12}O_3$ (?) [162°]. The soid $C_9H_9PO_8$ is converted by aniline into C_2 , H_{28} N₂PO₆[158°], whence phenyl-hydrazine forms C_2 ; H_{28} N₀2 [169°]. Tolylene-diamine converts C_6 ; H_9 PO₆ into C_9 ; H_9 PO₆2C, H_{10} N₂ [178°] (Messinger a. Engels, B. 21, 334, 2919).—19. Pyruvic acid (50 g.) heated with dry sodium succinate (92 g.) and Ac.O (58g.) at 110° forms di-methyl-maleic anhydride (pyrocinchonic anhydride) (Fittig a. Parker, A. 267, 204). Pyruvic acid (30g.) heated with dry sodium succinate (55 g.) at 110° forms uvic (pyrotritaric) acid.—20. Pyruvic acid (30g.) heated with sodium pyrotartrate (60 g.) and AcO (36g.) at 140° forms methyl-ethyl-maleïc anhydride (F. s. P.).—21. Benzonitrile and cold H₂SO₄ form an acid C₁₇H₁₆N₂O₄, crystallising in tables [172°], insol. water, v. sol. acetone (Böttinger, B. 14, 1599).—22. Phenyl-acetonitrile and H₂SO₄ form, in like manner, C₁₀H₂₀N₂O₄ [145°] (B.).—23. By heating with aldehydes R.CHO and baryta-water it is converted into s-alkyl-isophthalic acids of the form $C_6H_3R(CO_2H)_2$ (Doebner, B. 23, 2378).—24. Benzoic aldehyde and aniline in ether form, in the cold, C₂₂H₁₈N₂O or CHPh:CH.C(NPh).CONHPh [225°], insol. water, acids, and alkalis, sl. sol. alcohol and ether. Cuminic aldehyde and aniline give C₂H₂₄N₂O [216°] (Doebner a. Gieseke, A. 242, 290; 249, 102). Benzoic aldehyde and p-toluidine form, in like manner, C24H22N2O [205°] .- 25. Isobutyric aldehyde and aniline in alcoholic solution react forming CHPr.CH.C(NPh).CONHPh (?) [222°] crystallising from HOAc in needles, while isovaleric aldehyde and aniline in ether give C20H22N2O [160°] (Doebner, A. 242, 275). Isovaleric aldehyde in warm alcoholic solution forms the acid $C_{14}H_{15}NO_2$.—26. Thioglycollic acid gives rise to CO_2H .CMe(OH).S.CH₂.CO₂H [110°] with evolution of heat; while HCl passed through a mixture of pyruvic and thioglycollic acids produces CO₂H.CMe(S.CH₂CO₂H)₂ [182°] crystallising from ether (Bongartz, B. 19, 1938; 21, 494).—27. Phenyl mercaptan forms the compound CH₂.C(SPh)(OH).OO₂H, while p-bromo-phenyl mercaptan forms CH₂.C(SC₆H₄Br)(OH).CO₂H

[114.5°] (Baumann, B. 18, 263).—28. Ethylene mercaptan forms a substance [96°] which crystallises from benzene as a fine powder, and may be

condensed to $\overset{\text{CH}_2.S}{\text{CH}_2.S}$ CMe.CO₂H [102°], which

on oxidation gives ethylene ethylidene disulphone [198°] (Fasbender, B. 21, 1473).—29. Benzene and H₂SO₄ form CH₂CPh₂CO₂H, while phenol and H₂SO₄ yield C₁₂H₁₄O₄ (Böttinger, B. 14, 1595; 16, 2071).—30. Distillation of the K salt with KOAc forms acetone and pyrotartaric acid (Wichelhaus, Z.[2] 5, 254).—31. Pyruvic acid (1 pt.) heated with *urea* (2 pts.) at 100° forms pyvuril or 'pyruvic diureide 'O,H_sN₄O₃, which crystallises in tables, S. 10 at 100°, insol. alcohol and ether (Grimaux, C. R. 79, 526, 1304, 1478; 80, 53; A. Ch. [5] 11, 373). Pyvuril is decomposed by long heating at 160°, forming amorphous 'tetrapyruvic tetraureide' C_{1e}H_{1e}N_sO_s, insol. water. Boiling dilute HClAq converts pyvuril into urea and 'dipyruvic triureide' $C_0H_{12}N_0O_3$, which crystallises from hot water in needles, S. 4 at 100°, decomposed by boiling alkalis into urea, pyruvic acid, and pyvuril. Boiling conc. HClAq converts pyvuril into urea and 'pyruvic ureide'

 $C_1H_4N_2O_2$ or CO < N = CMe NH.COcrystalline

powder, m. sol. hot water, insol. alcohol. Nitric acid converts pyvuril into nitro-pyruvic ureide C,H₂(NO₂)N₂O₂ [above 200°], which is decomposed by boiling bromine-water into parabanic acid and CBr. NO2. Pyruvic acid heated with an equal weight of urea at 100° forms amorphous insoluble 'tripyruvic tetraureide' C13H16N8O, and dipyruvic triureide .- 32. Sodium hippurate and Ac₂O at 100° form the dibasic acid C₁₂H₉NO₄ [157°] (A. Hoffmann, B. 19, 2555).—33. solution of indoxyl forms on adding conc. HCl the indogenide $C_0H_4 < \stackrel{CO}{NH} > C:CMe.CO_2H$ [197°],

crystallising in red needles forming a blue solution in H₂SO₄ (Baeyer, B. 16, 2199).—34. Glycerin and KHSO₄ form 'glycuvic acid,' or 'pyruvin,' which is also got by heating glycerin with glyceric acid at 120°, and by distilling glycerin with citric acid. Glycuvic acid is probably

 $CH_3.CO.CO.O.CH_2.CH < \stackrel{CH_2}{O}$. It is crystalline [82°] (241°), and yields pyruvic acid when boiled with baryta (Böttinger, B. 10, 286; 14, 816; A. 263, 246; Jowanovitch, M. 6, 467; Erhardt, M. 6, 511; Schlagdenhauffen, C. R. 74, 672; De

Clermont, C. R. 105, 520).

VOL. IV.

Salts.—The salts crystallise well, provided heat is avoided in their preparation; otherwise The acid also is they become amorphous. changed by evaporation of its aqueous solution into a non-volatile syrupy mass.—NaA': large prisms.—BaA'2 aq: scales.—BaA'2 2aq: amorphous.—PbA'₂ aq: crystalline pp.—ZnA'₂ 3aq: white microcrystalline powder, sl. sol. water (Beckurts a. Otto, B. 18, 227).—CuA'₂ aq.— AgA': scales, sl. sol. water.

Combinations with bisulphites (Clewing, J. pr. [2] 17, 241).—HA'NaHSO, aq: crystals.-NaA'NaHSO, aq. — NaA'NaHSO, 1 aq. HA'KHSO,: octahedra. - KA'KHSO, aq. CaA'₂Oa(HSO₃)₂ Saq.—(CaA'₂)₃(Ca(HSO₃)₂)₂24aq.—(CaA'₂)₃(Ca(HSO₃)₂)₂Saq.—SrA'₂Sr(HSO₃)₂5aq.—BaA'₂Ba(HSO₃)₂.

Methyl ether MeA'. (c. 136°). S.G. P 1.154. Got from AgA' and MeI.

Ethyl ether EtA'. Oil. Converted by aniline into two crystalline bodies [144°] and [c. 250°]. The compound O(CMe(OH).CO,Et)2, formed by the action of nitrous acid on amidopropionic ether, is an oil (80°-86° at 120 mm.) (Curtius, J. pr. [2] 38, 472).

Isoamyl ether C.H.A'. Got by distilling the acid with isoamyl alcohol (Simon, Bl. [3] 9,

Amide CH3.CO.CO.NH2. [125°]. Got from acetyl cyanide and HCl (Claisen a. Shadwell, B. 11, 1566). Prisms or tables (from alcohol).

Nitrile v. ACETYL CYANIDE.

Phenyl hydraside CH, C(N₂HPh).CO₂H. [192°] (F.); [185°] (Japp a. Klingemann, B. 20, 3284; A. 247, 208; C. J. 53, 519). Ppd. when a solution of phenyl-hydrazine hydrochloride is added to a dilute (even 1 in 1000) solution of pyruvic acid (Fischer, B. 16, 2241; 17, 578). Prisms, sol. hot water and hot alcohol. Reduced by sodium-amalgam to phenyl-hydrazido-propionic acid [172°]. At 190° it gives off CO₂ and H_2 , and forms the phenyl-hydrazide of di-methyl diketone [242°] and the oily phenyl-hydrazide of aldehyde. The ether CH, C(N, HPh). CO, Et [117°], crystallises in yellow needles (from ligroin). o-Chloro-phenyl-hydrazide

C₆H₄Cl.NH.N:CMe.CO₂H. [178°]. Lemon-yellow needles, forming a crystalline ethyl ether [168°] (Hewett, C. J. 59, 211).

p-Nitro-phenyl-hydraside C_aH₄(NO₂).NH.N:CMe.CO₂H. Yel Yellow (Fischer a. Ach, A. 253, 64). V. sol. hot alcohol.

p-Sulpho-phenyl-hydraside SO₂H.C₆H₄.NH.N:CMe.CO₂H. Solid, insol. ether, v. sol. water and alcohol (Pfülf, A. 239, 217). NaA' aq: nodules.

Phenyl-methyl-hydrazide C₁₀H₁₂N₂O₂ NPhMe.N:CMe.CO₂H. [78°]. Yellowish i.e. NPhMe.N:CMe.CO₂H. [78°]. Yellowish needles (Fischer, B. 16, 2245; 17, 559). Converted by heating with HClAq into methyl-indole carboxylic acid and NH, The phenyl-ethylhydrazide is converted in like manner into ethyl-indole carboxylic acid.

Di-phenyl-hydrazide

NPh, N:CMe.ČO, H. [145°]. White needles (Fischer a. Hess, B. 17, 567). Sol. hot benzene and CHCl, forming deep-yellow solutions. Converted into phenyl-indole carboxylic acid by heating with HClAq.

o-Tolyl-hydraside $C_6H_4MeN_2H:CMe.CO_9H.$ [156°] (J.); [159°] (R.) Formed by saponifying the product of the action of o-diazotoluene chloride on sodium methylacetoacetic ether (Japp, A. 247, 213), and from o-tolyl-hydrazine hydrochloride and pyruvic acid (Raschen, A. 239, 228). Small yellow plates (from benzene). Yields o-tolyl-hydrazido-propionic acid [143°] on reduction. At about 160° it forms the di-o-tolyl-di-hydrazine of dimethyl-di-ketone, a yellow crystalline powder

p. Tolyl-hydrazide C,H,N,H:CMe.CO,H. [162°]. Formed in the same way as the o-isomeride. Yellow plates (from benzene). At 165° it is converted into the di-p-tolyl-hydrazide of di-methyl-di-ketone [230°]. Gives an ethyl ether C₁₂H₁₆N₂O₂ [106°], crystallising from ligroin

in yellowish plates.

BB

Methyl-p-tolyl-hydraside C11H14N2O2 Yellow prisms (Hegel, A. 232, 215).

Ethyl-p-tolyl-hydrazide C₁₂H₁₆N₂O₂. Needles, v. sol. alcohol and ether (H.).

ψ-Cumyl-hydrazide

C_eH₂Me₂NH.N:CMe.CO₂H. [148°]. Yellow needles (Ruhemann, C. J. 57, 55).

p-Benzoylphenyl-hydrazide

C.H.Bz.N.H:ČMe.CO.H. [200°]. Lemon-coloured crystals (Ruhemann a. Blackman, C. J. 55, 616). (a)-Naphthyl-hydrazide

C₁₀H, N₂H; CMe. ČO₂H. [159°] (Fischer, A. 232, 236). Yields EtA [100°] (Schlieper, A. 239,

(β)-Naphthyl-hydrazide [166°]. Yields

EtA' [131°] (Schlieper, A. 236, 176).

(B. 4)-Quinolyl-hydraside C_pH_eN.NH.N:CMe.CO_cH. [174°]. Formed from pyruvio acid and (B. 4)-hydrazido-quinoline acetate (Dufton, C. J. 59, 758). Light-yellow

pp., insol. water, sol. acids and alkalis.

Oxim CH₃.C(NOH).CO₂H. a-Nitroso-propionic acid. Formed by the action of cold KOHAq on its ether, which is got by adding a dilute solution of KNO2 (10 pts.) to methylacetoacetic ether dissolved in water (3 pts.), KOH (2 pts.), and some alcohol, the yield being 16 p.c. of the methyl-acetoacetic ether used (V. Meyer a. Züblin, B. 11, 692; Gutknecht, B. 13, 1116). Got also from pyruvic acid and hydroxylamine (V. Meyer a. Janny, B. 15, 1527: Schäfer, A. 264, 153; Hantzsch, B. 24, 50). Crystalline powder, sl. sol. ether, v. sol. water and alcohol. Decomposes suddenly at about 177°. Oxidised by potassium permanganate to ethyl-nitrolic acid CH₃.CH(NO)(NO₂). Yields acetonitrile on heating with hydroxylamine hydrochloride. Reduced by tin and HClAq to alanine. Salts .-KA' aq: pearly plates, sol. water.—BaA'; soluble prisms.—Cu₂OA'; aq.—AgA': white insoluble powder. Ethyl ether EtA'. [94°]. (233° cor.). Formed as above, and also by the action of HNO, on methyl-malonic ether and on propionyl-propionic ether (Bergreen, B. 20, 533). Needles or prisms, with weak acid properties.

References .- Bromo-, Chloro-, and Oxy-

PYRUVIO ACID.

PYRUVIC ALDEHYDE CH, CO.CHO. Methyl-glyoxal. Got by the action of dilute acids upon CH2.C(OH)(SO3Na).CH(SO3Na).NHSO3Na, which is got from nitroso-acetone and NaHSO. (Pechmann, B. 20, 2548).

Mono-oxim v. NITROSO-ACETONE.

Di-oxim CH, C(NOH).CH:NOH. Methylglyoxim. [153°]. Formed by the action of hydroxylamine on CH, CO.CH.NOH or upon CH, CO.CHCl. (V. Meyer a. Janny, B. 15, 1165; Treadwell, B. 15, 2787). Small prisms (from alcohol) or needles (by sublimation). Salt .-AgC,H,N,O, pp.—Diacetyl derivative

C₂H₄(NOAc)₂. [51°]. (Schramm, B. 16, 2187). [51°]. Prisms (from ligroin)

Phenyl-hydrazide of the oxim

CH, C(N, HPh). CH: NOH. [134°]. Formed from nitroso-acetone and phenyl-hydrazine (Pechmann a. Wehsarg, B. 21, 2994; A. 262, 278). Prisms or needles (from alcohol). H₂SO, forms a reddish-yellow solution, coloured deep blue by Ac₂O yields CH₃.C(N₂HPh).CH:NOAc [163°], crystallising in colourless needles.

Phenyl-methyl-hydrazide of the oxim CH₃.C(N₂PhMe).CH:NOH. [118°]. Orange

yellow prisms (from alcohol).

Phenyl-hydrazide CH₂.CO.CH:N₂HPh [150°]. Formed from sodium acetoacetic ether and diazobenzene chloride (Japp, A. 247, 198, 218; C. J. 53, 519). Plates (from benzene or MeOH). Yields CH₂.CO.CH:N₂AcPh [93°], orystallising from ligroin in needles. NaOEt and chloro-acetic ether yield, on saponification of the product, CH₃.CO.CH:N.NPh.CO₂H [162°], which crystallises from hot water in needles, and is reduced by tin and HClAq to phenyl-amido-acetic acid [127°].

Di-phenyl-di-hydraside CH. C(N. HPh). CH: N. HPh. [145°]. Formed by heating the phenyl-hydrazide of acetyl-carbinol with phenyl-hydrazine hydrochloride and NaOAc in alcohol at 100° (Laubmann, A. 243, 248). Formed also by warming pyruvic aldehyde or its mono-oxim, phenyl-hydrazide, or phenyl-hydrazide of the oxim with phenyl-hydrazine acetate (Pechmann, B. 20, 2543; 21, 2755; Japp, A. 247, 207). Yellow plates or needles (from dilute alcohol), sl. sol. alcohol. Conc. H₂SO₄ forms an olive-green solution, changing through slatyblue to violet. Yields a crystalline hydrochloride [197°]. Oxidised by K₂Cr₂O, and HOAC to ÇH :N.NPh crystallising in needles [107°], CMe:N.NPh

whence boiling HClAq produces CH:N>NPh a colourless oil (150° at 60 mm.), oxidised by

alkaline KMnO, to the acid C,H,N,O, [1927].

Acetyl derivative of the di-phenyldi-hydraside CH,C(N,HPh).CH:N.NPhAc. [229°]. Formed from the acetyl derivative of the phenyl-hydrazide and phenyl-hydrazine (Japp, C. J. 53, 519). Yellowish needles.

Phenyl-methyl-hydraside CH, CO.CH:N.NPhMe. [64°]. Formed from the phenyl-hydrazide, NaOMe, MeOH, and MeI (Japp). Flat needles (from MeOH), v. e. sol. alcohol. Converted by phenyl-hydrazine into CH, C(N, HPh).CH:N.NPhMe [152°], crystallising in pale-yellow needles.

Phenyl-ethyl-hydraside CH, CO.CH:N.NPhEt. [55°]. Prepared in like manner. Prisms (from ligroin).

PYRUVYL ALCOHOL v. AGETYL CARBINOL.

QUARTENYLIC ACID v. ISOCROTONIC ACID. CHCl₂. Occurs in quassia-wood (Winckler, Rep. QUASSIN C_{E2}H₄₂O₁₀ or C₂₂H₄₂O₁₀ or C₂₂H₄₂O₁₀ Pharm. 54, 85; Wiggers, A. 21, 40; Christensen, QUASSIN C. H., O, or C., H., O

(10 pts.) with boiling water (45 pts.), evaporating (to 10 pts.) at a gentle heat, ppg. by tannin, mixing the pp. with lead carbonate, and extracting with boiling alcohol. The product is recrystallised from water and alcohol (yield 01 pt.)

(Oliveri a. Denaro, G. 15, 6).

Properties.-Slender monoclinic needles, v. sol. alcohol and chloroform, sl. sol. ether. Sol. conc. HClAq and KOHAq, insol. Na₂CO₃Aq. Reduces Fehling's solution (Oliveri), or, according to Allen (An. 12, 107), has no reducing power.
Not coloured by FeCl₃. Ppd. by tannin. Quassin appears to be the dimethyl ether of quassic acid, and probably contains two hydroxyls, two CO₂Me, and two CO groups, and is perhaps a derivative of anthraquinone (Oliveri). Phenyl-hydrazine forms $C_{22}H_{40}O_8(N_2HPh)_2$ (?), a yellow amorphous powder, decomposing at about 230° without fusion (Oliveri, G. 18, 169).

Reactions.—1. Boiling dilute (4 p.c.) H₂SO₄ forms quasside' C₂₂H₄₂O₅, a white amorphous very bitter substance [194°], which reduces Fehling's solution, and is reconverted by boiling dilute alcohol into quassin. - 2. Ac₂O and NaOAc form, on boiling, the amorphous anhydride $C_{22}H_{40}O_{2}$ [150°-158°], sol. alcohol and ether.—3. Bromine forms $C_{32}H_{41}Br_{3}O_{9}$, a very bitter yellow powder [155°].—4. Fuming HClAq in a sealed tube at 100° forms MeCl and quassic acid.—5. PCl_s gives $O_{22}H_{30}Cl_sO_{8}$, a yellow powder [120°] decomposed on fusion.—6. HIAq and P at 150°–280° form durene (188°–195°) $C_{14}H_{16}$ (220°–240°) and anthracene (Oliveri, G. 17, 575).

Quassic acid $C_{50}H_{88}O_{10}$ or $C_{30}H_{40}O_{10}$. [245°]. S. .0043 at 23°. Formed, together with McCl (2 mols.), by heating quassin with HClAq. Monoclinic prisms (containing aq), sol. hot alcohol, sl. sol. ether. Alkalis give a reddish-yellow colour. FeCl, gives a greenish-yellow colour. Reduces Fehling's solution and ammoniacal AgNO, Hydroxylamine forms C₂₀H₂₆(NOH)₂O₂ (?), crystallising in rectangular prisms [230°].—BaA"7aq: reddish-yellow crystals.—PbA"6aq: white

amorphous pp.—Fe,A",: brownish-green pp.
QUEBRACHAMINE. [142°]. Occurs in white quebracho bark (Hesse, A. 211, 269). Plates, v. sol. alcohol, benzene, CHCl, and ether. alcoholic solution is alkaline and tastes bitter.

H₂SO, and K₂Or₂O, give a dark-violet colour. QUEBRACHINE O₂₁H₂₂N₂O₃. [216°]. = 62.5 in a 2 p.c. alcoholic solution at 15°; = 18.6 in chloroform. Occurs in white quebracho bark (Hesse, B. 13, 2308; A. 211, 254). Colour-less needles, sol. alcohol, ether, benzene, and CHOl₂, v. sl. sol. cold water, NaOHAq, and NH, Aq. Dextrorotatory. Its solutions are alkaline, bitter, and poisonous. Its solution in H₂SO₄ is turned blue by PbO₂ or K₂Cr₂O₇, and finally brown. FeCl₂ gives no colour.

Salts.—B'HCl.—B'₂H₂PtCl₄ 5aq: crystals.—

B'2H2SO, 8aq: cubes or short prisms, v. sol. hot water.—B',H,C,O,.—Tartrate B',H,C,O, 6aq.—Citrate B',C,H,O,: nodular groups of needles.

Hypoquebrachine C₁, H₂, N₂,O₂. [80°]. Strong base with bitter taste, v. sol. alcohol and ether, forming yellow amorphous salts. Platino-

othloride.—B'H.PtCl, 4aq (Hesse, A. 211, 264).

QUEBRACHITE C.H., MeO. [187°]. S.G.

2 1.54. (c. 210° in vacuo). [a]p = -80°. Occurs
in the seeds of Aspidosperma quebracho, from which it can be extracted with alcohol (Tanret,

C. R. 109, 908). Trimetric prisms with sweet taste, v. sol. water and alcohol, insol. ether. Does not ferment with yeast or reduce Fehling's Reduces ammoniacal AgNOs. solution. affected by dilute acids or alkalis at 100°. by ammoniacal lead acetate, but not by lead subacetate. Heated with HI it gives off MeI and a little benzene. Ac₂O and ZnCl₂ form an acetyl derivative [89°]. Gives on heating with HNO, the same reactions as inosite.

QUEBRACHO BARK. Quebracho blanco, or white quebracho bark, used as a febrifuge, contains about .8 p.c. of alkaloids, consisting of Aspr-DOSPERMINE, ASPIDOSPERMATINE, ASPIDOSAMINE, QUEBRACHINE, HYPOQUEBRACHINE, QUEBRACHAMINE,

and QUEBRACHOL (Hesse, A. 211, 251).
QUEBRACHO GUM. Occurs in quebracho colorado, the bark of Loxopterygium Lorentii (Jean, Bl. 28, 6; Arata, Anales de la Sociedad cientifica Argentina, July 1878; Feb. 1879; C. J. 34, 986). Brittle red concretions, with astringent taste, sol. boiling water and alcohol, insol. ether. Quebracho gum contains quebrachitannic acid, a pale-red amorphous mass, m. sol. hot water, giving a green colour with FeCl, turned dark red by NaOAc. Quebrachitannic acid ppts. Pb(OAc)2, gelatin, albumen, and alkaloids. yields pyrocatechin and a liquid (100°-120°) on distillation. Potash-fusion gives protocatechuic acid and phloroglucin; nitric acid yields oxalic and picric acids.

C20H31O. QUEBRACHOL [125°]. = -29.3°. Occurs in white quebracho bark (Hesse, A. 211, 272; 228, 288). Plates, v. e. sol. alcohol and ether, insol. water and alkalis. When the solution in chloroform is shaken with H,SO. the chloroform is coloured red (cf. Cholestern).

A cetyl derivative C₂₀H₁₂.OAc. [115°]. QUERCETAGETIN C₂₇H₂₂O₁₃ 4aq. Yell Yellow crystals, extracted by alcohol from the blossoms of the common marigold, Tagetes patula (Latour

a. Magnier, Bl. [2] 28, 337). QUERCETIN $C_{2i}H_{16}O_{11}$ 8aq (Liebermann a. Hamburger, B. 12, 1178) or $C_{15}H_{10}O_7$ (Herzig, M. 12, 172). (6)-Rhamnetin (Herzig, M. 10, 561). [above 250°]. S. (alcohol) 5.5 at 78°; 4 in the cold (Stein, J. 1862, 499). Mol. w. (by Raoult's method) 258 (calc. 302). Formed, together with isodulcite, by the action of dilute H2SO, on quercitron, a yellow dye-stuff consisting of the shavings of the bark of Quercus tinctoria, growing in the United States (Rigaud, A. 90, 289). Butin and robinin also yield quercetin when treated with dilute H₂SO₄ (Zwenger a. Dronke, A. 123, 153; Suppl. 1, 261; Schunck, C. J. 53, 262; Hlasiwetz, A. 112, 96; J. pr. 94, 65). Occurs in Persian berries (Kane, A. Ch. [3] 8, 380; Bolley, C. J. 13, 327), in the ripe fruit, flowers, and leaves of the horse-chestnut (Rochleder, A. 112, 112), in the berries of the seabuckthorn (Hippophae rhamnoides), in appletree bark, in tea-leaves, and in catechu (Loewy, Fr. 12, 127).

Properties. — Lemon - yellow crystalline powder, sl. sol. water, v. sl. sol. ether. Gives off water of crystallisation at 130°. May be sublimed as yellow needles. Dissolves in alkalis, forming yellow solutions. Dyes fabrics mordanted with alumina yellow; with iron, grey or black. Its alcoholic solution is coloured dark green by FeCl, the colour becoming dark red on warming.

Pb(OAc)₂ gives a brick-red pp. Reduces AgNO₂ in the cold, and Fehling's solution on heating.

Reactions.—1. On fusion with potash it yields phloroglucin and quercetic acid, and by prolonged fusion, paradatiscetin C15H10Os, quercimeric and protocatechuic acid.—2. Boiling dilute alcoholic potash gives phloroglucin and protocatechuicacid (Herzig, M. 6, 863).—3. KClO₃ and HCl yield protocatechnic acid. 4. Sodiumamalgam forms phloroglucin, a compound C13H12O5 crystallising in needles, sl. sol. water, and a compound C, H,O,, which forms granular crystals [130°], v. sol. water.—5. Bromins in HOAc forms $C_{24}H_{12}Br_{2}O_{11}$, crystallising in lemonyellow needles [237°] and yielding $C_{24}H_{12}Ac_{3}Br_{3}O_{11}$. Excess of Brin HOAc forms $C_{24}H_{11}Br_{3}O_{11}$, whence C₂,H₂Ac₂Br₂O₁₁ may be got, crystallising from HOAc in needles [253°]. Liebermann obtained the compounds C₂,H₁,Br₂O₁₁, C₂,H₁₂Ac₂Br₂O₁₁ [218°], C₂,H₁₂Br₄O₁₁, and C₂H₁,Ac₂Br₄O₁₁ [228°], all crystallising in needles.—6. Phenyl cyanate at 160° forms C₂₄H₁₁O₆(O.CO.NHPh)₅, a white amorphous powder [200°-205°] (Tesmer, B. 18, 2609).-7. Ammonia at 150° forms amorphous 'quercetamide' (Schützenberger a. Paraf, Z.

Salts.-C₂₇H₂₀K₂O₁₃ (?).-C₂₇H₂₀Na₂O₁₃ (?).-

 $C_{27}H_{20}ZnO_{14}$ (H. a. F.).

Acetyl derivative C₂H₂Ac₄O₁₁ or C₁₃H₂Ac₄O₇ (Herzig, M. 5, 72; 6, 890; 9, 537; 10, 561; 12, 174; cf. Liebermann, B. 12, 1178; A. 196, 319). [191°]. Formed by boiling queretin or phomother in the control of the contro cetin or rhamnetin with Ac.O, and NaOAc. Needles (from alcohol). Liebermann obtained

C₂₄H₁₄Ac₂O₁₁ [198°], crystallising in needles.

Methyl ether O₁₂H₂O₂(OMe). Rhamnetin.

Formed, together with isodulcite, by heating xanthorhamnin with dilute H₂SO₄ (Gellatly, N. E. P. J. 7,256; Liebermann a. Hörmann, B. 11, 1618; Herzig, M. 6, 889; 9, 560; 12, 175. Lemon-yellow powder, nearly insol. water, alcohol, and ether, v. sol. hot phenol. Forms a yellow solution in KOHAq. Reduces ammoniacal AgNO, and hot Fehling's solution. Yields protocatechuic acid and phloroglucin on fusion with potash or on treatment with sodium-amalgam (Smorawski, B. 12, 1595). Converted by HI into quercetin and MeI. Dyes iron mordants black and alumina yellow. Yields C15H5Ac4O6(OMe) KOH and EtI give ethyl-rhamnetin [108°], whence Ac2O and NaOAc form an acetyl-ethyl-rhamnetin [157°]. Rhamnetin yields also (tetra-?) propionyl [158°-162°] and (tetra-?) benzoyl [212°] derivatives, and an acetyl-dibromo- derivative [212°].

derivative $C_{24}H_{10}Me_6O_{11}$ Methyl [157°]. Formed from quercetin, H₆Me₄O₇. KOMe, and MeI (Herzig, M. 5, 83) and got also by heating xanthorhamnin with KMeSO, and MeOH at 120°, and from rhamnetin, KOH, and MeI (Liebermann a. Hörmann, A. 196, 317; Herzig, M. 6, 889; 9, 552). Golden needles, sl. sol. alcohol, converted by alcoholic potash at 140° into the di-methyl derivative of protocatechuic acid. Boiling Ac.O and NaOAc give C₂₄H₄Ac₂Me₄O₁₁ or C₁₈H₄AcMe₄O₇ crystallising in needles [167°].

Ethyl derivative C_{2.}H_{1,E}Et₆O₁, or C_{1.}H₂Et₄O₇, [122°]. Got by boiling quercetin with alcoholic potash and EtI (Herzig, M. 9, 587). Yellow needles, m. sol. alcohol. Yields

C_eH_s(OEt)₂CO₂H when heated with potash at 140°. Ac.O and sodium acetate give the colourless acetyl derivative C₂₄H₈Ac,Et₈O₁₁ or C₁₅H₁₅AcEt₄O₇ crystallising in needles [153°]. Mol. w. (by Raoult's method) c. 426 (Herzig, M. 12, 172).

A compound of quercetin and rhamnetin which sometimes occurs in Persian berries (Herzig, M. 10, 561) yields an acetyl derivative [171°], an acetyl-ethyl derivative [142°], and an ethyl derivative [102°].

Paradatiscetin C15H10O6. Formed from quercetin by potash-fusion (Hlasiwetz a. Pfaundler, J. pr. 94, 65). Yellowish needles (from dilute alcohol), nearly insol. water, m. sol. ether. Acid in reaction. FeCl, colours its alcoholic solution violet. Potash forms a yellow solution, turning green in air. Reduces AgNO, and Fehling's solution on heating. Potashfusion yields phloroglucin but not protocatechuic acid. Boiled with water and BaCO, it yields

BaA', 2aq.-SrA', 2aq.

Quercetic acid C15H10O, 3aq (?). Formed. together with phloroglucin, by heating quercetin (1 pt.) with moist KOH (3 pts.) till a sample no longer gives a flocculent pp. with HCl and the residue quickly turns dark red at the edges (Hlasiwetz, A. 112, 96; 119, 213; J. pr. 94, 65). Slender silky efflorescent needles, sl. sol. cold water, v. sol. alcohol and ether. Its aqueous solution turns yellow, and finally crimson, in air. H₂SO₄ forms a brown solution, whence water gives a red pp. forming a purple solution in NH, Aq. FeCl, gives a blue-black colour. Reduces AgNO. Potash-fusion gives protocatechuic acid. AcCl forms C15H8Ac2O2, crystallising from alcohol in prismatic needles.

Quercimeric acid C,H,O, aq (?). Formed Colourless from quercetin by potash-fusion. prisms, v. sol. water, alcohol, and ether. Tastes bitter. Alkalis colour its solutions purple-red. FeCl, gives a dark-blue colour. Reduces AgNO, and Fehling's solution. Potash-fusion gives

protocatechuic acid.

QUERCIN C_eH_e(OH)_e. [340°]. Occurs in oak bark, being obtained from the mother-liquors in the preparation of quercite (Gerber, A. 48, 348; Vincent a. Delachanal, C. R. 104, 1955. Frieddl. 105 of Physics 100. 1855; Friedel, 105, 95; Bl. [2] 48, 113). Efflorescent monoclinic prisms (containing x aq), sl. sol. water, insol. boiling alcohol. Inactive to light. Does not ferment with yeast, nor reduce Fehling's solution. Does not react with phenyl-hydrazine. Reduces ammoniacal AgNOs after addition of NaOH. Gives a gelatinous pp. with lead sub-acetate. After evaporation with HNO, it gives a rose-red colour with NH,Aq and CaCl, coloured by boiling NaOHAq.

Acetyl derivative.—C.H.(OAc). [301°].
QUERCITANNIC ACID C1.H1.C0, C₁₀H₁₆O₁₀. S. 6. S. (ether) 035. Occurs in oak bark, from which it is got by powdering, sifting from bast fibres, extracting with dilute alcohol, and shaking the filtrate with ether and EtOAc. The acetic ether when evaporated deposits ellagic acid. The filtrate is evaporated to dryness, and the residue washed with ether and then extracted with Et₂O and EtOAc (Etti, Sits. W. [2] 81, 495; M. 1, 264; 4, 514; of. Stenhouse, A. 45, 16; Böttinger, A. 202, 270; 240, 331; 263, 112; Lowe, Fr. 20, 210). Occurs

atso in tea (Rochleder, A. 63, 205). Reddishwhite powder. FeCl, colours the alcoholic solution dark blue. Pb(OAc)₂ ppts. the alcoholic solution. Quercitannic acid is not a glucoside (Etti). At 130° to 140° it forms an anhydride which gives C₃₄H₂₈BaO₁₇ (Etti). On evaporating with NaCl it is converted into a mixture of anhydrides which give with bromine-water an amorphous pp. C₁₉H₁₄Br₂O₁₀, m. sol. alcohol and EtOAc, v. e. sol. a mixture of these solvents, reacts with hydroxylamine, and is converted by Br into C₁₉H₁₀Br₄O₁₀, whence Ac₂O forms C...HAc.Br.G...

338, 1460; Hofmann, A. 190, 282).

Preparation. — Acorns are extracted with cold water, the extract concentrated at 40°, ppd. by lead subacetate, filtered, mixed with yeast to remove sugar, and, after fermentation, freed from lead by H₂S, and evaporated to crystallisation.

Properties. — Monoclinic prisms, $a:b:c = 800:1: 766, \beta = 68°57'$, insol. ether, benzene, and chloroform, v. sl. sol. hot alcohol. Does not undergo alcoholic fermentation. Dextrorotatory. Does not render borax solution acid.

Reactions.—1. At 100° it slowly loses water, forming C₂₄H₄₆O₁₉ (?). At 240° in vacuo it gives C₁₂H₂₂O₅ [230°], v. sl. sol. water. Rapidly heated to 290° it swells up and gives off quinhydron and hydroquinone.—2. Potash-fusion gives quinone, hydroquinone, CO₂, formic acid, and oxalic acid.—3. H₂SO₄ and MnO₂ give quinone.
4. Conc. HClAq forms, at 100°, C₆H,Cl(OH)₄ [200°], sol. ether, and viscid C₆H₅ClO₃, which is converted by baryta-water into amorphous quercitan C₆H₁₆O₄, sol. water and alcohol, insol. ether. HClAq at 120°-140° gives C₆H,Cl₂(OH)₂ [155°], converted by further treatment with HCl into C₆H₇Cl₅ [102°].—5. Distillation with HIAq forms benzene, phenol, iodo-phenol, quinone, hydroquinone, and hexane.—6. Nitric acid forms mucic and tri-oxy-glutaric acids (Kiliani a. Scheibler, B. 22, 517).—7. Phenyl cyanate at 165° forms C₆H₇(O.C.N.HPh)₅ [120°-140°], a white amorphous powder (Tesmer, B. 18, 2606).
8. Benzoic acid at 200° forms solid 'benzoquercite' C₅H₁₆Bz₂O₅ (?) insol. water, sol. alcohol and ether. Stearic acid forms a similar body, while tartaric acid gives 'quercitartaric acid C₂₂H₃₂O₂, (Berthelot, C. R. 44, 452; A. Ch. [3] 54, 82).

Salts.—(C₆H₁₁O₅)₂Ba 3aq: amorphous, sol. water and alcohol.—(C₆H₁₂O₅)₂CaSO₄ 2aq.

Acetyl derivatives. The compounds $C_8H_{11}AcO_5$, $C_8H_{10}Ac_2O_5$, $C_8H_8Ac_2O_5$, and $C_8H_7Ac_5O_5$ have been prepared.

Butyryl derivatives. Heating with butyric acid gives rise to $C_6H_{11}(C_4H_7O)O_5$, $C_8H_8(C_4H_7O)_2O_5$, and $C_8H_7(C_4H_7O)_5O_5$, all being amorphous with bitter taste.

Pentanitrate C.H. (NO.). Formed from queroite (1 pt.), H.SO. (10 pts.) and HNO.

(4 pts.) (Hofmann, A. 190, 288). Resin, insol. water, sol. alcohol and ether. Explodes when heated. Zinc-dust and alcoholic soda give off the N as NH₂.

QUERCITRIN $C_{16}H_{18}O_{20}$ or, more probably, $C_{21}H_{22}O_{12}$ (Herzig, M. 14, 53). [168°]. S. 04 in the cold; '7 at 100°. S. (alcohol) 4 in the cold 29 at 78° (Stein). S. (ether) '8 (Schunck, C. J 53, 264). Occurs in quercitron-bark, the bark of Quercus tinctoria (Bolley, A. 37, 101; 62, 136; Rochleder, J. pr. 77, 34; Rigaud, A. 90, 283; Zwenger a. Dronke, A. Suppl. 1, 266; Stein, J. pr. 85, 351; Hlasiwetz, A. 112, 109). Occurs also in fully-developed horse-chestnut leaves (Rochleder a. Kawalier, Sitz. W. 55 [2] 46), in leaves of the ash (Gintl, Z. [2] 4, 732), and in leaves of Andromeda japonica (Eykman, R. T. C. 2, 200).

Preparation.—Quercitron bark is exhausted with alcohol, and the evaporated extract dissolved in water and shaken with ether. The ethereal solution is evaporated, the residue dissolved in alcohol, and the quercitrin ppd. by water and crystallised from boiling water (Löwe,

Fr. 14, 233; cf. Herzig, M. 6, 877).

Properties.—Yellow needles or plates (containing 3aq). Neutral and tasteless. Nearly insol. cold water, sol. alkalis and HOAc. FeCl₂ colours its solution dark green. FeSO₄ gives no colour. Ppd. by Pb(OAc)₂. Reduces aqueous AgNO₃ in the cold, and Fehling's solution after long boiling. Boiling dilute acids split it up into quercetin (q. v.) and isodulcite (Liebermann a. Hamburger, B. 12, 1178). Bromine forms C₃₆H₂₄Br₄O₂₀, which is crystalline and is decomposed by acids into isodulcite and tetra-bromoquercetin.

Salt.— $C_{36}H_{36}K_2O_{20}$: yellow pp. •

Violaquereitrin C₂₂H₄₂O₂₄. Occurs in Viola tricolor (Mandelin, J. 1883, 1369). Yellow needles (from water). Split up by dilute acids

into glucose and quercetin.

QUILLAJIC ACID C₁₉H₂₀O₁₀. Extracted by water from the bark of Quillaja Saponaria (Kobert, C. C. 1888, 972). White flakes, solwater and alcohol, insol. ether. Coloured dark red by H₂SO₄. Boiling dilute acids split it up into sapoginin and an unfermentable glucose. The Na salt violently attacks the mucous membrane, and is very poisonous when injected into the blood.

QUINACETOPHENONE is DI-OXY-ACETO-PHENONE.

QUINALDINE v. METHYL-QUINOLINE.
QUINALDINIC ACID v. QUINOLINE CARBOXYLIC ACID.

QUINAMINE v. vol. ii. p. 179.

QUINAMICINE v. vol. ii. p. 180.

QUINAMIDINE v. CINCHONA BASES.

QUINANISOLE v. Methyl derivative of Oxymoline.

QUINAZOLE v. METHYL-INDAZINE.

QUINAZOLINE. This name is given to the ring $C_eH_e < \begin{array}{c} CH:N\\N=CH \end{array}$; v. Oxy-quinazoline.

QUINAZOLINE DIHYDRIDE $C_sH_sN_2$ i.e. $C_sH_s \sim C_H.NH$ [127°]. Formed by reducing o-nitro-benzyl-formamide by Zn and HClAq (Gabriel a. Jansen, B. 23, 2814; 24, 3097). Yellowish crystals, sol. warm water, forming an

alkaline solution.—B'HCl.—B'C₆H₂N₂O₇. [215°]. Small crystals, sl. sol. water. -B'2H2PtCl3.

QUINENE v. vol. ii, p. 181.

QUINETHONIC ACID C₁,H₁₈O₈ i.e. C.H₄(OEt).C₆H₄O₇. [146°]. Occurs in urine after a dose of C₆H₃.OEt (Kossel, H. 4, 296; 7, 292; Lehmann, H. 13, 181). Crystalline. Lævorotatory. Does not reduce Fehling's solution. Converted by dilute H2SO, into crystalline C₈H₁₀O₂. HIAq forms hydroquinone. Oxidising agents yield quinone. — KA'aq: monoclinic crystals.—AgA'aq. Forms with C₄H₃.O.SO₂H the double salt PhO.SO₃BaA'aq (dried at 110°) and with cresol and indoxyl the corresponding C,H,O.SO,BaA'aq and C,H,NSO,BaA', all three being crystalline (Hoppe-Seyler, H. 7, 424). QUINHYDRONE $C_{12}H_{10}O_4$. Formed by mix-

ing aqueous solutions of quinone and hydroquinone; by oxidising hydroquinone; and by reducing quinone (Wöhler, A. 51, 153; Liebermann, B. 10, 1614, 2000; Hesse, A. 200, 248; Nietzki, A. 215, 130; Wichelhaus, B. 5, 840; 12, 1500; Stenhouse a. Groves, B. 13, 1305). Brownishred prisms, with green metallic lustre. May be sublimed. Sl. sol. cold water, v. sol. alcohol and ether, forming yellow solutions. Decomposed by boiling water, giving off quinone and leaving a solution of hydroquinone. Its ammoniacal

solution is green. Reduces ammoniacal AgNO₃.

QUINIC ACID C,H₁₂O₆ i.e. C₆H₁(OH)₄CO₂H.

Mol. w. 192. [162° cor.]. S.G. 8.5 1.637 (Henry a.

Plisson, B. J. 10, 186). S. 40 at 9°. [a]_D = -44° at 20° in a 20 p.c. solution (Thomsen, J. pr. [2] 35, 156). H.C. 833,700. H.F. 238,300 (Berthelot a. Recoura, C. R. 105, 144; Bl. [2] 48, 703; A. Ch. [6] 13, 342). R_{∞} 66.52 in an 18 p.c. aqueous solution (Kanonnikoff, J. pr. [2] 31, 348). Occurs as calcium salt in cinchona bark (Hoffmann, Crell's Ann. 2, 314; Vauquelin, A. Ch. 59, 162; Pelletier a. Caventou, A. Ch. [2] 15, 340; Liebig, P. 21, 1; 29, 70; Baup, A. Ch. [2] 51, 5; A. 6, 7; Woskresensky, A. 27, 260; Hesse, A. 110, 194; 112, 52; 114, 292; 176, 124; Clemm a. Will, A. 110, 345). Occurs also in the bilberry plant (Vaccinium Myrtillus) (Zwenger, A. 115, 108; 129, 203; Suppl. 1, 77), in the leaves and beans of the coffee-plant (Z.), and in hay (O. Loew, J. pr. [2] 19, 310; 20, 476).

Preparation.—Cinchona bark is extracted

with dilute H2SO4, the extract ppd. by milk of lime, filtered, and evaporated. The residue is boiled with alcohol and the calcium quinate left undissolved is crystallised from water and de-

composed by oxalic acid.

Properties .- Monoclinic prisms. V. e. sol. water, m. sol. alcohol, nearly insol. ether.

Lævorotatory.

Reactions .- 1. Heated to 200°-225° it gives off aq, forming quinide C,H10O5, a crystalline anhydride, v. sol. water, sl. sol. dilute alcohol. Quinide is acid in reaction, and is reconverted by bases into quinic acid. On dry distillation quinic acid gives hydroquinone, phenol, benzoic acid, and pyrocatechin.—2. Ac₂O at 170° forms tri-acetyl-quinide C.H. (OAc), CO [132°] and tetra-acetyl-quinic acid (Erwig a. Königs, B. 22, 1458; cf. Hesse, A. 200, 233). By further heating with Ac.O at 240°-250° monoclinic

orystals of iso-tri-acetyl-quinide [139°] are obtained.—3. Bromine added to an aqueous solution forms protocatechuic acid (Hesse).-Hot H₂SO₄ forms CO and hydroquinone disulphonic acid .- 5. Hydroquinone is formed by boiling the aqueous solution with PbO2. Distillation with MnO₂ and dilute H₂SO₄ yields quinone (detection of quinic acid in bark: Stenhouse, A. 59, 100).—6. HNO, gives oxalic acid.
7. KClO, and HCl give chlorinated quinones and chlorinated acetones (Städeler, A. 69, 300; 111, 293).—8. Conc. HIAq at 120° reduces it to benzoic acid (Lautemann, A. 125, 9).—9. Conc. HClAq at 150° gives p-oxy-benzoic acid and hydroquinone (Hesse, A. 200, 232). Dilute (3 p.c.) HClAq at 100°-120° forms phenol, hydroquinone, and p-oxy-benzoic acid (Chadounski, C. C. 1888, 1029). - 10. Fuming HBrAq at 130° gives protocatechuic and benzoic acids (Fittig, A. 193, 197).—11. PCl, forms m-chloro-benzoyl chloride (Graebe, A. 138, 197). 12. Gives protocatechuic acid on fusion with KOH or NaOH.-13. Boiling with iodine and KOHAq yields iodoform.—14. Calcium quinate fermented by schizomycetes in presence of air yields protocatechuic acid, in absence of air it gives propionic, acetic, and formic acids (Löw, B. 14, 450).

Salts .- NaA' 2aq .- NaA' 5aq. S. 200.— BaA', 6aq: dodecahedra, very soluble in water.-CaA', 10aq. S. 17 at 16°. Plates, insol. alcohol. —Cah'Ac aq (Gundelach, B.9,852).—SrA', 10aq. —SrA', 15aq.—MgA', 6aq.—CdA', S. 4 in the cold.—ZnA', .—CoA', 5aq.—NiA', 5aq: crystals. CuA', 5aq: blue needles.—CuC, $H_{10}O_6$ 2aq. S. 09 at 18°.—PbA', 2aq: needles, sol. alcohol, v. e. sol. water.—Pb₂C₂H₈O₆. Amorphous, insol. water.—MnA'₂. S. 5 in the cold.—FeC₁H₂O₁, —AgA': mammellated groups of crystals.

Ethyl ether EtA'. Viscid mass, with

bitter taste, v. sol. water and alcohol. Converted by boiling Ac₂O into C₆H₇(OAc)₄.CO₂Et crystallising from water in plates [135°], sl. sol. boiling water.

Tetra-acetyl derivative $C_6H_7(OAc)_4.CO_2H.$ [130°-136°]. Formed by heating quinic acid with Ac, O and ZnCl, (Erwig a. Koenigs, B. 22, 1461). Crystalline crusts, sl. sol. cold water, insol. ligroïn.—AgA': needles.

Anilide C₁₈H₁₇NO₅. [174°]. Formed by

Anilide C₁₈H₁₇NO₅. [174°]. Formed by heating quinic acid with aniline at 180°. Small silky needles (containing aq), v. sol. water and alcohol, sl. sol. ether.

QUINICINE v. CINCHONA BASES. QUINIDE v. QUINIC ACID. QUINIDINE v. vol. ii. p. 180. QUINIENE v. vol. ii. p. 181.

QUININE $C_{20}H_{24}N_2O_2$: [173° cor.] (Lenz, Fr. 27, 559; Hesse, A. 258, 133). The trihydrate melts at 57° (H.). S. 05 at 15° (Regnauld, J. Pharm. Chim. [4] 21, 8). S. (of the trihydrate) ·06 at 15° (Hesse, B. 10, 2152); ·06 20°. 11 at 100° (Sestini, Fr. 6, 359). S. (of the anhydrous base) '051 at 15° (H.); '07 at 20°, '13 at 100° (S.). S. (ether) 100 at 10° (Hesse, A. 135, 327); 4.4 at 18° (van der Burg, J. 1865, 438). A conc. ethereal solution often gelatinises, the quinine thus separated being less sol. ether (S. about 5 at 15°). S. (chloroform) 57.5 (Pettenkofer, J. 1858, 363). S. (benzene) 5 at 15° ; 3.8 at 80° (Oudemans, J. 1874, 867). S. (xylene) 11 at 15° ; 645 at 188° (Swaving, R. T. C. 4, 186). [a]_D = -166° at 15° ; -162° QUININE. 875

at 25° in alcoholic solution (Hesse, A. 166, 217) = $-145^{\circ}2 + 657p$ in a solution of p grammes in 100 c.c. of 97p.c. alcohol at 15° (Hesse, A. 176, 206; 182, 131). $[a]_{\rm b} = -158^{\circ}7 + 1^{\circ}911p$ in an ethereal solution containing from 1.5 to 6 p.c. base. In solutions containing not more than 1.6 p.c. quinine, Oudmans, jun. (Ar. N. 10, 193), found $[a]_{\rm b} = -167^{\circ}5^{\circ}$ (in alcohol); -136° in benzene; -127° in toluene; -117° in chloroform. Dispersive power: Grimbert, J. Ph. [5] 16, 295, 345.

Occurrence.—In cinchona bark, v. vol. ii. p. 175. According to Grimaux (Bl. [3] 7, 304), cupreïne heated with NaOMe, MeOH, and McOl or MeNO₂ yields 10 to 15 p.c. of quinine, while MeI gives chiefly quinine di-methylo-di-iodide.

Preparation.—The bark is extracted with dilute H₂SO₄, and the solution ppd. by NaOHAq. The pp. is dissolved in ether, the ether shaken with dilute H₂SO₄, and the boiling solution neutralised by ammonia. Quinine sulphate separates on cooling. The sulphate is decom-

posed by ammonia.

References. — Vide references in articles CINCHONA BARK and CINCHONIDINE, and also Pasteur, C. R. 36, 26; 37, 110, 162; Schützenberger, A. 108, 347, 350; Robiquet, A. Ch. [2] 17, 316; Stratingh, R. P. 15, 139; Pelletier, J. Ph. 11, 249; Duflos, B. J. 27, 1, 110; Strecker, A. 91, 155; Thiboumery, J. Ph. [3] 16, 369; Alluard, J. Ph. [3] 46, 192; Körner, Z. [2] 1, 150; Bouchardat, A. Ch. [3] 9, 213; De Vrij, N. J. P. 14, 268; Laudrin, C. R. 108, 750.

Properties.-Ppd. by ammonia from solutions of its salts in an amorphous anhydrous form, which quickly changes, especially in presence of free ammonia, into the crystalline hydrate (containing 3aq), consisting of minute four-sided prisms terminated by pyramids. The hydrate gives off its water of crystallisation over \dot{H}_2SO_4 . Quinine is v.e. sol. ether and alcohol, v. sol. \dot{CS}_{21} m. sol. benzene, v. sl. sol. ligroïn, sol. volatile and fixed oils. It is alkaline Its solutions are lævorotatory. in reaction. A 5 p.c. solution of the sulphate gives $\alpha = -22^{\circ}$ in a depth of 200 mm. (Rozsnyay, Fr. 23, 589). Its solutions in dilute H₂SO₄, HNO₃, H₂PO₄, and HOAc exhibit blue fluorescence, destroyed by HCl, HBr, HI, or H, FeCy. Fluorescence of the sulphate is prevented by a large excess of sulphate of cupreïne (Grimaux a. Arnaud, Bl. [3] 7, 304). Chlorine-water and ammonia give a green pp. dissolving in excess of ammonia to an emerald green liquid. On addition of an acid the colour changes through blue (when neutral) to red, the green colour returning on adding ammonia (Brandes, Ar. Ph. 13, 65; André, J. Ph. 22, 132). Excess of 05; André, J. Ph. 22, 132). Excess of chlorine-water and of ammonia should be avoided. Chlorine does not render a solution of quinine sulphate turbid (Lepage, J. Ph. 26, 140). Bromine-water and ammonia give the green colour even in exceedingly dilute solutions (Flückiger, Fr. 11, 318). The green colour may be also got by mixing 01 g. of the salt to be tested with an equal bulk of KClO, and a drop of conc. H₂SO, followed by excess of ammonia (Mylius, *C. C.* 1886, 602). Chlorine-water followed by K₄FeCy_e gives a red colour in a solution of quinine sulphate (Vogel, A. 78, 221; 86, 122). Quinine gives the usual alkaloidal

reactions. It is coloured green by potashfusion (Lenz, Fr. 25, 31). Potassium sulphocyanide gives a white pp., sol. excess (Schrage, Ar. Ph. [3] 13, 25). Boiled with dilute H_sSO_s and PhO_s quinine forms quinetin, a redsubstance (Marchand). Dilute HNO, gives no colour. Sunlight acting on an aqueous solution, even in an atmosphere of H, ppts. brown floculent quiniretin, insol. water, alcohol, and ether (Flückiger, Ph. [3] 8, 885). ICl gives a light-brown crystalline pp., sl. sol. HClAq. Quinine is antiseptic, hindering putrefaction and the alcoholic, lactic, and butyric fermentation. Quinine is a febri-

fuge. Its salts taste bitter.

Reactions.-1. The sulphate is oxidised by KMnO, to pyridine tricarboxylic acid, oxalic acid, and NH, (Hoogewerff a. van Dorp, B. 12, 158). At 0° the first product is chitenine C₁₈H₂₂N₂O₄, which crystallises in colourless prisms [240°–286°], insol. ether and alcohol, sol. dilute acids and alkalis (Skraup, B. 12, 1104; M. 10, 39). Quinine (5 g.) is apparently oxidised by KMnO, (50 g.) to pyridine dicarboxylic acid (Ramsay a. Dobbie, C. J. 33, 102). Boiling nitric acid also forms pyridine dicarboxylic (cinchomeronic) acid. Aqueous CrO, oxidises quinine to quininic or methoxyquinoline carboxylic acid.—2. A mixture of HNO₃ and H₂SO₄ reacts, and on diluting an amorphous pp., apparently C₂₀H₂₂(NO₂)₂N₂O₂H₂O is got (Rennie, C. J. 39, 469).—3. PCl₃ converts it into colourless quinine-chloride C20H23N2OCl [151°] which by boiling with alcoholic KOH gives chinen $C_{20}H_{22}N_2O$ (Comstock a. Königs, B. 17, 1988).—4. Reduced by zinc and dilute H_2SO_4 to hydroquinine $C_{20}H_{26}N_2O_2$, an amorphous bitter resin (containing aq), sol. alcohol and ether, giving a green colour with chlorine-water and ammonia (Schützenberger, A. 108, 347).-5. Heated with water at 250° it yields quinoline (Reynoso, C. R. 24, 795). The same body is formed by heating with conc. KOHAq at 190° (Gerhardt a. Wertheim, J. 1840, 370).—6. Fuming H₂SO, forms a sulphonic acid. Conc. H₂SO₄ dissolves quinine, forming isocinchonine. Dilute H₂SO₄ at 125° converts quinine into quinicine.—7. *Iodic acid* gives off much gas on warming (Brett, J. Ph. [3] 27, 116).—8. Conc. HClAq at 150° forms McCl and apoquinine (Hesse, A. 205, 317). Concentrated HClAq at -17° slowly forms hydrochloroquinine C₂₀H₂₅ClN₂O₂[187°], while hydrogen bromide gives C₂₀H₂BrN₂O₂, which forms B'H₂Br₂ (Comstock a. Königs, B. 20, 2510).—9. Heated for a long time with 12 pts. of HIAq. S.G. 1.96 for 3 hours at 100° it gives methyl iodide and a yellow salt $C_{19}H_{22}N_2O_23HI$ [238°] sol. alkalis. Alcoholic NH₂Aq converts it into C₁H₂N₂O₂HI mixed with a little C₁H₂N₂O₂HI, which yields an oxalate [187] (C₁H₂N₂O₂HI)H₂C₂O₄ mixed with (C₁H₂N₂O₄H₂I₂J₂H₂O₄ (Schubert a. Skraup, M. 12, 684). HBrAq acting on quinine at 100° C₁₀H₂₇BrN₂O₂HBr aq, crystallising in forms needles, from which sodium carbonate sets free C₁₈H₂₇BrN₂O₂, a powder [210°] (Julius, M. 6, 751). 10. Quinine dried at 120°, heated with HIAq (S.G. 1.7) at 100°, forms C₂₀H₂₁N₂O₂3HI [230°], a yellow crystalline mass, v. sl. sol. water, sl. sol. alcohol, insol. ether and NaOHAq. Alcoholic NH_2 converts this salt into $C_{20}H_2$, N_2O_2HI ? which is white, v. sol. alcohol, sl. sol. ether [155°-

160°]. Its solution in dilute H₂SO, shows blue fluorescence, and gives a brownish-yellow pp. with chlorine-water and ammonia. It gives rise to the salts ((C₂₀H₂₄N₂O₂)₂H₃I₃₂BH₂C₂O₄? and C₂₀H₂₄N₂O₂H₂I₄HNO₂? [217°] (Sohubert a. Skraup, M. 12, 678; cf. Lippmann a. Fleissner, M. 12, 329).—11. Bromine added to a solution of quinine in dilute H₂SO₄ ppts. B'Br₂ 2aq, B'₂Br₆, and B'Br, as bitter yellow curdy pps., melting at 160°-180° (Colson, C. R. 108, 678).

Estimation.—V. CINCHONA BARK. V. also Lenz, Fr. 27, 549-631. Quinine may be ppd. as chromate (De Vrij, C. C. 1889, 708; Ar. Ph. [3] 24, 1073; Vulpius, Ar. Ph. [3] 24, 1022; Schlikum, Ar. Ph. [3] 25, 128; Hesse, Ph. [3] 17, 585). A solution of quinine sulphate requires more ammonia to redissolve the pp. first formed than solutions of the sulphates of the alkaloids usually present with it (Kerner a. Weller, Ar. Ph. [8] 25, 712, 749; Fr. 1, 159; 27, 115; cf. Schäfer, Ar. Ph. [3] 25, 1033; Ruddiman, C. N. 58, 202, 216, 226; Jungfleisch, J. Ph. [5] 15, 5). Use may also be made of the sparing solubility of quinine oxalate in presence of K₂C₂O₄ (Schäfer, Ar. Ph. [3] 25, 64,1041).

Constitution.—Chitenine C19H22N2O4 may be got by oxidation of both quinine and cinchonine. On further oxidation chitenine yields quinic, pyridine tricarboxylic, and cincholeuponic acids. From this it may be surmised that one half of the quinine molecule has the same composition as one half of the cinchonine molecule (Skraup, M. 10, 39, 220). Quinine, quinidine, and quinicine are probably stereo-isomeric, since they give the same products of oxidation; they are derivatives of (B. 2)-methoxy-quinoline while cinchonine is a derivative of quinoline. Cincholeuponic acid C₈H₁₈NO₄ [226°] yields C₈H₁₈NO₄HCl [194°] and C₈H₁₂AcNO₄, and is converted by nitrous acid into the dibasic nitroso- derivative $C_sH_{12}(NO)NO_4$. Oxidation of quinine also yields a base, cincholeupone $C_sH_{17}NO_2$ oxidised by CrO_3 to cincholeuponic acid and other products. Cincholeupone when heated with zinc-dust yields (\$\beta\$)-ethyl-pyridine. By nitrous acid cincholeupone is converted into C₂H₁₆(NO)NO₂, which is an acid. Ac₂O forms acid $\dot{O}_0H_{1s}\Delta c\dot{N}O_2$. Oxy-quinoline (cynurine) is also a product of oxidation of quinine. From these experiments Skraup concludes that quinine and cinchonine contain a quinoline nucleus united to an ethyl-pyridine nucleus; quinine being methoxy-cinchonine. Cupreïne heated with NaOMe (1 mol.), MeI (6 mols.), and MeOH forms quinine di-methylo-di-iodide (Hesse, A. 266, 244; cf. Grimaux a. Arnaud, C. R. 112, 774; A. 267, 879). The chief product is, however, cupreïne mono-methylo-iodide. \mathbf{W} hen only 1 mol. MeI is used, no quinine is got.

Salts.—C. H₂₂AgN₂O₂. Gelatinous pp. got by adding AgNO, to a solution of quinine in alcoholic NH₂ (Skraup, M. 2, 613).—B'HCl 2aq: groups of white needles (Hesse, A. 176, 210; 267, 142). Melts (when anhydrous) at 160°, without undergoing any change. S. 2.5 at 10°. V. e. sol. alcohol and ether. $[a]_D = -134^\circ$ in a 2 p.c. solution at 17° (Oudemans). Its solution is not ppd. by silver nitrate (1 mol.) until added in excess (Vulpius, Ar. Ph. [3] 20, 861).-

or gelatinous mass. S. 100 in the cold. Melts at 15° (A. Clermont, J. Ph. [5] 15, 15).— B'H. PtCl, aq: yellowish flocculent pp., soon becoming orange and crystalline (Gerhardt, B. J. 23, 354). S. 07 in the cold; 8 at 100° (Duflos). -B'2H2PtCl, 3aq : orange amorphous pp. (Hesse, A. 207, 308).—B'H2HgCl4. Ppd. by mixing alcoholic solutions of quinine, HCl, and HgCl, (Hinterberger, A. 77, 201).—B'2H,ZnCl, 2aq: prisms (from alcohol) (Gräfinghoff, Bl. [2] 4,391). - B'₂H₆ZnCl₈ 3aq: crystalline. - B'H₂Br₂ 3aq: crystals, v. e. sol. water. - B'HI: lemon-yellow prisms (Herapath), or heavy white powder. Sl. sol. water, v. e. sol. alcohol (Winckler, Jahrb. pr. Pharm. 20, 321). Melts at 150°-155°, but softens at about 100° and then blackens (Lippmann a. Fleissner, M. 13, 436).—B'HIEt₂O (from other). Crystals.—B'H₂I₂5aq (Regnault). Crystals. V. Reaction 9, supra.—B'H.I., [230°]. Converted by NH,Aq into BHI.—B',H.SO, 8aq (Hesse, A. 119, 361; 225, 97; Carles, Bl. [3] 7, 108; cf. Cownley, Ph. [3] 7, 189). Commercial specimens are somewhat effloresced. Monoclinia efflorescent prisms. Loses 6aq over H2SO4. S. (of B'.H.SO.) ·126 at 6°: 3 at 100° (Howard). S. (alcohol of S.G. 85) 1. S. (glycerin) 2.5. Insol. chloroform and fatty oils. $[a]_D = -163^\circ$ in a 2 p.c. solution in 80 p.c. alcohol.—B'H2SO47aq. [100°]. Rectangular prisms (from hot solutions) or small needles. S. 9 at 13°; 12.5 at 22°. Loses 6aq over H_2SO_4 . $[\alpha]_b = -164.9 + .31p$ in a solution of p g. in 100 c.c. water, where p is between 1 and 6 (Hesse, A. 176, 215; 182, 134).—B'(H₂SO₄)₂ 7aq: prisms, v. e. sol. water, m. sol. alcohol. Its hot alcoholic solution deposits gelatinous B'(H_2SO_4)₂5aq. [a]₅ = -170 + 94p.— B'₂H₂Cl₂H₂SO₄3aq. Mass of small needles, v. sol. water. S. (of anhydrous salt) 86. [120°] (hydrated); [165°-170°] (anhydrous) (Grimaux, Bl. [3] 7, 819).—B'₂H₂Br₂H₂SO₄3aq. S. 26 at 21°.-B'2H2L2H2SO, 2aq: yellow crystals, forming a colourless solution. Crystallises also with Mg 2 Colories Studies Crystallies also with Aq. ... B'₂H₂Cl₂H₃PO₄9aq: small needles, sol. water. - B'₂H₂E₁H₂PO₄ 9aq: small needles, sol. water. - B'₂H₂I₂H₂PO₄ 6aq. -B'₂H₂I₃S₂O₃2aq. S. ·3 (Wetherill, A. 66, 150; How, N. Ed. P. J. [2] 1, 47).—B'HNO₃aq: prisms (Strecker, A. 91, 159).—B'AgNO₃aq: crystalline pp. Chlorate.—B'HClO, 2aq (Tichborne, Z. 1866, 665; cf. Serullas, A. Ch. [2] 45, 279).—B'(HClO₄)₂ 7aq: trimetric octahedra.— B'(HClO₄) 2aq. [210°] (Boedeker, A. 71, 61).— B'HIO, 9aq: needles (Langlois, A. Ch. [3] 31, B'HIO, vaq: needles (Liangiois, A. Ch. [2] C. 274).—B'₄(HCl)₃(HI)₄I₁: small brown crystals.—B'₃(HCl)₅(HI), I₁₀: blackish-green flat needles (Jörgensen, J. pr. [2] 15, 79).—B'HI₂ (Bauer, Ar. Ph. [3] 5, 214).—B'HI₃. Black prisms.—B'₄I₅.—B'₄(H₂SO₄)₅(HI)₂I₄ 6aq. Herapathite. Formed by adding an alcoholic solution of iodine to a solution of quinine sulphate in HOAc (Herapath, P. M. [4] 3, 161; 4, 186; 6, 171, 846; 7, 352; 9, 366; 14, 224; C. J. 11, 130; Haidinger a. Stokes, Sitz. W. 10, 106; Jörgensen, J. pr. [2] 14, 230). Large colourless plates with metallic green lustre. Polarises light, so that two plates at right angles are opaque. S. (90 p.c. alcohol) 125 at 16. Decomposed by cold water. Dried over H2SO, they become B',(H,SO,),I, 3sq (Hauer, Z. [2] 1, 481).

B',(H,SO,),(HI),I,: thin lustrous plates.

B',(H,SO,),(HI),I, 2sq. Crystals, resembling B'HCll 1 aq. Large monoclinic octahedra, deposited at 0°.—B'2HCl. Groups of white needles, herapathite, decomposed by hot alcohol into

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iodine and herapathite.—B's(H2SO4)e(HI)4I144aq: brownish needles.—B'2H2SO4(HI)2I2: red needles. -B'2H2SO4(HI)2I4: brown rectangular plates with olive-green reflex.— $B'_2H_2SO_4$ _2(HI)_2 I_3 . black crystals with green reflex.— $B'_3(H_2SO_4)_2(HI)_2I_3$.— $B'_3(H_2SO_4)_2(HI)_2I_3$. aq. — $B'_3(H_2SO_4)_2(HI)_2I_3$.— B'H, SeO, 7aq: trimetric tables, insol. alcohol (Hjortdahl, J. 1879, 794). $-B'_4(H_2SeO_4)_8(HI)_2I_4$. Isomorphous with herapathite, which it greatly resembles. Almost insol. cold alcohol (Jörgensen, J. pr. [2] 15, 65, 418).—B'₂H₂CrO₄. S. ·042 at 15° (André); ·037 at 14°; ·05 at 16° (De Vrij, Ar. Ph. [3] 24, 1073; ·625 at 100° (A.). Ppd. by adding K₂CrO₄ to a solution of the neutral or acid sulphate. Tufts of golden needles.—B'2H2CrO, 2aq (Hesse, Ph. [3] 17, 585, 665). Becomes anhydrous at 80°, but re-absorbs 2aq on exposure to moist air.—B'H, CrO, 7aq. Ppd. by adding K2Cr2O, to a solution of quinine in excess of dilute H2SO4 (André, J. Ph. [3] 41, 341). Orange pp., decomposed by light, and turned brown when heated to 60°-65° or when boiled with water. More soluble than B'2H2CrO4. -B'H₂CO₈ aq. Efflorescent needles, with al-kaline reaction, deposited from a solution of quinine in aqueous CO₂. Sol. alcohol, insol. ether (Langlois, A. Ch. [3] 41, 89).—B'₂H₃PO₄8aq. S. 127 at 10° (Hesse). Tufts of long needles.— B',2H,PO,5aq (Anderson, A. 66,59). Crystals.— B',2H,PO,12aq (A.).—B',2H,PO,2aq (Gerhardt, Gerh. iv. 118).—B'H,PO, 8. 1-33 at 15.5° (L. Smith, Z. 1862, 159).—B'HVO, (Ditte, A. Ch. [6] 13, 236).—B'H2SiF6. Got by passing SiF4 into a solution of quinine in absolute alcohol (Cavazzi, G. 17, 563). Minute crystals, insol. sther and CS₂, sl. sol. hot alcohol. Its aqueous solution is fluorescent.—B'2H3AsO48aq. Prisms, v. sol. hot water (Hesse).—B'2H3AsO46aq (Sestini).—B'H₄ASO, 2aq.—B'H₄FeCy₈ 3aq (Dollfus, A. 65, 227).—B'H₅FeCy₈ 1₃aq: golden plates.—B'H₂PcCy₄aq (Wertheim, A. 73, 210). Crystals.— B' H.PtCy, 2aq. S. .054 at 18°; 1.8 at 57°. S. (alcohol) 20 at 16° (Schwarzenbach, Pharm. Viertelj. 8, 210; Van der Burg, Fr. 4, 312).— B'H.PtCy, 2aq. 074 at 18.5°; 5 at 100°.— B'H.PtCy, —B'2HCyS. Lemon-yellow monoclinic crystals (W.).—B'HCyS aq. S. 18 at 20° (Hesse, A. 181, 48).—B'₃(HCyS)₄HgCl₂.—B'₂(HCyS)₂HgCy₂.—B'(HCr(CyS)₄(NH₃)₂)₂2aq. Formed by ppg. a solution of the acid sulphate with Reinecke's salt (Christensen, J. pr. [2] At. 16 Hencek's sait (Christensen, J. pr. [2] 45, 366). Red crystals, v. sl. sql. hot water.

Nitro-prusside. S. '04 (Davy, Ph. [3] 11, 756).—Cyanurates B'H₃C₂N₃O₃ 9aq. [237°]. Sl. sol. hot water (Claus, J. pr. [2] 38, 227).—B'(H₃C₃N₃O₃)₂ 7aq. [248°].—Oxalates.—B'₂H₃C₂O₄6aq. S. 1 at 10°. Prisms (Hesse, A. 176 213; et Parayut 4, 26, 27). B'H O co. 176, 218; cf. Regnault, A. 26, 37).—B'H₂C₂O₄ aq. Prisms, m. sol. cold water. — B'H₂C₂O₄ 3q. [α] = -131°.—B'HOAc. [140°]. Long needles, v. sol. hot water (R.).—B'Cu(OAc)₂. Green crystals (Skraup, M. 2, 611).—Chloro-acetate B'C₂H₂ClO₂ 2 aq. S. 1.6 at 21° (Mazzara, G. 13, 525).—Di-chloro-acetate B'C₂H₂Cl₂O₂ 2aq. S. 24 at 22°.—Formate: colourless needles (Bonaparte, J. Chim. Méd. 18, 680).-Valerate B'C,H₁₀O₂1½aq (Bonaparte, *J. Chim. Méd.* 18, 680; 19, 330; Chatin, *J. Ph.* [4] 1, 268).—
B'C,H₁₀O₂ (Stalmann, *A.* 147, 132; Schmidt a. Sachtleben, *A.* 193, 100). S. 9 in the cold; 2.5 at 100° (Wittstein, Repert. 87, 295; Landerer,

N. Br. Arch. 119, 240). — Succinate B'C,H,O, 8aq. S. 1 at 10°. Prisms, v. sol. hot water.—Tartrate B'₂C,H,O, 2aq. Crystalline powder (Hesse, A. 243, 134; cf. Arppo, J. pr. 53, 334).—B'C,H,O, aq (Pasteur, J. 1853, 421). The levo-tartrate is more soluble than the dextro- tartrate, and has a different crystalline form.—B'C,H₄(SbOH)O₅ 2aq. Resin (Clarke, B. 15, 1540).—Citrate B'₂C₆H₈O₇ 7aq. S.·11 at 12° (Hesse); 44 at 100° (Mandelin, J. 1879, 796; Scribani, G. 9, 284).—B'₂C₆H₈O₇: minute prisms. S. 11 in the cold; 2.4 at 100°.— B'C₆H₈O₇. Small prisms. S. 16 in the cold, 2.6 at 100°.—Chloro-crotonate B'C₄H₈ClO₈. [201°]. Sl. sol. ether (Daccomo, J. 1884, 1885). -Trichlorocrotonate B'C3H3Cl3O3. Sl. sol.
-Tri-chloro-lactate B'C3H3Cl3O3. Sl. sol.
-Tri-chloro-nyruvate B'C3H2Br2O3. ether.—Di-bromo-pyruvate $B'C_3H_2Br_2O_3$. [93°] (D.). — Mucate $B'C_4H_{10}O_3$. Needles (Ruhemanna. Dufton, C.J.59,754).—Benzoate B'HOBz. S. 27 at 10°. Small prisms.—Salicylate B'C,H₀O₃. S. 44 at 16°. S. (ether) 83 at 16°. Prisms (from alcohol). — Melitate $B'_2C_{12}H_uO_{12}$. Crystalline powder, v. sl. sol. cold water (Karmrodt, A. 81, 170). — Tannates $B'(C_{11}H_{10}O_{9})_2$ 4aq and $B'(C_{14}H_{10}O_{9})_3$ 8aq (Jobst, Ar. Ph. [3] 12, 331; Neumann, Fr. 28, 664).-Meconate B'C,H₄O₇. Crystals (Austen, Ph. [3] 3,1016).—Urate B'C₅H₄N₂O₅. Minute prisms. S. 117 in the cold, 2.7 at 100°. S. (alcohol of S.G. ·823) ·063 in the cold, 2·2 on boiling (Elderhorst, A. 74, 77; Andrew, Pharm. Viertelj. 10, 382). Dextro-tropate. [187°].—Lævo-tropate. [178°] (Ladenburg, B. 22, 2590). Dextro-iso-propyl-phenyl-glycollate [193°]. S. 18 at 19°; S. (alcohol) 1.44 at 20°. [α]_D = -79°. — Lævo-iso-propyl-phenylglycollate [205°]. S. 09 at 15°; S. (alcohol) 54. $[\alpha]_D = -118^\circ$ (Fileti, J. pr. [2] 46, 560).

Compound with benzene B'C.H... Needles (from benzene) (Oudemans, J. 1874, 867).—Compound with toluene B'C,H8. Needles. — Compounds with phenol B'PhOH. Crystals (from alcohol). S. 25 at 16° (Romei, Z. [2] 5, 383; Jobst, N. R. P. 24, 193). -B'_H_Cl_PhOH 2aq: prisms. S. 1 at 15° (Jobst a. Hesse, A. 180, 248).—B'H_SO,PhOH 2aq (Hesse, Pharm. Zeit. 34, 191).—B'_2H_SO,PhOH aq. S. 147 at 15°. Prisms.—B'_2H_SO,PhOH 6aq (Cotton, S. 147 at 15°. Prisms.—B'_2H_SO,PhOH 6aq (Cotton, S. 15°). Bl. [2] 24, 535) .- Compound with tribromo-phenol B'C₆H₂Br₂O. Silky needles (from alcohol) (Purgotti, G. 16, 528).—Compound with anethole B'₂C₁₆H₁₂O 2aq. Monoclinic crystals (from ether), sl. sol. cold alcohol (Hesse, A. 123, 382). Compound with eugenol B'C₁₀H₁₂O₂. [110°]. S. (ether) 8.5 at 10°. Long silky prisms (Hesse, A. 185, 329).—Compound with pyrocatechin $B_2'C_8H_8O_2H_2SO_4$ aq: colourless needles (Hesse, C. C. 1889, 519).—Compounds with resorcin B'C₆H₆O₂H₂SO, $\frac{1}{2}$ aq: needles (Malin, A. 138, 77). $-B'_{2}C_{6}H_{6}O_{2}H_{2}SO_{4}$ aq (Hesse). — Compound with phloroglucin B'C,H,O,H,SO,2aq: stellate groups of needles (Hlasiwetz, Z. [2] 1, 618). -Compound with orcin B'C,H,O,H,SO, aq. Needles (Hlasiwetz a. Barth, A. 134, 290; 138, 77) .- Compound with chloral B'C2HCl2O. [149°]. Amorphous, sl. sol. cold alcohol (Mazzara, G. 16, 270).—Compound with m-nitrobenzoic aldehyde B'C₅H₄(NO₂).CHO. [118°_ 118°]. Yellow powder (Mazzara, G. 13, 368).—

Compound with nitro - camphor $B'(C_{10}H_{15}(NO_2)O)_2$ aq. Needles. [c. 181°]. [a]_D = +46° in a 2.7 p.c. alcoholic solution (Cazeneuve, Bl. [2] 49, 97).—Compound with urea B'CON₂H₄H₂Cl₂ Saq. S. 6. Small prisms (Drygin, C. C. 1878, 622; 1881, 245).—Compounds with quinidine B'C₂₀H₂₄N₂O₂ 2\frac{1}{2}aq.—B'C₂₀H₂₄N₂O₂ 3aq.—B'C₂₀H₂₄N₂O₂C₆H₆ 2aq (Wood a. Barret, C. N. 45, 6; 48, 4; Hesse, A. 243, A. 146). - Compound with hydroquinidine $B'C_{20}H_{28}N_2O_2$ $2\frac{1}{2}aq$: slender white needles, v. sl. sol. water, m. sol. ether.—Compounds with cinchonidine $B'2C_{10}H_{22}N_2O$. Rhombohedra, very slightly sol. ether (Hesse, A. 243, 131).

— B'7C₁₉H₂₂N₂O. Crystals (from alcohol).

— Crystals (from alcohol). $B'_{2}(C_{19}H_{22}N_{2}O)_{4}(H_{2}SO_{4})_{2}$ 20aq. S. 597 at Needles. — $B'_{2}(C_{19}H_{22}N_{2}O)_{4}(C_{4}H_{4}O_{6})_{3}$ 6aq. S. 597 at 15°. Needles. — $B'_2(C_{19}H_{22}N_2O)_4(C_4H_8O_8)_3$ 6aq. — $B'_2(C_{19}H_{22}N_2O)_4(H_2CrO_4)_3$ 18aq: crystals. — $B'_2C_{19}H_{22}N_2O(H_2C_2O_4)$ 6aq (?). Long colourless needles. — Compounds with cupreïne $B'C_{10}H_{22}N_{2}O$ 4aq. [177°]. $[\alpha]_{D} = -236^{\circ}$. Trimetric prisms (Howard a. Hodgkin, C. J. 41, 66; Hesse, A. 225, 98; 226, 242; 230, 72). Its solution in dilute H2SO, shows blue fluorescence. -B'C, H₂₂N₂OH, PtCl, 2aq : orange-red prisms. B'C₁₉H₂₂N₂OH₂SO₄ 6aq: six-sided prisms. S. 3 at 100°.—B'C₁₉H₂₂N₂OC₄H₂O₅ 2aq: needles.

Acetyl derivative C₂₉H₂₇AcN₂O₂. [108°].

Formed by heating quinine with Ac2O (Hesse, A. 205, 317). Prisms, sol. alcohol and chloroform, sl. sol. ether. $[a]_D = -54^\circ$ in a 2 p.c. solution in alcohol (of 97 p.c.) at 15°. $[a]_D = -115^\circ$ in presence of 3HCl. Decomposed by alcoholic potash into quinine and HOAc.—B'H2PtCl, 2aq. $-B'(HAuCl_1)_2$ aq : yellow flocculent pp.

Propionyl derivative C₂₂H₂₂(C₃H₃O)N₂O₂. [129°]. Six-sided prisms, sol. ether and alcohol. $[a]_{D} = -109^{\circ}$ in a 2 p.c. solution at 15° (Hesse).—B'H2PtCl, 2aq.— B'(HAuCl.), 2aq: yellow amorphous pp.

Benzoyl derivative C₂₀H₂₃BzN₂O₂. Amorous (Schützenberger, C. R. 47, 334). phous

B'H,PtCl

Methylo-iodide B'MeIaq or

C₁₀H₇(OMe)N.C₈H₁₄NMeIO (Grimaux, Bl. [3] 7, 573). [233°-236°]. Formed by boiling quinine (1 mol.) with MeI (6 mols.) in MeOH (Strecker, A. 91, 164; Claus a. Mallmann, B. 14, 76). Formed also from cupreïne, NaOEt and MeI (Hesse, A. 266, 240). Colourless needles (from water), v. sol. hot water and alcohol. Not attacked by alkalis in the cold. Boiling KOHAq or moist Ag₂O converts it into methyl-quinine C₂₀H₂₂MeN₂O₂, an oil which yields an amorphous hydroiodide, crystalline B'H. PtCl, aq (Hesse) or B'H₂PtCl_e 2aq (Lippmann, M. 12, 512), and C_{2e}H₂₃MeN₂O₂MeI aq crystallising in needles, decomposed at 218°.

Methylo-periodides B'MeI. needles (Jörgensen, J. pr. [2] 3, 145; 14, 261).— B'_Me_I_H_SO_. Reddish-brown needles (from alcohol).—B'2Me2I3H2SO4. Brown plates.--B',Me,I,82H,8O,. Almost black laminæ.-Almost black needles with $B'_1Me_1I_{22}2H_2SO_4$.

green lustre.

Methylo-chlorids B'MeClaq. [182°]. Needles. -B'2MeHPtCl_s: orange prisms.

B'MeBr aq. [126°]. Methylo-bromide

Slender needles, sl. sol. cold water.

B'Me, I, 3aq. Di-methylo-di-iodide [158°-162°]. Formed by heating B'MeI with

MeI and MeOH in sealed tubes at 100°. Got also. together with B'MeI from cupreïne by the action of NaOMe followed by MeI (Hesse, A. 266, 240). Yellow tables (from water). Converted by cold NaOHAq, or better NaOH in MeOH, into a resin and a smaller quantity of a yellow crystalline body. S. (boiling MeOH) ·1, melting about 280°, sol. acids and reppd. by alkalis as a jelly. Its solutions are yellow with green fluorescence. A similar body is got by the action of alkalis on the methylo-iodide of methoxy-quinoline (obtained from quinine). - B'Me2PtCl22aq. -B'2MeAuCl4.

Ethylo-iodide B'Etlaq (Howard, C. J. 26, 1180). [211°]. Trimetric needles (from ether), with very bitter taste. Lævorotatory.

 $-B'EtI_s$: black needles.

Ethylo-chloride B'EtCl 3aq. needles.—B'EtHPtCl: yellow pp. Ethylo-bromide B'EtBr 2aq. Slender

Ethylo-sulphates B'zEtzSO, 8aq. -B'EtHSO, 2aq: needles, v. e. sol. water.

Ethylo-cyanide B'EtCN. [90°]. White

needles (Claus a. Merck, B. 16, 2746).

Di-ethylo-di-iodide B'Et₂I₂3aq. [115°]. Prepared by heating quinine with EtI and alcoholic potash (Skraup, M. 2, 610). Monoclinic tables, sol. alcohol, insol. ether.

Methylo-ethylo-di-iodide B'MeEtI2 aq. [208°]. Formed by boiling B'MeI with EtI and alcohol (Claus, B. 14, 76). An isomeride [157°-

160°] is got from B'Etl and Mel.

Benzylo-chloride B'C,H,Cl. Amorphous.

B'(C,H),HPtCl₂2aq: crystalline (Mazzara, G. 13, 530).—B'(C,H),PtCl₆: yellow powder.
Tolyl-quinine C₂₀H₂₂(C,H,)N₂O₂. Two modifications are formed by heating quinine with o toluidine, and two from p-toluidine. The o and p (a) modifications form oils soluble in ether; the o and p (8) modifications form amorphous yellow powders insoluble in ether, soluble in chloroform and alcohol. B'₂H_{*}Cl₂PtCl₄aq: yellow crystalline powders (Claus a. Bottler, B. 14, 80)

Quinopropyline C₂₄H₂₈N₂O₂. [164°]. Got by heating sodium-cupreine with PrNO₃ and PrOH at 110° (Grimaux a. Arnaud, Bl. [3] 7, 310). White powder.—B'₂H₃SO₄ 1¹/₂aq. Silky needles.

 $[224^{\circ}]$. $[a]_{D} = 229 \text{ at } 22^{\circ}$

Quinoisopropyline [154°]. Formed in like manner from PrNO₂.-B'₂H₂SO₄ aq. S. 3 at 10°.

 $[a]_{D} = 229^{\circ}$.

Quinoamyline C₂₅H₃₂N₂O₂. [167°]. Formed from sodium-cupreïne and amyl chloride in amyl alcohol at 105°. Amorphous.—B' H2SO, 2aq. Needles. S. 025 at 11°. Its solution in dilute H2SO, fluoresces.

Quinine sulphonic acid C20H23(SO3H)N2O2. [209°]. Formed by moistening quinine tetrasulphate with Ac2O, and treating the product with hot water (Hesse, A. 267, 141). Small white prisms (containing aq when air-dried). [a]_D = -182° in a 2 p.c. solution containing HCl (3 mols.).—H₂A'₂H₂PtCl_u 8aq. Quinine (Iso)-sulphonic acid

C₂₀H₂₂N₂O₂(SO₂H). Formed from quinine and fuming H₂SO₄ (Hesse, A. 267, 138). V. sol. water, forming a slightly acid lavorotatory solution with blue fluorescence. Gives a dark-green colour with chlorine-water and NH,Aq.— HA'HAuCl .: yellow flocculent pp.

Apoquinine $C_{10}H_{22}N_2O_3$. [160°]. [α]_D = -178° in a 2 p.c. solution of alcohol (of 97 p.c.) at 15°. $[\alpha]_{D} = -247^{\circ}$ in water containing 3HCl. Formed by heating quinine or cupreïne with HClAq at 140° (Hesse, A. 205, 323, 341; 230, 65). Amorphous powder (containing 2aq), sol. alcohol, ether, CHCl_s, and hot water. A solution of its sulphate shows no fluorescence. Chlorine-water and ammonia give a dark-green colour in the The alcoholic solution gives a dark cold. brownish-red colour with FeCl₃. Fuming HClAq at 150° forms $C_{19}H_{23}ClN_2O_2$ 2aq [160°], which gives the salts B'H₂Cl₂8aq, B'H₂PtCl₆2aq, and

 $C_{19}H_{21}Ac_2ClN_2O_2$ [184°]. Salts. — B'H₂PtCl₆ 3aq. — B'HI. crystals (Lippmann a. Fleissner, M. 12, 331).

Acetyl derivative C19H20Ac2N2O2: amorphous powder, sol. ether and alcohol.

Isospoquinine C₁₉H₂N₂O₂. [176°]. Formed by the action of boiling alcoholic potash on C₁₉H₂₂N₃O₂3HI [238°] (v. QUININE, Reaction 8). Crystalline (Lippmann a. Fleissner, M. 12, 331).

-B'H₂PtCl₈ aq: granules, sl. sol. water. Isoquinine $C_{20}H_{24}N_2O_2$. [186°]. [a]_D = -181° in a 4 p.c. solution. Formed from the compound $C_{20}H_{24}\dot{N}_2O_23HI$ [230°] (v. Quinine, Reaction 9) with alcoholic potash (Lippmann a. Fleissner, M. 12, 332). Needles (containing aq), v. e. sol. benzene, sl. sol. ether, v. sl. sol. boiling water. Coloured green by chlorine-water and ammonia. -B'HCl 2aq : needles, v. sol. water.—B'2HCl, m.sol. water.—B'H2PtCl6: yellow crystalline pp.-B'2H2SO, 10aq: needles, v. e. sol. water.— B'AgNO_s: needles.

 ψ -Quinine $C_{20}H_{24}N_2O_2$. [191°]. Formed, together with nichine $C_{10}H_{24}N_2O_2$ (?), by heating quinine hydroiodide with alcoholic potash (Skraup, B. 25, 2911). Lævorotatory. Yields B'HCl, B'HNO3, and B'2HNO3, all sl. sol. water

and readily crystallised.

Chinene or Quinene v. QUINIENE, vol. ii. p. 181.

Homoquinine v. CINCHONA BASES.

QUININIC ACID v. Methyl derivative of (B. 3)-OXY-QUINOLINE CARBOXYLIC ACID.

QUINISATIC ACID C₆H₄(NH₂).CO.CO.CO₂H. o-Amido-bensoyl-glyoxylic acid. Formed by oxidation of (Py. 1,2,3)-tri-oxy-quinoline with FeCl₂ (Baeyer a. Homolka, B. 16, 2219). Yellow prisms, v. sol. water. On reduction with zincdust and HOAc and exposure of the filtrate to the air it forms a dark-blue colouring matter.

Anhydride C.H. CO.CO.NH.CO. [255°-260°]. Formed by heating the acid at 120°. crystals, forming a red solution in alcohol. Readily combines with water, with re-formation of the soid. Dilute NaOHAq forms a yellow solution, very quickly becoming colourless. NaOEt gives a body crystallising in indigo-blue Gives an oxim $C_6H_4 < \begin{array}{c} CO.C(NOH) \\ N=0.OH \end{array}$ needles.

crystallising in orange prisms [208°].

QUINITANNIC ACID. An acid occurring in cinchona bark (Schwarz, J. pr. 56, 76; cf. Pelletier a. Caventou, A. Ch. [3] 15, 837). Hygroscopic yellow mass. Its alkaline solutions absorb oxygen. Converted by boiling dilute acids into glucose and a brownish-red powder C28H2014, which gives acetic and protocatechnic acid when fused with potash (Rembold, A. 143, 270).

QUINITE C.H.2O2 i.e.

 $CH(OH) < \frac{CH_2 \cdot CH_2}{CH_2 \cdot CH_2} > CH(OH).$ Hydroquinone hexahydride. Formed, by reduction with sodiumamalgam in a current of CO2 from the product of the action of dilute H₂SO₄ on the dihydride of di-oxy-terephthalic ether (succinyl-succinic ether) (Baeyer, B. 25, 1037, 1840). V. sol. water and alcohol. Occurs in two modifications, trans [144°] and cis [90°], which yield acetyl derivatives melting at 106° (trans) and 32° (cis). Both acetyl derivatives boil at 244°. Quinite is converted by HBr into di-bromo-benzene hexahydride (trans [114°], cis an oil), which on heating with quinoline give oily benzene dihydride (81°), which forms a tetrabromide [182°].

QUINIZARIN v. DI-OXY-ANTHRAQUINONE. QUINIZINE. Substances supposed to be derived from the hypothetical quinizine were subsequently found to be derivatives of phenylpyrazole. Thus oxy-methyl-quinizine is oxyphenyl-methyl-pyrazole (q. v.). QUINOL v. Hydroquinone.

QUINOLIC ACID C.H. N.2O. Got by oxidation of cinchonine with HNO, (Weidel, A. 173, 91; B. 12, 1152). Woolly crystals, almost insol. water, v. sl. sol. alcohol, sol. HClAq. Alkalis give a transient crimson colour. Br and water at 180° give hexa-bromo-quinoline. HNO₃ at 170° forms cinchomeronic acid.—AgA'.— 170° forms einchomeronic acid.—A HA'HCl.—H₂A'₂H₂PtCl₆: orange needles. QUINOLINE C₆H₇N i.e.

ÇH:CH.C.CH:ÇH Chinoline. Leucoline. Mol. CH:CH.C.N :CH' w. 129. (237.5°) (Young, C. J. 55, 485); (234°) (Schiff, B. 19, 566); (241° cor.) (Kretschy, M. 2, 80). S.G. 2 1.106 (O. de Coninck, Bl. [2] 37, 208); 2 1·108 (Skraup). S.V. 139·8 (Schiff). Vapour-pressure: Young. Heat of neutralisation: Colson, A. Ch. [6] 19, 409. Absorption of ultra-violet spectrum: Hartley, C. J. 41, 47. Occurs in coal-tar (Runge, P. 31, 68; Hofmann, A. 47, 76; 53, 427; 74, 15; Greville Williams, Tr. E. 21 [2]; [3] 377; O. Fischer, B. 16, 720; Jacobsen a. Reimer, B. 16, 1084).

Formation .- 1. By distilling quinine, cinchonine, or strychnine with potash (Gerhardt, A. 42, 310; 44, 279; O. de Coninck, A. 42, 310; 44, 279; O. de Coninck, C. R. 94, 87; Bl. [2] 35, 296). Quinoline free from homologues is got from cinchonine by treatment with potash and CuO (Wyschnegradsky, B. 13, 2318). -2. By passing allyl-aniline over heated PbO (Königs, B. 12, 453).—3. From hydrocarbostyril (vol. i. p. 180) by treatment with PCl, and reduction of the product with HI and HOAc (Baeyer, B, 12, 1320).—4. By distillation of acrolein-aniline (Königs, B. 18, 911).-5. By distilling anil-uvitonic acid with soda-lime (Böttinger, B. 13, 2165).—6. By heating a mixture of nitro-benzene, aniline, glycerin, and H.SO, for three hours with inverted condenser (Skraup, M. 1, 316; 2, 139, 535).—7. By heating its carboxylic acids with lime .--8. By adding a few drops of NaOHAq to a cold dilute solution of equimolecular quantities of o-amido-benzoic aldehyde and acetic aldehyde (Friedländer a. Gohring, B. 16, 1833).—9. By heating the hydrochloride of quinoline tetrahydride with water and Hg(OAc)₂ at 150° (Tafel, B. 25, 1623).—10. By fusing methyl-acetanilide with ZnCl₂ at 290° (Piotet, B. 23, 1903).

Preparation.—A mixture of aniline (38 pts.), glycerin (120 pts.), nitro-benzene (24 pts.) and H₂SO₄ (100 pts.) is heated with inverted condenser. Potash is then added, and the base distilled over with steam and fractionally distilled. It may be further purified by means of the acid sulphate and freed from aniline by oxidation or by treatment in acid solution with NaNO₂. The

yield is 60 p.c. (Skraup).

Properties. - Colourless liquid with peculiar odour, slowly becoming yellow. V. sl. sol. water, miscible with alcohol, ether, and CS₂. Quinoline is very hygroscopic, forming the hydrate (C,H,N), 3aq, which becomes turbid on warming (Hoogewerff a. Van Dorp, R. T. C. 1, 1, 107). It is antiseptic, antipyretic, and antizymotic (Donath, B. 14, 178). Quinoline separated from solutions of its salts dissolves in excess of NH, or ammonium carbonate, but not in NaOH or Na₂CO₃. Iodine in KI gives a brown pp., insol. HClAq. Phosphomolybdic acid a yellowishwhite pp., sol. NH₃Aq. Picric acid a yellow amorphous pp. HgCl₂ a white pp., sol. HClAq. K_2 HgI, a yellow amorphous pp., changed to yellow needles on adding HCl. K_2 Cr₂O, gives crystals, sol. excess. Potassium ferrocyanide an amorphous orange pp. (Donath, B. 14, 1769). On shaking with conc. CuSO, Aq a green pp. (CuSO₄)₂(CuO₂H₂)₈4aq is got (Borsbach, B. 23, 924; cf. Lachovitch, M. 10, 884). NaOBr gives no reaction (Deniges, C. R. 107, 662). Quinoline forms very soluble crystalline compounds with NaHSO_s (Brunck a. Graebe, B. 15, 1785). combines with iodoform (Rhoussopoulos, B. 16,

Reactions.—1. On passing through a red-hot tube it yields (β)-diquinolyl (Zimmermann a. Müller, B. 17, 1965).—2. On boiling with sodium or sodium-amalgam it is converted into 'diquinoline,' which forms a scarlet hydrochloride which dyes silk a transient orange (Greville Williams, Pr. 31, 536; C. N. 37, 85).—3. Reduced by tin and HClAq to quinoline tetrahydride (244°) (Wyschnegradsky, Bl. [2] 34, 339).-4. Zincdust and NH3Aq or alcohol and sodium-amalgam give the tetrahydrides of quinoline and of diquinoline (Königs, B. 14, 99).—5. Oxidised by bleaching-powder or other hypochlorites to oxy-quinoline (carbostyril) and chloro-oxy-quinoline (Erlenmeyer, B. 19, 489; Einhorn a. Lauch, A. 243, 342). Chlorine passed into its solution in HOAc forms tri-chloro-oxy-carbostyril.—6. On heating with SbCl_s at 170°-400° and passing in chlorine, the products are C₂Cl_s and C₆Cl₆ (Smith a. Davis, C. J. 41, 413).—7. Bromine (2 pts.) added to a mixture of quinoline (1 pt.) and water (3 pts.) forms the tetrabromide C₂H₂NBr₄, which crystallises from chloroform in very unstable red needles, and is converted, by heating with alcohol, into C.H.NBr.HBr [86°] crystallising in red prisms, insol. chloroform, v. sol. alcohol and ether, decomposed at 180° into HBr and bromo-quinoline-hydrobromide (Grimaux, Bl. [2] 38, 125; C. R. 95, 85). Bromine added to an ethereal solution of quinoline forms the dibromide C,H,NBr, yielding B'HCl [100°-105°] and B'HBr [88° cor.], which form red crystals (Claus a. Collischonn, B. 19, 2765).—8. KMnO. in alkaline solution forms pyridine dicarboxylic acid (Hoogewerff a. Van Dorp, R. T. C. I, 1, 107; of Dewar, Pr. 26, 65). KMnO₄ and conc. H₂SO₄

also give quinolinic acid. KMnO, and dilute H₂SO, form quinoline (B. 3)-carboxylic acid, diquinolyl being an intermediate product (Georgievitch, M. 12, 312).—9. H₂SO₄ at 220° forms the (B. 4)-sulphonic acid; at 250° the (B. 2)-sulphonic acid is produced (Georgievitch, M. 8, 578, 641).—10. Oxygen passed through platinised asbestos containing quinoline and quinoline hydrochloride at 190° forms (a)-diquinolyl. Oxygen acting on a mixture of aniline and quinoline hydrochloride forms (Py. 3; B. 2)-diquinolyl and p-amido-(Py. 3)-phenyl-quinoline, while quinoline and o-toluidine hydrochloride are converted by oxygen into amido-phenyl-methyl-quinoline (Weidel, M. 8, 120; 9, 99).—11. p-Amido-phenyl-quinoline is also formed by heating quinoline hydrochloride with aniline.-12. BzCl at 240° gives (β)-diquinolyl.—13. Alloxan added to a solution of quinoline saturated with SO₂ forms B'C₄H₂N₂O₄SO₃H₂ crystallising in yellowish prisms (Pellizzari, A. 248, 150).— 14. SiCl, forms (C₂H,N)₂SiCl₄ (Harden, C. J. 51, 40). SiF₄ forms (C₂H,N)₂SiF₄, crystallising in needles (Comey a. Jackson, Am. 10, 176).—15. Nitroso-di-methyl-aniline and hydrogen cyanide form B'(C₆H₄(NO).NMe₂)₂HCy, crystallising in golden plates (Lippmann a. Fleissner, M. 6, Glycerin dichlorhydrin 543). — 16. (C,H,N)C,H,Cl, which is deliquescent and forms (B'C₃H₃Cl)₂PtCl₄ and B'C₃H₃AuCl₄.—17. Ethylens chloride at 100° forms B'₂C₂H₄Cl₂, crystallising in needles, yielding B'₂C₂H₄PtCl₆ (Rhoussopoulos, B. 16, 879).—18. Ethylene bromide (1 mol.) heated with quinoline (1 mol.) at 80° forms the bromo-ethylo-bromide B'BrC₂H₄Br, crystallising from alcohol in needles and yielding B'ClC2H4Br and (B'ClC,H,Br),PtCl, (Berend, B. 14, 1349). Ethylene bromide (1 mol.) with quinoline (2 mols.) at 40° forms B'C₂H₄Br₂ aq crystallising from alcohol in needles.—19. By heating quinoline (16 g.) with glycolic chlorhydrin (10 g.) and water (10 c.c.) for 3 days at 100° there is formed B'Cl.C2H4OH, crystallising from alcohol-ether in splendid prisms (Wurtz, Pr. 33, 452; C. R. 95, 263; 96, 1269). It is hygroscopic, v. sol. water and alcohol, insol. ether. On boiling with moist Ag₂O it forms a caustic liquid that soon turns crimson. It yields B'(C2H4OH)AuCl4, crystallising in minute pointed hexagons, and also B'(C₂H₂OH)Cl6HgCl₂, and (B'(C₂H₂OH)Cl)₂PtCl₄.

20. Iodoform (1 mol.) heated with quincline (2 mols.) for 8 days at 100° forms B'2CH2I2, crystallising in long needles [132°] (Rhoussopoulos, B. 16, 202, 880, 2004). It is converted by AgCl into B'2CH2Cl2 [168°], which gives B'₂CH,PtCl_s. Iodoform (1 mol.) added to quinoline (3 mols.) in ethereal solution forms B'₂CHI₂ crystallising in needles [65°], not affected by AgCl.—21. Chloro-acetic acid forms quinoline-betaine C₁₁H₂NO₂ [171°], which separates from alcohol in thick crystals (containing aq) and forms (C₁₁H₉NO₂)₂H₂PtCl₆2aq (Gerichten, B. 15, 1254; Rhoussopoulos, B. 15, 2006). -22. Chloroacetic ether forms C,H,NCl.CH,CO,Et crystallising in needles, v. e. sol. water, converted by moist Ag₂O into quinoline. It yields the salt C₁₃H₁₄NO₂)₂H₂PtCl_e. — 23. Chloral in ether forms CoH, NC, HCl, O aq [66°], crystallising from benzene (Rhoussopoulos, B. 16, 881). It is insolwater and decomposed by alcohol. It yields (C.H.NC.HCl.O aq) BPtCl. - 24. Resorcin at

100° forms (C₀H,N)₂C₀H₆O₂, [102°], S. 25 in the cold (Hock, B. 16, 886). It crystallises from dilute alcohol in silvery plates, decomposed by HClAq.—25. Phthalic anhydride at 150° forms 'quino-phthalone' C₁₇H₂NO₂ [235°] (Traub, B.

16, 297).

Salts. - B'HCl. [94°]. Deliquescent nodules, v. e. sol. alcohol, chloroform, and hot ether (O. de Coninck, Bl. [2] 37, 208). B'₂H₂PtCl₆ 2aq [225°] (Skraup); [218°] (Lellmann, A. 237, 323).—B'₂H₂PtCl₆ aq. S. 07 at 11°.—B'₂PtCl₂: insoluble powder.—B'₂H₂PtCl₆.— B'HAuCl., Canary-yellow needles.—B'HCdCl, aq.
—B'CdCl₂: white pp.—B'HgCl₂. Pearly plates
(Bromeis, 4. 52, 136).—B'₂H₄HgCl, 2aq. [91°].
Monoclinic crystals.—B'₄H₂PdCl₄. Crystals.— B'H2SnCl, 2aq: needles.—B'HSnCl, [127°]. Thin needles.— B'₂H₂SnCl₃. [above 240°].— B'₂H₂UrO₂Cl₄: yellow prisms.— B'SbCl₃.— B'HSbCl₃.—B'₃H₃BiCl₃.—B'₂H₂ZnCl₄.—B'₂ZnCl₂. -B'HClICl. [118°] (Östermayer). Yellow pp., converted by ammonia into explosive B'NH₂I (Dittmar, B. 18, 1613).—B'ICl. [160°]. Small white needles (Pictet a. Krafft, Bl. [3] 7, 73).—B'Br₂HCl. [100°-105°]. Orange crystals (Claus, B. 19, 2766).—B'HMnCl₃: rose-red needles (Borsebach, B. 23, 433).—B'HFeCl. Formed by adding HCl to a solution containing quinoline and FeCl₃ (B.). Small needles, decomposed by boiling water.—B'₂CoCl₂: blue triclinic crystals.—B'₂CuCl₂: nearly black crystals.—B'Cu(OAc)₂: [above 240°]. Green crystalline B'Cu(OAc), [above 240°]. Green crystalline powder.—B', ZnI, white powder, sl. sol. cold water.—B'₂ZnBr₂.—B'CdBr₂.—B'CdI₂.—B',CdI₂.—B',CdI₃.—B'HgBr₂. [204°].—B'HgI₂. [168°].—B'Br₂. [92°-100°]. Red crystals (Lubavin, J. R. 18, 434).—B'Br₄. Red needles.—B'HBr₅. [86°].—B'Br₂HCl. [100°-105°].—B'I₂. [90°]. Formed by adding I to a solution of quinoline in CS, (Claus a. Istel, B. 15, 824). Dark-green lustrous needles.—B'HI₃. [67°]. Ppd. by adding I in KI to a solution of quinoline sulphate (Dafert, M. 4, 509). Green crystalline pp., sol. alcohol and benzene.—B'H₂SO₄. [164°]. S. (alcohol) 2 at 18°; 11 at 78° (Krakau, J. R. 17, 364). Deliquescent crystals. — $B'_2H_2Cr_2O_7$. [c. 167°]. Yellow needles (from hot water). S. 36 at 10°. -B'HNO₂. Needles (from alcohol), v. sol. water, insol. ether. - B'₂AgNO₃: needles. -B'2Hg(NO3)2: white crystalline precipitate. B'₂(HNO₃)₂Ce(NO₃)₄: orange-red plates (G. Williams, C. N. 58, 199).—B'₂H₂SiF₈. Long needles (Comey a. Jackson, Am. 10, 176).—B'H₂C₂O₄. Silky needles (from alcohol). Decomposes at Silky needles (from alcohol). Decomposes at 100° (Williams).—B'2HgCy2: long needles (from water) (H. Schiff, A. 131, 112).—Tartrate B',4C,H₀O₆. [125°] (Friese, B. 14, 2805).— Cyanurate B',H₃C,N₃O₅. Crystals (Claus, anurate B',H,C,N,O,. Crystals (Claus, pr. [2] 88, 226). — B'HSCy(Cr(SCy),NH,). Red lamellæ, sl. sol. hot water (Christensen, J. pr. [2] 45, 365). — Picrate: [203°]. — o-Oxybenzoate B'C, H,O, Crystalline powder.

According to Decker Alkylo-iodides. (B. 24, 690) the alkylo-iodides B'RI are converted by alkalis into the hydroxides B'ROH which are readily oxidised by the air to crystalline oxy- ν -alkyl-quinolines $C_eH_4 < \stackrel{CH:CH}{\sim}_{NR.CO}$, which melt about 100°. The cyanines may be considered as derived from one molecule of alkylquinoline and one molecule of oxy-v-alkyl-quinoline. The alkylo-hydroxides are converted by warming with alcohols R'OH into compounds B'ROR'

Methylo-chloride B'MeClaq. On heating with ZnCl₂ at 180° it is converted into a base $C_{20}H_{20}N_2\tilde{O}$ [72°-75°] (above 360°), crystallising in large trimetric prisms, and yielding a hydrochloride [112°] and the double salts C₂₀H₂₀N₂OHAuCl₄ and (C₂₀H₂₀N₂O)₂H₂PtCl₆ [190°] nearly insol. water (Ostermayer, B. 18, 593).—B', Me, PtCl. [230°]. Yellow plates.—B'MeAuCl. [205°].—B'MeClICl. [112°]. Formed by adding ICl to the methylo-chloride in aqueous solution. Large yellow plates.

Methylo-tribromide B'MeBr. [128°]. Orange-red plates. Converted by picric acid

solution into B'MeOC_eH₂(NO₂)₃ [164°].

Methylo-iodide B'MeI. [73°] (La Coste, B. 15, 192; Pictet, B. 23, 1903). Large crystals. Converted by moist Ag₂O into a strongly alkaline solution of the hydroxide B'MeOH. Aqueous NaOH in the cold converts B'MeI into the oxide B'2Me2O, a white powder, reconverted by HI into B'MeI (possibly oxy-v-methyl-quinoline). (La Coste; Claus, B. 15, 475; Bernthsen, B. 18, 29; Decker, B. 24, 690). Potash, even in cold. aqueous solution, forms an oil probably containing methyl-quinoline (lepidine) (Skraup, Sitz. W. [2] 81, 593). By the action of KOH on a mixture of B'MeI (2 pts.) and the methyloiodide of (Py. 1)-methyl-quinoline (1 pt.) there is formed di-methyl-cyanine lodide C₂₁H₁₉N₂I, crystallising in oblong green tablets, [291°], sl. sol. water, forming a reddish-blue solution (Hoogewerff a. Van Dorp, R. T. C. 2, 317; 3, 337). The alcoholic solution is violet. with green fluorescence. Its solution in acids is yellow. The aqueous solution is decolourised by CO₂, the colour re-appearing on boiling. The corresponding $C_{21}H_{19}N_2Cl$ 5aq [c. 300] forms $C_{21}H_{19}N_2Cl(HPtCl_3)$ aq as yellow crystals. Quinciling mathelatics oline methylo-iodide is oxidised by alkaline K₂FeCy₆ to oxy-methyl-quinoline or ν-methylquinolone C₆H₄ CH :CH :CH [72°] (324° at 728-

mm.) (Decker, J. pr. [2] 46, 31; cf. Bernthsen a. Hess, B. 18, 37, who took it for di-methyl-diquinolyl; Ostermayer, B. 18, 591). This body is also got from carbostyril and MeI (Friedländer a. Müller, B. 20, 2009). It deliquesces with water, forming an oil. Hydrochloride [112°].

Ethylo-chloride B'EtCl aq. [93°]. Large trimetric tables (Claus a. Tosse, B. 16, 1277). B'₂Et₂PtCl₆. [226°]. Yellow pp. Ethylo-bromide B'EtBr aq. [80°]. Tri-

metric tables. By the action of (Py. 1)-methylquinoline ethylo-bromide it is converted into diethyl-cyanine bromide C23H23N2Br or C23H23N2Br, crystallising in small needles, not melted at 290°.

forming a blue solution in alcohol.

Ethylo-iodide B'EtI. [118°] (Spalteholz, B.16, 1851); [160°] (Hoogewerff a. Dorp, R. T. C. 2, 321). Monoclinic crystals. At 280° it yields (γ)-ethyl-quinoline di-ethyl-quinoline and (Reher, B. 19, 2996). Moist Ag₂O forms strongly alkaline B'EtOH, which is sol. water and resinified by atmospheric oxidation. On heating: B'EtI (2 pts.) with (Py. 8)-methyl-quinoline-ethylo-iodide (1 pt.) and alcoholic potash, there is-formed di-ethyl-isocyanine iodide C.H., N.I.

(Hoogewerff a. Van Dorp, R. T. C. 8, 344) or | C22H25N2I (Spalteholz, B. 16, 1851). This body forms lustrous green prisms (containing aq or EtOH or EtOH 1 aq) melting at 152°. It is insol. ether, sl. sol. water, and forms a crimson solution in alcohol. By the action of KOH on a mixture of B'EtI and (Py. 1)-methyl-quinoline ethylo-iodide there is formed di-ethyl-cyanine iodide C₂₁H₂₃N₂I [273°], crystallising in green prisms (Hoogewerff a. Van Dorp, R. T. C. 2, Quinoline ethylo-iodide is oxidised by 817). alkaline K_3 FeCy_s to C_6H_4 < $\frac{CH}{NEt.CO}$ (Decker, J. pr. [2] 47,36), which body is also got from carbostyril and Etl (Friedländer a. Weinberg, B. 18, 1530). Solidifies at -20° . $C_0H_7NEtIHgCy_2$. [138°]. Needles (from water). C_sH,NEtBrHgOy₂. [155°]. Crystals (from alcohol) (Claus, A. 269, 271).

Ethylo-nitrate B'EtNO_s. [89° uncor.].

Large trimetric crystals (Claus, B. 16, 1277).

Propylo-chloride B'PrClaq. [c. 95°] (hydrated); [135°] (anhydrous). Colourless crystals (Claus a. Collischonn, B. 19, 2504). Crystallises from chloroform in prisms (containing CHCl₃) [79°]. V. e. sol. water.—B'PrClBr₂. [85° uncor.]. Orange-red radiating crystalline mass. - B'BrClI2.

[62°]. Brown crystals.

Propyl-bromide B'PrBr 2aq. [66° uncor.] (hydrated); [148° uncor.] (anhydrous). Formed by heating quinoline with PrBr and a little (10 p.c.) alcohol at 100°. Tables, v. e. sol. water and alcohol. The anhydrous crystals (obtained from absolute alcohol) are very deliquescent. Crystals obtained from chloroform (containing CHCl,) melt at 65°-129°.—B'PrCl,Br. [c. 60°]. Formed by passing dry Cl into a solution of B'PrBr in chloroform. Small sulphur-Unstable.—B'PrBr_s. plates. Formed by adding Br to the aqueous or alcoholic solution of B'PrBr. Triclinic garnet-red crystals. Yields (Py. 3)-bromo-quinoline on heating. B'PrBrl. [60° uncor.]. Formed by evaporating a mixture of I in ether and B'PrBr in alcohol. Lustrous brown needles.—B'PrBrI₄. [49°]. Slender, nearly black, needles with green reflex.

Propylo-iodide B'PrI. [145°]. Small yellow crystals. Not hygroscopic. Crystallises from chloroform with CHCl, partially melting at 92°. — B'PrCl.I. [87°]. Sulphur-yellow needles.—B'PrCl.I. [145°]. Converted by long boiling with water into B'PrCl.-B'PrBr.I. [77°]. Triclinic orange tables.—B'PrBr.I. [48°-58°]. Very unstable orange powder.—B'PrI_s. [62°]. Very stable bronzy brown prisms.—B'PrI_s. [62°].]. Stable tables.

Iso-amylo-chloride B'C,H,Cl.-

B's(C,H₁₁₎₂PtCl₂. [220°]. Yellow crystalline pp. Iso-amylo-bromide B'C₅H₁₁Braq. [87°] (hydrated); [140°] (anhydrous) (Claus a. Tosse,

B. 16, 1277).

Yellow monoclinic needles (Perkin, C. J. 14, 246; Nadler a. Merz, J. pr. 100, 129; Greville Williams, C. N. 1861, 219; Hoogewerff a. Van Dorp, R. T. C. 2, 40: 8. 852 4, 62). Does not form cyanine on treatment with potash unless it contains the isoamyloiodide of (Py. 1)-methyl-quinoline, in which case di-isoamyl-cyanine iodide C.H., N.I

[c. 100°] is produced. This body crystallises in lustrous green monoclinic crystals (containing $1\frac{1}{6}$ aq); a:b:c=2.379:1:1.669; $\beta=88^{\circ}45'$. It also forms brass-yellow crystals. It is very hygroscopic, nearly insol. ether, sl. sol. CHCl₃, acetone, and isoamyl alcohol. It is v. sol. alcohol, forming a deep-blue solution. Iodine converts it into $C_{29}H_{35}N_2I_3$ [189°], which is sl. sol. alcohol, v. sol. acetone, forming blue solu-The solution of di-isoamyl-cyanine iodide in HClAq is colourless, and deposits on evaporation colourless scales of C29H85N2I2HCl, which at 100° gives off HCl, leaving bronzed C₂₀H₃₁N₂HCl. Moist Ag₂O forms uncrystallisable C₂₀H₃₁N₂(OH). Alcohol and AgCl produce C28 H35 N2Cl 4aq, crystallising in long blue prisms, and yielding (C₂₀H₃₅N₂Cl)₂PtCl₄. Alcohol and AgNO₃ give C₂₀H₃₅N₂(NO₃) aq, which forms lustrous needles giving a blue aqueous solution and yielding colourless $C_{2p}H_{33}N_2(NO_s)2HCl$ and blue $C_{25}H_{35}N_{2}(NO_{3})HCl$. Alcoholic ammonium sulphide converts the nitrate into reddishyellow monoclinic crystals of $C_{58}H_{68}N_4S_3O_2$, which yields $C_{56}H_{68}N_1S_3Cl_42PtCl_4$ as an orange pp. (N. a. M.). H₂SO₄ converts di-isoamyl-cyanine iodide into $(C_{26}H_{26}N_2)_2$ SO₄ 2aq, crystallising from hot water in blue needles.

Allylo-iodide B'C,H,I. [177.5°] (Pictet,

C. R. 95, 800).

Benzylo-chloride B'C,H,Cl 3aq. [65°]. Triclinic tablets. It crystallises also as B'C,H,Cl 2aq [130°] in trimetric crystals (Claus, B. 13, 2045; 16, 1279; 18, 1305). V. sol. water and alcohol, insol. ether. KMnO, oxidises it to benzyl-o-amido-benzoic acid and its formyl derivative. By treatment with KOH or Ag₂O it is converted into a strong base, v. sol. water. Excess of KOH ppts. an oil which turns red in air. The benzylo-hydroxide yields quinoline on distillation (Bernthsen a. Hess, B. 18, 35).

Mercuric chloride yields a crystalline pp. [142°].—(B'C,H,c)lBr₂. [246°]. Yellow pp.—

B'C,H,ClBr₂. [92°]. Crystalline.—B'C,H,BrCl₂. [80°]. Yellow needles.—B'C,H,Br,. Orange prisms.—B'C,H,BrI, [110°]. violet needles.

Phenacylo-bromide B'Br.CH,Bz. [115°-165°]. Needles, v. sol. water and alcohol, v. sl. sol. ether (Bamberger, B. 20, 3340).—B'HNO₃.

References .- AMIDO-, BROMO-, BROMO-AMIDO-, BROMO-NITRO-, CHLORO-, CHLORO-NITBO-, IODO-, NITRO-, OXY-AMIDO-, and OXY- QUINOLINES.

Diquinoline v. DiquinolyL.

Isoquinoline C₉H₇N i.e. C₈H₄ CH:ÇH. [23°].

(241° i.V.). Occurs, as well as quinoline, in coal-tar (Hoogewerff a. Van Dorp, R. T. C. 4, 125, 285; 5, 305).

-1. By the action of HI at 230° Formation. or of zinc-dust in a current of H at a low red heat upon the di-chloro- derivative [123°] obtained by the action of POCl, on the imide of C.H. CO.NH o-carboxy-phenyl-acetic acid

(Gabriel, B. 18, 3470; 19, 1656, 2361; Le Blanc, B. 21, 2299).-2. Together with di-isoquinolyl, which yields B'₂H₂PtCl₂ [177°] by distilling benzylidene-ethyl-amine (Pictet a. Popovici, Is. 25, 733).—8. By heating isocarbostyril or its carboxylic acid with zinc-dust in a current of hydrogen (Bamberger a. Kitschelt, B. 25, 1146;

Zincke, B. 25, 1497).

Properties. - Hygroscopic tables. Yields phthalic and cinchomeronic acids on oxidation by KMnO4. Tin and HClAq give a tetra-hydride. Forms crystalline compounds with HgCl, and AgNO₂. Bromine in ether gives C₀H₇NBr₂, [82°], which forms very unstable C,H,NBr,HBr, [c. 133°], converted at 200° into bromo-iso-

quinoline [40°] (Edinger, J. pr. [2] 43, 191).
Salts.—B'₂H₂PtCl₂2aq. [264°]. Reddishyellow crystals.—B'HAuCl₄. [225°]. Yellow needles.—B'HgCl₂. [165°]. White needles.—B'H₂SO₄. [206°].—B'H₂Cr₂O₇: needles.—B'Erate B'C₆H₃N₃O₇. [223°]. Yellow needles.

Methylogicidal Riving 1100° [100°].

Methylo-iodide B'Mel aq. [160°]. Flat yellowish needles. Treatment with AgCl and platinic chloride gives (B'MeCl),PtCl, [240°] (Claus, J. pr. [2] 38, 492). KOHAq added to B'MeI forms an oily base which yields (B'MeCl)₂PtCl₄ 6aq. Oxidised alkaline by K_s FeCy_s to ν -methyl-isoquinoline C_sH_s < CH:CH CO.NMe [40°] (319°), sol. water, and yielding
 B'H_PtCl_e 2aq (Decker, J. pr. [2] 47, 37).
 Ethylo-iodide B'Etl. [147°]. Yellowishyielding

red tables. Yields ethyl-phthalimide on oxida-

tion. Gives (B'EtCl),PtCl.

Benzylo-chloride B'C,H,Cl. V. sol. water and alcohol, sl. sol. ether (Goldschmidt, M. 9, 678). Yields benzyl-phthalimide [115°] on oxidation by KMnO, in a solution kept neutral by HCl.

Phenacylo-bromide B'BrCH2Bz. [205°]. Formed from isoquinoline and ω-bromo-acetophenone dissolved in benzene (G.). Prisms (from alcohol), v. sol. water. AgNO forms B'NO_s(CH,Bz) crystallising in needles.

Diquinoline v. DIQUINOLYL

QUINOLINE (B. 1)-CARBOXYLIC ACID

ÇH:C(CO,H).C.CH:ÇH

Quinoline ana-carb-- O−N:ĊH° CH:CH oxylic acid. Quinoline-m-carboxylic acid. [857°].

Formation.—1. Together with a smaller quantity of the (B. 3)-acid by heating m-amidobenzoic acid with glycerin, H,SO,, and m-nitrobenzoic acid (Schlosser a. Skraup, M. 2, 518; 7, 519).—2. By saponifying its nitrile, which is got by distilling sodium quinoline (B. 1)- or (B. 4)sulphonate with KCy (Bedall a. Fischer, B. 14, 2574; 15, 683, 1980).—3. By heating quinoline (B. 1,4)-di-carboxylic acid (Skraup a. Brunner, M. 7, 153).

Properties .- White crystalline powder, sol. dilute acids and alkalis, insol. ether and benzene,

v. sl. sol. water and alcohol.

Salts .- HA'HCl 1 aq : white needles.-(HA'),H,PtCl.: yellow plates. — CaA', 2aq. — CaHA', 6aq: needles. — HO.CuA' 2aq: minute violet plates, insol. water. AgA': crystalline pp.

Nitrile C.H.N.CN. [88°]. (above 360°). Formed as above and also from (B. 1)-amidoquinoline by diazotisation and treatment with CuCy₂ (Freydl, M. 8, 581). Crystals, sl. sol. Aq.

Tetrahydride C_{1e}H₁₁NO₂. [147°]. Formed by reducing the acid with tin and HCl (Fischer a. Körner, B. 17, 765). Yields a crystalline nitrosamine C_{1e}H_{1e}N₂O₂. [186°].

+-Isomeride C_{1e}H₂NO₃. [338°]. Perhaps identical with the preceding. Formed by warm-

ing quinoline with m-amido-benzoic acid, nitrobenzene, glycerin, and H.SO, (Lellmann a. Alt, A. 237, 318). Got also by saponifying its nitrile which is formed by fusing sodium quinoline ψ-ana-sulphonate with KCy in a current of hot air (Lellmann, B. 20, 1449; 21, 397). Snowy powder.—HA'HCl aq.— $H_2A'_2H_2PtCl_6$: needles.— ZnA'2.—AgA'.

Tetrahydride C10H11NO2. [147°]. Got by reducing the acid or bromo-quinoline carboxylic acid with tin and HClAq. - HA'HCl aq.

Nitrile C.H.N.Cy. [89°]. Needles (containing 1½ aq), melting at 70°. Effloresces, and then contains and melts at 74.5°. - $B'_{2}H_{2}PtCl_{s}$: needles.

Quinoline (B. 2)-carboxylic acid

CO₂H.C=CH.C.CH:CH

CH:CH.C-N:CH Quinoline p-carboxylic acid. [291°]. Formed by heating p-amido-benzoic acid with glycerin, p-nitro-benzoic acid, and H₂SO₄ (Schlosser a. Skraup, M. 2, 518). Formed also by heating its nitrile with HClAq at 140° (Fischer a. Wittmack, B. 17, 440). Thick prisms, sl. sol. hot water, v. sol. acids and alkalis.—HA'HCl.—(HA')2H2PtCl. — CaA'2 2aq. -AgA'.

Nitrile C₁₀H₈N₂. [131°]. Formed by distilling potassium quinoline p-sulphonate (from sulphanilic acid) with KCy. Needles, forming a

red solution in HClAq.

Quinoline (B. 3)-carboxylic acid

CH:CH.C.CH:CH Quinoline-m-carb-CO2H.C=CH.C-N:CH

oxylic acid. [248°].

Formation.—1. By oxidation of (β)-diquinolyl with CrO, and HOAc (O. Fischer a. Loo, B. 17, 1901; 19, 2473).—2. By oxidation of (B. 3)-methyl-quinoline (Skraup a. Brunner. M. 7, 142).—3. By heating the stannochloride of c-amido-phthalic acid with c-nitro-phthalic acid, glycerin, and H.SO. (Tortelli, G. 16, 367).

Properties.—Needles (from water).

Salts.-Cu(OH)A' aq.-AgA'.-HA'HCl aq: triclinic prisms. - H,A',H,PtCl,: orange prisms. Quinoline (B. 4)-carboxylic acid

CH:CH -- C.CH:CH CH:C(CO.H).Ö—N:CH Quinoline-o-carboxylic

acid. [1876].

Formation.-1. By heating o-amido-benzoic acid with o-nitro-benzoic acid, glycerin, and H₂SO₄ (Schlosser a. Skraup, M. 2, 518).—2. By heating its nitrile with conc. HClAq at 150° (Lellmann a. Reusch, B. 22, 1392).—3. Together with the (B. 1)-isomeride, by heating quinoline B. 1,4)-di-carboxylic acid (Skraup a. Brunner, M. 7, 153).

Properties.—Needles, m. sol. cold water and alcohol. FeSO, gives a purple colour in neutral solutions. - HA'HCl : yellowish prisms. -H,A',HCl: golden prisms. - H,A',H,PtCl,: orange needles.—CaHA',: needles.—AgA': pp.

Methylo-iodide HA'MeI. Slender yellow needles (La Coste, B. 15, 196). Moist Ag₂O forms a solution of the hydroxide which reproduces HA' on evaporation.

Nitrile C₁₀H₄N₂. [84°]. Needles. Yields B'₂H₄PtCl₆, crystallising in orange needles.

Quinoline (Py. 1)-carboxylic acid C₆H₄ < $\stackrel{C(CO_2H)}{N}$: $\stackrel{CH}{=}$ Cinchoninic ac Cinchoninic acid. [254°]. Formation.—1. By exidation of einchonine,

cinchonidine, cinchotenidine, cinchotenine, cinchonitine, and cincholepidine (Weidel, A. 173, 84; M. 3, 79; Skraup, A. 201, 303; M. 2, 601; Forst a. Böhringer, B. 14, 436; Schniderschnitsch, M. 10, 55).—2. By oxidation of (Py. 1)-methyl-quinoline (lepidine) (Hoogewerff a. Van Dorp, R. T. C. 2, 11).

Preparation.—By boiling cinchonine with nitric soid of S.G. 1-3 (Königs, B. 12, 97; Claus,

B. 18, 362).

Properties.—Slender needles (containing aq) (Skraup) or triclinic crystals (containing 2aq) a:b:c=.746:1:.817; $\alpha=81^{\circ}$ 37'; $\beta=120^{\circ}$ 2'; $\gamma = 86^{\circ} 37'$ (Muthmann a. Nef, B. 20, 637). Βv slow evaporation of an aqueous solution it is obtained in monoclinic forms (containing 2aq), a: $b:c=276:1: \cdot 583$; $\beta=98^{\circ}3'$ (Ditscheiner, A. 173, 84; Claus, B. 20, 1604). V. sl. sol. water and alcohol, insol. ether. Yields quinoline on distillation with lime, and oxy-quinoline carboxylic acid on fusion with potash. On heating with conc. H_2SO_4 it yields quinoline p-sulphonic acid (Georgievitch, M. 8, 644). Sulphuric acid and P₂O₅ at 175° gives an (a)-sulphonic acid C₅H₂N(SO₃H)(CO₂H), which forms very bitter triclinic crystals (containing aq), nearly insol. cold water, giving the salts (NH4)2A" 2aq, CaA" 2½aq, BaA" 3aq, PbA" aq, and CuA" aq. H₂SO, and P₂O₅ at 260° give a (β)-sulphonic acid C₁₀H₂NSO₅ crystallising in bitter needles (containing 2aq), m. sol. cold water, forming the salts (NH₄)HA"2aq, BaA"aq, and PbA" 4aq (Weidel, M. 1, 845; 2, 565; Georgievitch, M. 8, 644).—KMnO, forms pyridine tricarboxylic acid.

Salts.—HA'HCl.—(HA')₂H₂PtCl₃: triclinic prisms.—HA'HClICl 2aq (Dittmar, B. 18, 1618). —HA'HNO₃.—(HA')₂H₂SO₄: long prisms.— KA' ½aq.—CaA', 1½aq: prisms (Weidel).—CaA', (Skraup).—CuA', dark-violet plates, sl. sol. hot water.—AgA': crystalline pp.

Dibromide C, H, NO, Br, [188°]. Long red needles, decomposed by boiling water (Claus,

B. 18, 1307).

Di-iodide C10H7NO2I2. [c. 242°]. Steel-

blue tables, nearly insol. cold water.

Ethyl ether EtA'. [13°]. Trimetric crystals, v. sl. sol. water, v. sol. ether, sl. sol. alcohol (Van der Kolf a. Van Leent, R. T. C. 8, 217).—EtA'HgCl₂. [153°]. Small needles.— (EtA')₂H₂PtCl₈. [204°]. Orange-yellow needles.

Amide C₁₀H₈N₂O. [181°]. Needles, v. e. sol. hot water.—B'2H2PtCl₈. [250°-255°].

Benzylo-bromide C, H, NO, C, H, Br. [130°]. Silky needles, v. sol. water and alcohol, insol. ether. By heating the aqueous solution or by treatment with alkalis, it is converted, by loss of HBr, into the anhydride $\overset{C_0H_0N.O_7H_7}{CO-O}$, which crystallises in dimetric tables (containing 3aq) [84°] and has a bitter taste. This anhydride is converted by alkalis into C.H.N(C,H.)CO.H [218°] (Claus a. Muchall, B. 18, 863, 1810).

Tetrahydride C10H11NO2. Formed from cinchonic acid, tin, and HClAq (Weidel, M. 2, 29; 8, 61). Yields a nitrosamine [137°]. -B'HCl $1\frac{1}{2}$ aq: monoclinic crystals; a:b:c=932:1:1.942; $\beta = 90^{\circ}$ 41'. Heated with H₂SO, it gives disulpho-cinchonic acid and methylquinoline (cincholepidine). -B',H,PtCl.

Acetyl derivative of the Tetrahydride, H₁₀AcNO₂. [164°]. Trimetric crystals; C₁₀H₁₀AcNO₂. [164°]. Trimetric crystals; a:b:c = 848:1: 569. Sl. sol. cold water and alcohol.—CaA', 2aq: minute needles.

Quinoline (Py. 2)-carboxylic acid

C,H,<N=CH:C.CO,H. [273°].

Formation.—1. By heating acridinic acid at 125° (Graebe a. Caro, B. 13, 100). -2. By oxidation of (Py. 2)-ethyl- (or methyl-) quinoline (Riedel, B. 16, 1609; Doebner a. Miller, B. 18. 1643).

Properties.—Slender needles, sol. alcohol, sl. sol. cold water. Dissolves in acids and alkalis. Oxidised by KMnO, to pyridine tricarboxylic

Salts.—HA'HCl: needles, v. sol. water.— Picrate: [216°].—CuA'2.—AgA': small prisms. -(HA'), H, PtCl, : orange needles.

Quinoline (Py. 3)-carboxylic acid

C₆H₄ CH:CH Quinaldinic acid. [156°]. Formed from (Py. 3)-methyl-quinoline by oxidation with chromic acid mixture (Doebner a. Miller, B. 16, 2472). Got also by oxidation of (a)-diquinolyl (Weidel a. Strache, M. 7, 300). Needles (containing 2aq), v. sol. hot water, acids, and alkalis.—CuA'₂2aq.—CaA'₂.—AgA': amorphous precipitate.—AgHA'₂HNO, aq: silky needles.—HA'HCl aq.—(HA')₂H₂PtCl₆2aq.— (HA')2H2Or2O,: sparingly soluble red crystals.

Quinoline (B. 1,4)-di-carboxylic acid ÇH:C(CO₂H).C.CH:CH [270°]. Formed by $CH:C(CO_2H).C.N = CH$ heating amido-terephthalic acid with glycerin, H.SO₄, and o-nitro-phenol (Skraup a. Brunner, M. 7, 147). Long needles (containing 2aq), v. sl. sol. cold water. Salts.—HA'HCl 11aq.-(HA')₂H₂PtCl₈.—Cu₃A'₂(HO)₂ aq: bluish pp. Quinoline (Py. 2,3)-dicarboxylic acid

C₆H₄ CH:C.CO₂H₂. Acridinic acid. Formed by oxidation of acridine with KMnO, (Graebe a. Caro, B. 13, 99). Long needles (containing 2aq). sl. sol. cold water, v. sol. alcohol. Does not combine with acids. Decomposed by heat into

CO₂ and a mono-carboxylic acid [275°].

Quinoline (Py. 1,3)-dicarboxylic acid

C₄H₄ C(CO₂H):CH

C.CO₂H. [246°]. Former C.CO₂H. [246°]. Formed from (Py. 8)-styryl-cinchoninic acid by oxidation with alkaline KMnO₄ (Doebner a. Peters, B. 22, 3009). Needles, sl. sol. cold water and alcohol.-CuA" aq.—Ag₂A": white pp

Quinoline (a)-dicarboxylic acid

C₂H₂N(CO₂H)₂. [270°]. Formed by saponifying its nitrile. Slender needles (containing aq), sl. sol. alcohol and ether.

Nitrile C, H, NCy2. $[222^{\circ}].$ Formed by heating potassium quinoline (a)-disulphonate with KCy (La Coste a. Valeur, B. 20, 99). Needles (from alcohol), insol. Aq, v. sol. alkalis.

Quinoline (B. 2, Py. 3)-dicarboxylic acid CO₂H.C=CH.C.CH:CH [275°-280°].

OH:CH.C.N=C.CO₂H. [275°-280°]. Formed by oxidation of benzylidene-quinoline (B. 2)-carboxylic acid (Von Miller, B. 28, 2261). References .- Bromo-, NITRO-, and

QUINOLINE CARBOXYLIC ACIDS.

QUINOLINE (Py. 8)-CARBOXYLIC ALDE-HYDE C.H.N.CHO. [71°]. Formed by oxida-

tion of (Py. 3)-quinolyl-acrylic acid with KMnO₄ (Miller a. Spady, B. 18, 8404; 19, 130). Monoclinic plates (from ligroin), sl. sol. water. Reduces ammoniacal AgNO₃. Yields a phenyl-hydrazide C₁₈H₁₃N₂ [198°] crystallising in yellow plates.

QUINOLINE HYDRIDES.

Folymeride of quinoline dihydride C₁₈H₁₈N₂. [162°]. Formed by reducing quinoline with zinc-dust and HOAc, zinc-dust and NH, zinc and HClAq, or alcohol and sodium-amalgam (Königs, B. 12, 101, 252; 14, 99; Wyschnegradsky, B. 12, 1461; Lellmann, B. 22, 1337). Amorphous powder. Its solution in conc. HClAq is ppd. by water and by NaOAc. Yields a nitrosamine.

Quinoline dihydride (?). (220°-226°). Found in small quantity among the products of the action of KOH on quinoline (O. de Coninck, C. R. 94, 87).—B'₂H₂PtCl₆.

Quinoline tetrahydride C.H.IN i.e.

C₈H₄ CH₂, CH₂. [c. 10°]. (240° uncor.).

Formation.—1. By reduction of quinoline with zinc or tin and HClAq (Wyschnegradsky, Bl. [2] 34, 339; B. 12, 1481; 13, 2400), with zinc-dust and NH,Aq, with alcohol and sodiumamalgam (Königs, B. 14, 99; 16, 727), or with sodium (Weidel a. Glaser, M. 7, 328).—2. From carbostyril, alcohol, and Na (Knorr a. Klotz, B. 19, 3302).

Properties.—Needles, sol. water, volatile with steam. Secondary base. ICl forms in presence of HCl an amorphous pp. C₀H_{*}L_{*}N (Dittmar, B.

18, 1619).

Reactions.—1. The vapour passed through a red-hot tube yields indole and quinoline.—2. Chromic acid mixture yields quinoline.—3. Nitro-benzene yields quinoline. p-Chloro-nitrobensene at 200° does the same (Lellmann a. Reusch, B. 22, 1889).—4. HI and P reduce it to C_0H_{18} , an oil (147° at 720 mm.) (Bamberger a. Lengfeld, B. 23, 1158).—5. Br forms oily CoH10BrN and CoH9Br2N.—6. Bensene sulphonic chloride and NaOH form C₉H₁₀N.SO₂C₈N₈ [67°], overstallising in plates, sl. sol. alcohol, insol. water (Schotten a. Schlömann, B. 24, 8695).—7. Methyl chloroformate gives C, H, N.CO, Me [c. 85°], which may be oxidised by KMnO, in the cold to an acid [156°] and at 100° to a compound [175°] crystallising in red needles.—8. Diazobenzene chloride and aqueous NaOAc form C.H. N.N. C.H., which is split up by dilute H₂SO₄ into phenol and quinoline tetrahydride.-9. Potassium cyanate acting on the hydrochloride forms C₀H₁₀N.CO.NH₂ [150·5°], crystallising in needles, sl. sol. cold water. It is converted by conc. H2SO, and HNO, at 0° into $C_9H_8(NO_2)_2N.CO.NH_2$ [191°], sl. sol. alcohol, and C_pH_u(NO₂)₂N [161°], v. sol. alcohol, which forms a nitrosamine [127°] (Thomas, R. T. C. 10, 148).—10. Bensoic aldehyde and ZnCl₂ yield CHPh(C₉H₁₀N)₂ [153°], while p-nitro-benzoic

aldehyde forms a (Einhorn, B. 19, 1243).
Salts. — B'HCl. [181°]. — B',H,PtCl. — B'H,SO,. [187°]. Prisms (from alcohol) or tables (from water).

Acetyl derivative C₂H_{1e}NAc. (295°). Yields CO₂H.CO.NH.C_eH₄.CO₂H [1:2] on oxidation by KNnO₄. Bensoyl derivative C.H. 10NBz. [75°]. Tables (from alcohol). Oxidised by KMnO₄ to benzoyl-isatic acid (Schotten, B. 24, 772).

Nitrosamine C₀H₁₀N(NO). Oil. Converted by alcoholic HCl into the isomeric (B. 2)-nitroso-quinoline tetrahydride [134°], which separates from benzene in steel-blue crystals, converted by NaNO₂ and HOAc into C.H.N.O. [98°] (Ziegler, B. 21, 864). Phenyl-hydrazine forms C₂H₁₀N₂O.PhN₂H₁ [120°]. Zinc-dust and HOAc reduce the nitrosamine to the hydrazine from ligroin, and forms B'₂H₂SO₄2aq, crystallising in yellow plates. The hydrazine is oxidised in ethereal solution by ppd. HgO to the tetrazone C₁₈H₂₀N₄, which forms colourless needles [160°], sol. ether, sl. sol. alcohol.

Methylo-chloride v. Hydrochloride of

v-METHYL-QUINOLINE TETRAHYDRIDE.

Ethylo-iodide B'EtI. Changed by alkalis into rethyl-quinoline tetrahydride, which is a liquid (254°-258°) yielding (C₀H₁₀NEt)₂H₂CdCl₄, [105°], and (C₀H₁₀NEt)₂H₂PtCl₆ [c. 160°] (Wyschnegradsky, B. 13, 2400; Claus a. Stegelitz, B. 17, 1329).

Quinoline tetrahydride C₉H₁₁N. (213°). Formed by distilling cinchonine (1 pt.) with KOH (3 pts.) (O. de Coninck, C. R. 94, 87; A. Ch. [5] 21, 478). Got also by distilling brucine with potash (De Coninck, C. R. 99, 1077). Slightly hygroscopic oil, v. sol. acids. Reduces FeCl₂. May be oxidised to quinoline.—B'HCl: deliquescent needles, v. sol. water.—B'₂H₂PtCl₄: —B'₂H₂PtCl₄: pale orange plates.

Quinoline hexahydride C₂H₁₃N. (227°) at 720 mm. A product of the action of HI and P at 230° on the hydrochloride of quinoline tetrahydride (Bamberger a. Lengfeld, B. 23, 1155). Liquid, turning brown in air, m. sol. hot water. FeCl₂ gives a brown colour on warming. Diazo benzene sulphonic acid gives a light-red colouring matter. Phosphomolybdic acid gives a white pp. Potassio-mercuric iodide gives no pp. NaNO₂ in acid solution gives an oily pp. of the nitrosamine.—B'HCl. [170°]. Silky needles. B'HBr. Long silky needles.

Quinoline decahydride C.H., N i.e.

CH₂·CH₂·CH₂·CH₂·CH₂·CH₂·CH₂·CH₃·CH₄·CH₂·CH₂·CH₂·CH₄·CH₂·CH₂·CH₄·CH₂·CH₂·CH₄·CH₄·CH₂·CH₄·CH₄·CH₂·CH₄·

Reactions.—1. Phenyl-cyanats forms, in ethereal solution, C₈H₁₈N.CO.NHPh, crystallising in needles [148°].—2. Phenyl thiocarbimids in ether forms C₈H₁₈N.CS.NHPh, crystallising from chloroform in tables [1345°].—3. An ethereal solution of CS₂ gives C₈H₁₈N.CS.SNH₂C₈H₁₈, crystallising in needles [120°], not decomposed by boiling alcohol, but converted by iodine into

(C_pH_{1,7}N)HI and S₂(CS.NC_pH_{1,8})₂, which crystallises in colourless needles [81°], sol. alcohol. Salts.—B'HCl. [276°]. Small tables.—

Salts.—B'HCl. [276°]. Small tables.— B'HI. [253°]. Silky needles.—B'HAuCl. [96°]. Golden needles.—B'₂H₂PtCl₈. [207·5°]. Acetyl derivative C₂H₁₂NAc. Oil.

Benzoyl derivative C.H., NBz. [44°]. ISOQUINOLINE TETRAHYDRIDE C.H., N. (282°). Formed by the action of tin and HCl upon isoquinoline (Hoogewerff a. Van Dorp, R. T. C. 5, 305). Liquid, smelling like isoquinoline, sl. sol. water. Absorbs CO2 from the air. forming a crystalline carbonate. The aqueous solution reduces AgNO2. Yields an oily nitrosamine.—B'HCl. [196°]. Tables, sol. alcohol. Not hygroscopic.—B'H, SO4: sol. alcohol.—B'2H, PtCl2. [232°]. Reddish-yellow tables, decomposed by hot water.

QUINOLINE-HYDROQUINONE v. DI-0XY-QUINOLINE.

QUINOLINE-IODOFORM v. QUINOLINE, Reaction 20.

QUINOLINE-QUINONE C.H.NO. i.e.

CH.CO.C.CH:CH. Formed by the action of cold chromic acid mixture on (B. 4,1)-oxy-amidoquinoline (Fischer a. Renouf, B. 17, 1644). Flat needles with green reflex (from alcohol), decomposing at 110°-120°. Its salts are decomposed by water. Reduced by sulphurous acid to dioxy-quinoline. Aniline in alcoholic solution forms, on boiling, the dianilide C₁H₁₀N₂O₂ crystallising from dilute alcohol in small copperred plates with green reflex [c. 190°], forming a violet solution in HClAq.

Mono-oxim CH.C(NOH).C.CH:CH CH.CO — C.N=CH

Formed by the action of NaNO₂ and HClAq on (B. 4)-oxy-quinoline (Von Kostanecki, B. 24, 152). Needles (from alcohol). Gradually blackens below 220°, decomposing completely at 245°.

CH.C(NOH).C.CH:CH

Di-oxim CH.C(NOH).C.N=CH

by adding hydroxylamine hydrochloride to an aqueous solution of the mono-oxim (Von Kostanecki a. Reicher, B. 24, 156). Minute crystals. Ac₂O yields a crystalline di-acetyl derivative decomposing below 160° with evolution of gas. Dyes cotton mordanted with iron salts greenish-black.

Quinoline-o-quinone C₀H₂NO₂. Formed by oxidising (B. 2)-oxy-(B. 1 or 3)-amido-quinoline with FeCl₃ in acid solution (Mathëus, B. 21, 1887). Unstable.—B'HCl: long reddish-yellow needles. Decomposed by Na₂CO₃, forming a brown solution. The sulphate forms yellow plates.

Mono-oxim C₂H₄N(OH)(NOH). Formed from p-oxy-quinoline, NaNO₂, and HCl (M.). Golden needles (from HOAc), sl. sol. ether.

Di-oxim C₂H₂N(NOH)₂. Needles, decomposing at 190° (Von Kostanecki a. Reicher, B. 24, 158). Converted by conc. KOHAq into the anhydride C₂H₄N(N₂O), which forms colourless needles, readily volatile with steam.

QUINOLINE (B. 1)-SULPHONIC ACID CH:C(SO,H),C.CH:CH A product of the culphonation of quinoline (O. Fischer, B. 15, 658, 1979; 20, 731; La Coste a. Valeur, B. 20,

97). Formed also from amido-benzene m-sulphonic acid, nitrobenzene, glycerin, and H_2SO_4 at 160° (Lellmann a. Lange, B. 20, 1446). Monoclinic needles (containing aq). Yields oxy-quinoline [224°] by potash-fusion.—CaA' $_2$ 5aq: needles, v. e. sol. water. According to Lellmann, there exists an isomeric quinoline (B. 1)-sulphonic acid, also got by sulphonation of quinoline, with H_2SO_4 (containing 50 p.c. SO_3 extra) at 260° (Lellmann, B. 20, 2172). This ψ -acid is converted by conc. H_2SO_4 at 250° -300° into the (B. 2)-acid (Lellmann, B. 22, 1391). The ψ -acid forms anhydrous crystals.

Tetrahydride C_pH₁₀N.SO₃H aq. [315°]. Formed by reducing the above acid (B. 4)-bromoquinoline (B. 1)-sulphonic acid and also (B. 4)-oxy-quinoline (B. 1)-sulphonic acid with tin and HClAq (Lellmann a. Lange, B. 20, 3087; Claus, J. pr. [2] 42, 344). Trimetric crystals (from dilute solutions) or monoclinic crystals (from conc. solutions). FeCl₃ gives a brown colour passing through purple to grass-green on warming.

Quinoline (B. 2)-sulphonic scid SO₃H.C=CH.C.CH:CH Formed

CH:CH:C.N=CH Formed by heating a mixture of amido-benzene-p-sulphonic acid, glycerin, nitrobenzene, and H₂SO₄ (Happ, B. 17, 191). Formed also (to the extent of 70 p.c.) by heating quinoline with conc. H₂SO₄ at 240°-300° and by heating the (B. 1)- or (β 4)- acid with conc. H₂SO₄ at 240°-300° (Georgievitch, M. 8, 578, 639). Monoclinic needles (containing 1½aq), sl. sol. alcohol and cold water. Not melted at 260°-Potash-fusion yields (B. 2)-oxy-quinoline [193°]. Bromine-water gives a yellow pp. of di-bromo-quinoline [126°] and at 100° Br (3 mols.) forms tri-bromo-quinoline [170°] (Claus a. Küttner, B. 19, 2884). Heating with KCy gives C₂H₄N.Cy [131°] which on saponification yields the carboxylic acid [288°].

Salts. – KA' (dried at 120°). Tables.—

Salts.—KA' (dried at 120°). Tables.—BaA'₂ (dried at 120°).—AgA': slender needles.—Ag₂HA'₃: needles.— (HA')₁₈K₄I₉6aq: lustrous green crystals.

Ethyl ether EtA'2aq. Crystals, v. sol. water (Claus a. Happ, B. 18, 366).—EtA'KI₃: unstable brown needles, v. sol. water (Claus a. Stegelitz, B. 19, 921).—EtA'KBr₃.—EtA'HgCl₂.

Benzyl ether C₇H,A' 2aq. Monoclinic

Benzyl ether C,H,A'2aq. Monoclinic crystals.—C,H,A'KI₂: needles with blue reflex.

Quinoline (B. 8)-sulphonic acid CH:CH.C.CH:CH

CH:CH.C.CH:CH
SO₃H.C=CH.C.N=CH
SO₅H.C=CH.C.N=CH
by heating quinoline with H₂SO₄ (containing 15
p.c. SO₃ extra) at 130° (Claus, J. pr. [2] 37,
260). Small needles, v. e. sol. water, v. sol.
alcohol. Potash-fusion yields oxy-quinoline
[165°-200°]. Bromine forms di-bromo-quinoline
[255°] and tri-bromo-quinoline [199°].—NaA'3aq.
KA' 2aq.—CaA'₂ 4aq.—BaA'₂ 4aq.—PbA'₃.—
ChA'₂ 2ag: dark-green ervstalline nowder.

CuA', 2aq: dark-green crystalline powder. Ethyl ether EtA'. [275°]. Needles (from alcohol).

Chloride C,H,N.SO,Cl. Viscid mass. Amide C,H,N.SO,NH, [119°]. Crystalline. Quinoline (B. 4)-sulphonic acid

CH:CH — C.CH:CH Quinoline o-sulphonic acid. Formed, together with a smaller quantity of the p-acid, by heating quinoline with fuming

H₂SO, at 170° (O. Fischer a. Bedall, B. 15, 683, 1979; 20, 731; Riemerschmied, B. 16, 721; La Coste a. Valeur, B. 20, 95; Claus, B. 19, 925, 2882; J. pr. [2] 37, 260). Formed also by heating quinoline with conc. H,SO, at 220° (Georgievitch, M. 8, 641). Monoclinic prisms, v. sl. sol. water. Oxidised by KMnO, to pyridine dicarboxylic (quinolinic) acid (Fischer, B. 17, 755) and a small quantity of (1,2,3)-sulpho-amidobenzoic acid (Zürcher, B. 21, 180). Changed by conc. H_2SO_4 at 250° to the (B. 2)-isomeride (Georgievitch, M. 8, 578). Bromine-water forms a yellow insoluble pp. and, at 100°, tri-bromoquinoline [198°].

Salts.—NaA' 5aq (Spies, J. pr. [2] 37, 265). -KA' 2aq.—CaA'2 9aq. Monoclinic crystals. PbA'2. CuA'2 2aq: small green needles.

Ethyl ether EtA'. [66°]. Needles (from chloroform). Decomposed by boiling water.

Chloride C,H,NSO,Cl. [124°]. Needles. [184°]. Amide C₉H₈N.SO₂NH₂. Needles (from water) (Hoogewerff a. van Dorp, R. T. C. 8, 184). Converted by Br and KOHAq into C₉H₆NSO₂NBrK 2aq, crystallising in yellow needles, m. sol. water, and converted by HOAc into C9H6NSO2NBrH, crystallising from acetone in needles, yielding (C.H.NSO2NBr)2Ba 2aq.

Quinoline (a)-disulphonic acid $C_0H_1(SO_3H)_2N$. Formed, together with the (β)isomeride, by heating quinoline (B. 4)-sulphonic acid with fuming H₂SO₄ at 250° (La Coste a. Valeur, B. 19, 995; 20, 98). Yellow needles (containing Saq), v. sol. water, insol. alcohol.-K₂A"3½aq: satiny plates, v. e. sol. water.—

BaA" 3aq. S. 4.08 at 15°.

Quinoline (β) -disulphonic acid $C_9H_4N(SO_9H)_2$. Formed as above (La Coste a. Valeur, B. 19, 998; 20, 3199). Needles (containing 1 aq), v. sol. water, insol. alcohol. K2A" aq : crusts, sl. sol. cold water.—BaA" 2aq : white powder, sl. sol. cold water (difference from (a)- isomeride).

References. — Bromo- and Oxy-Quinoline

SULPHONIC ACIDS.

ISOQUINOLINE SULPHONIC ACID

 $C_9H_6N(SO_9H)$. Formed by heating isoquinoline (1 pt.) with fuming H_2SO_4 (2 pts.) at 100° (Hoogewerff a. van Dorp, R. T. C. 5, 305).— , 9aq. Needles, v. sol. water.

QUINOLINIC ACID v. Pyridine dicarb-OXYLIC ACID. The name is also given by Dewar (Pr. 30, 164) to an acid $C_0H_0NO_3$ [143°] formed in small quantity in the oxidation of quinoline (derived from cinchonine) by boiling aqueous KMnO. (a)-DIQUINOLYL C₁₈H₁₂N₂ i.e.

C_eH₄ CH:CHCH:CH N=C-C-N>C_eH₄. Diquinoline.

[176]. (above 400°). V.D. 8.73 (calc. 8.86). Formation.—1. By heating quinoline with sodium at 192° (Weidel, M. 2, 491; 7, 327).—2. In small quantity (8 p.c.) by passing dry oxygen into a mixture of quinoline and quinoline hydrochloride in presence of platinum-black (Weidel, M. 8, 120).

Properties. — Monoclinic plates, a:b:c= 1.37:1:1.32; $\beta = 109^{\circ} 58'$. Insol. water, sol. alcohol, ether, and benzene. Oxidised by KMnO. and HOAc to cyclothraustic (C1,H12N2O2), quinoline (Py. 3)-carboxylic, oxyisocinchomeronic (oxy-pyridine dicarboxylic), and o-amido-benzoic acids (Weidel a. Strache, M. 7, 280, 306; 8, Alkaline KMnO, forms isocinchomeronic, o-amido-benzoic, and pyridanthrilic (C15H10N2O, MeOH and H2SO, form B"MeSO,H (Ostermayer, B. 18, 333, 597).

Salts. — B"H₂Cl₂ 4aq: white needles.— B"H₂PtCl₃ aq.—B"HAuCl₄ 2aq.—B"H₂SO₄ aq. Methylo-chloride B"Me2Cl2 6aq: needles.

-B"Me,Cl,2ICl. [238°]. Lemon-yellow pp. Methylo-iodide B"MeI. [283°].

Ethylo-iodide B"EtI. Yellow needles (from water), sl. sol. water (Roser, B. 17, 2769).

Sulphonic acid C₁₈H₁₁N₂(SO₂H). Formed by heating diquinolyl with H.SO₄ (containing 5 p.c. SO, extra) (Weidel a. Gläser, M. 7, 308). Minute needles, almost insol. hot water, sol. conc. HClAq and H2SO4. Converted by potashfusion into oxy-diquinolyl [208°].—KA'2aq: needles, v. e. sol. water.—CuA', 2aq.

(a)-Disulphonic acid $C_{18}H_{10}(SO_3H)_2N_2$.

Formed by heating diquinolyl (10 pts.) with H₂SO₄ (11 pts.) and SO₃ (84 pts.) at 170° (Weidel, M. 2, 503). Minute needles, sl. sol. water. Yields, by potash-fusion, di-oxy-di-quinolyl [239°].—K₂A" 5aq: needles.—CuA" 6aq: bluish-green crystals, nearly insol. water.

(β)-Disulphonic acid C18H10(SO3H)2N Formed, together with the mono-sulphonic acid, by heating (a)-diquinolyl (100 pts.) with H.SO (332 pts.) and SO. (18 pts.) for 20 hours at 190° (Weidel a. Gläser, M. 7, 322). Small needles, converted by potash-fusion into di-oxy-di-quinolyl [above 305°].— K_2A'' (dried at 120°): crystalline powder.—CuA'' (dried at 120°).

(β)-Diquinolyl

CH:CH.C.CH:CH:CH C₆H₄. [192°].

Formation.—1. By heating quinoline with BzCl at 250° (Japp a. Graham, C. J. 39, 174).— 2. In very small quantity by distilling cinchoninio acid with lime (Weidel, M. 2, 501). -3. By passing quinoline through a red-hot tube (Zimmermann a. Müller, B. 17, 1965).—4. By distilling quinoline (B. 4)-sulphonic acid; the yield being 10 p.c. (O. Fischer a. Loo, B. 17, 1899; 19,

Properties.—Iridescent plates (by sublimation) or colourless prisms (from ether), sol. hot benzene, alcohol, and ether, insol. water. Weak base, the salts being decomposed by water. Oxidised by CrO, and HOAc to quinoline (B. 3)carboxylic acid. CrO₃ and H₂SO₄ give pyridylquinoline carboxylic acid C₆H₄N.C₅H₄N.O₂H [273°]. Bromine forms C₁₈H₁₂N₂Br₄ [192°].

Salts.—B"H₂SO₄: sparingly soluble concentric needles.—B"H₂PtCl₅: granular pp.

Ethylo-iodide B"EtI. Very unstable red crystals, decomposed by hot water.

C18H10(SO3H)2N2 Disulphonic acidFormed by heating the base (1 pt.) with conc. H₂SO₄ (10 pts.) at 180°. V. e. sol. water.— K,A" Baq: prisms.

(γ)-Diquinolyl ÇH:N—Ç.CH:ÇH ÇH:CH.Ç.N=ÇH Di-p-CH:CH.C.CH:C-C=CH.C.CH:CH

quinolyl. [178°].

Formation.—1. By boiling a mixture of benzidine, nitrobenzene, glycerin, and H.SO. (Roser, B. 17, 1817, 2767; O. W. Fischer, M. 5, 418; Colson, C. R. 108, 677). A better yield is got by using o- or p- nitrophenol instead of nitrobenzene (Ostermayer a. Henrichsen, B. 17, 2444). When crude benzidine is used there is also formed a base C₂H₁₈N₂O₃, which yields B'2SnCl₂ [185°] and B"H₂PtCl₃2aq (Colson, C. R. 107, 1008; 108, 677).—2. By heating azobenzene with glycerin and H₂SO₄ (Claus a. Stegelitz, B. 17, 2380).

Properties .- Monoclinic crystals, insol. water, sl. sol. ether, v. sol. hot alcohol and benzene. MeOH and H2SO4 at 180° form B"(MeHSO4)22aq, crystallising in needles. Bromine in alcohol forms unstable C₁₈H₁₂N₂Br₂ and C₁₈H₁₂N₂Br₄, converted by HClAq at 190° into C. HijBrN, [150°-

155°7

Salts. - B"H2Cl2 4aq: white needles. B"HAuCl, 2aq: slender needles.—B"H2SnCl,: colourless needles. -- B"H2PtCl4 (dried at 105°). B"H₂Cl₂2ICl: cheesy pp. (Dittmar, B. 18, 1618). — B"H₂SO₄ 3aq. — B"2H₂SO₄. — B"2H₂Cr₂O₇. —

B"(O_H₄(NO₂)₂OH: minute yellow prisms.

Methylo-iodides B"MeI. Crystals.—
B"Me₂I₂. [c. 300°]. Yellow needles, sol. water. Ethylo-iodide B"2Etl. [270°]. Needles. Disulphonic acid C₁₆H₁₀(SO₂H)₂N₂. White powder, insol. water and alcohol.— Na₂A" 5aq: small needles. The same, or an isomeric, disulphonic acid is got by heating benzidine disulphonic acid with o-nitro-phenol, glycerin, and H_2SO_4 . It crystallises in plates or needles, and yields K_2A'' aq.

(8)-Diquinolyl QH:N—C:CH:CH CH:CH:CH:CH:CH

ČH:CH.C.CH:Č--C.N=CH[148°]. Formed by heating o- p- di-amido-diphenyl hydrochloride with nitrobenzene, glycerin, and H.SO. (O. W. Fischer, M. 6, 546). Leaflets, insol. water, sol. hot alcohol and benzene. Bromine in presence of water forms orange needles of $C_{18}H_{12}N_2Br_4$.—B"H₂Cl₂: needles, v. e. sol. water.—B"H₂PtCl₃ aq: needlesh-yellow crys-

talline pp.—Picrate. (168°). Yellow needles.

Methylo-iodide B"MeI. [126°]. Needles.

Disulphonic acid C₁₈H₁₀(SO₂H)₂N₂. Foursided tables, not melted at 300°. Sl. sol. hot

water.—BaA" 3aq: needles, m. sol. hot water. (Py. 3, B. 1 or 3)-Diquinolyl $C_{1s}H_{12}N_2$. [159° Formed (7 pts.), together with the isomeride [115°], by heating a mixture of m-amido-(Py. 3)phenyl-quinoline (20 pts.), glycerin (70 pts.), o-nitro-phenol (15 pts.), and H₂SO₄ (60 pts.) (Miller a. Kinkelin, B. 18, 1910). Small monoclinic tables, sl. sol. ether, cold alcohol, and chloroform.—B"H₂Cl₂2aq: plates.—B"H₂PtCl₂. $-B''H_2SO_4$: easily soluble needles.— $B''C_6H_3N_4O_7$. [240°]. Crystalline powder.

Methylo-iodide B'MeI. [263°]. Needles. (Py. 3, B. 3 or 1)-Diquinolyl $C_{18}H_{12}N_2$. 5°]. Formed as above (M. a. K.). Lightyellow triclinic crystals, v. sol. alcohol, ether, and benzene, sl. sol. ligroin.—B"H2Cl2Saq: tables, v. e. sol. water. - B"H2PtCls: crystalline.

(Py. 3, B. 1)-Diquinolyl C.H. CH:CH CH.OH:CH.C.CH:CH [144°]. -C.N = CHFormed from p-amido-(Py. 3)-phenyl-quinoline, glycerin, nitro-benzene, and H.SO. (Weidel, M. 8, 140). It is also a product of the action of oxygen on a mixture of quinoline hydrochloride, aniline, and platinised asbestos at 180°. Monoelinic tables (from alcohol).—B"H_PtCls: crystalline powder.—B"HCl: yellow.—B"H2Cl2.

Methylo-iodide B"MeI aq. [282°]. Small needles (from water), sl. sol. boiling water.

Sulphonic acid C, H, (SO, H)N, soluble powder. Potash-fusion yields an oxy-

diquinolyl [187°]

(B., Py. 1)-Diquinolyl C₁₈H₁₈N₂. [122°]. Got by heating (a)-amido-(Py. 1)-phenyl-quinoline [150°] with glycerin, nitro-benzene, and H₂SO₄ (Koenigs a. Nef, B. 20, 632). Large crystals (from benzene). May be distilled. V. e. solalcohol, v. sl. sol. ether. Strong base. -B"H2PtCl5: granules, sol. conc. HClAq.

(B., Py. 1)-Diquinolyl $C_{18}H_{12}N_2$. [117°]-t by heating (β)-amido-(Py. 1)-phenyl-Got quinoline [198°] with glycerin, nitro-benzene, and H₂SO₄ (Koenigs a. Nef, B. 20, 634). Tables (from benzene), v. e. sol. alcohol. May be distilled. Strong base.—B"H_PtCl_a: ye needles.—Picrate. [248°]. Bulky flakes. (Py. 2.3).Diquinolyl yellow

 $C_0H_4 < CH_1O - C=N \\ C_0H_4 < CH_1OH CH_1CH > C_0H_4$. [176°]. Formed by the action of (Py. 3)-quinolyl-acetic aldehyde on o-amido-benzoic aldehyde in alcoholic solution (Carlier a. Einhorn, B. 23, 2895). Plates (from alcohol) .- Aurochloride: [248°]. -Platinochloride: [278°].-B"H2Cl2 4aq.

Methylo-iodide B"MeI.

Diquinolyl tetrahydride C₁₈H₁₈N₂. [118°]. Formed by heating (Py. 3)-chloro-quinoline with quinoline tetrahydride (Friedländer a. Weinberg, B. 18, 1533). Crystals, sl. sol. ligroïn, insol. water. May be distilled. HClAq forms a deep-yellow solution.

Reference. - OXYDIQUINOLYL. (Py. 3)-QUINOLYL-ACETIC ACID

CeH. CH: CH [275°]. Formed from the aldehyde by potash-fusion or by oxidation (Carlier a. Einhorn, B. 23, 2896). Formed also by oxidation of a-oxy-quinolyl-propionic acid with KMnO. Needles (from alcohol).—Hydrochloride: [243°]; needles. — Silver salt: white scales, yielding quinaldine on distilling with lime.

(Py. 3)-QUINOLYL-ACETIC ALDEHYDE C₁₁H_aNO i.s. C₂H_aN.CH₂.CHO. [104°]. Formed by careful oxidation of a-oxy-quinolyl-propionic

acid (Einhorn, B. 18, 3467; 19, 908). Crystals (from alcohol). Yields a phenyl-hydrazide [199°].—B'₂H₂PtCl₆2aq.—B'C₆H₂N₃O₇. [212°].

QUINOLYL-ACETONE v. ACETONYL-QUINOL-

(Py. 3)-QUINOLYL-ACETYLENE C.H. N.C.CH. Formed by distilling a solution of di-bromo-quinolyl-propionic acid with Na₂CO₂ (Carlier a. Einhorn, \hat{B} . 23, 2896). Volatile oil. Br forms C.H.N.CBr:CHBr.

(Py. 8)-QUINOLYL-ACRYLIC ACID

 $C_{12}H_{\bullet}NO_{2}$ i.e. $C_{\bullet}H_{\bullet} < \stackrel{CH:CH}{\sim} \stackrel{CH:CH:CH.CO_{2}H}{\sim} [193^{\circ}]_{\bullet}$ Formed by the action of boiling potassium carbonate solution upon 'quinolyl-acrylo-trichloride' C.H.N.CH:CH.CCl. aq [145°], which is got by heating (Py. 3)-methyl-quinoline with chloral at 110° (Miller a. Spady, B. 18, 3402, 3465; 19, 130; Einhorn, B. 19, 908). Plates. Oxidised by KMnO₄ to aldehydo-quinoline. HBr forms \$\beta\$-bromo-quinolyl-propionic acid. — *BaA'₂: needles, sl. sol. cold water.—*HA'HCl: needles...

QUINOLYL-ACRYLO-TRICHLORIDE

TRI-(B. 2)-QUINOLYL-CARBINOL

(C.H.N), C.OH. [108°]. Formed by oxidising tri-quinolyl-methane with H₂CrO₄ and HOAc (Noelting a. Schwarz, B. 24, 1608). Needles, v. sol. alcohol, sl. sol. ligroin.

(B. 3, Py. 8)-QUINOLYLENE DIACRYLIC

ACID ÇH:CH.Q.CH:ÇH

CO₂H.CH:CH.Ö=CH.Ö.N=C.CH:CH.CO₂H. [over 300°]. Formed by boiling tri-chloro-propenyl-quinolyl-acrylic acid with K₂CO₂Aq (Eckhardt, B. 22, 284). Sol. acids and alkalis. Insol.

water, sl. sol. hot alcohol.

DI-(B. 2)-QUINOLYL-ETHANE C₂₀H₁₈N₂ i.e. C₅H₈N.CH₂.CH₂.C₆H₈N. [124°]. Formed by heating di-p-amido-di-phenyl-ethane with nitrophenol, glycerin, and H₂SO₄ (Comey, B. 23, 1115). Hexagonal plates, insol. water, sl. sol. ether.—B"H₂Cl₂ 4aq: needles, v. sol. water.—B"H₂Cl₂.—B"H₂PtCl₃.—B"2HAuCl₄.—Sulphate: prisms, m. sol. cold water.

(Py. 3)-QUINOLYL-ETHYLENE $C_{11}H_9N$ i.e. C_6H_4 CH:CH Formed by boiling β -

bromo-quinolyl-propionic acid with conc. K,CO,Aq (Emhorn, A. 246, 172). Oil. — B'.H.PtCl. 4aq. [187°].—B'HAuCl.. [159°].

Formed by heating m-amido-(Py. 3)-styryl-quinoline with o-nitro-phenol, glycerin, and H₂SO₄ (Wartmann, B. 23, 3650). Oil, v. sol. alcohol, sl. sol. ether.

Methylo-iodide B'MeI 1½aq. [226°]. Golden needles, v. sol. hot water and alcohol.

Yields $C_{20}H_{14}N_2Br_2MeI$. [210°].

QUINOLYL-DI-ETHYL-PHENOL C, H, NO i.e. C, H, N.C, H, Et, OH. This is probably the constitution of apocinchene (Comstock a. Königs,

B. 20, 2674; v. vol. ii. p. 174).

(B. 4)-QUINOLYL-ĤYDRAZINE C₅H₅N₅ i.e. CH:CH.C.CH:CH.CH.CH
CH:N.—C.——C.N₂H³· [64°]. Formed from (B. 4)-amido-quinoline by diazotisation and treatment with SnCl₂ and HCl (Dufton, C. J. 59, 757; Böttinger, B. 24, 3277). Needles (from ether) or prisms (from ligroin). — B"H₂Cl₂: prisms. Converted by potassium cyanate and HCl into C₅H₅N.NH.NH.CO.NH₂ [235°], and by pyruvic acid into C₅H₅N.N₂H:CMe.CO₂H crystallising in orange needles, converted by boiling HClAq into quinindole (a)-carboxylic acid C₁₅H₁₀N₂O₂ [286°].

(B. 1)-Quinolyl-hydrazine [151°]. Formed in like manner from ana-amido-quinoline (Dufton, C. J. 61, 785). Yellow needles (from water). Yields B'HCl [248°] and is converted by cyanic acid into C.H., N.NH.NH.CO.NH. [255°], by benzoic aldehyde into C.H., N.NH.N:CHPh [194°], by acetone into C.H., N.NH.N:CMe. [140°] and by pyruvic acid into the acid C., H., N.NH.N:CMe. CO., H. [185°], whence boiling HClAq forms the corresponding quinindole carboxylic acid, which decomposes at about 300°.

DIQUINOLYLINE is DIQUINOLYL.

DI-(B. 2)-QUINOLYL KETONE (C.H.N).CO. [174°]. Formed from di-p-amido-benzophenone, glycerin, H.SO., and picric acid (Noelting a. Schware, B. 24, 1608). Needles, v. sol. alcohol,

sl. sol. hot water. — B"H₂Cl₂. [over 200°]. Needles, v. e. sol. water and alcohol.

(Py. 8)-QUINOLYL-MERCAPTAN

C_eH_s CH:CH [174°]. Formed by heating (Py. 3)-oxy-quinoline with P_sS_s at 145° (Roos, B. 21, 620). Yellow plates, sl. sol. hot water, v. sol. hot alcohol. Yellow dye. Combines with acids and bases. Does not react with hydroxylamine or phenyl-hydrazine.

Ethyl derivative C.H.N.SEt. Formed from quinolyl-mercaptan, NaOEt, and EtI. Oil, decomposed by distillation.—B'HI. [154°].—

B'₂H₂PtCl, aq. [c. 190°]. Octahedra. TRI-(B. 2)-QUINOLYL-METHANE

CH(C_pH_eN)₃. [202°]. Formed from p-rosaniline, glycerin, H_eSO₄, and picric acid at 150° (Noelting a. Schwarz, B. 24, 1606). Needles (containing EtOH), melting at 98°. Melts at 202° when anhydrous. — B"'3HCl: plates, sol. water. — B"C_eH₄N₂O₇: yellow needles. — B"'4H₁Pt₂Cl₂: yellow crystals.—B"'8MeI. [266°].

(B. 2)-QUINOLYL-METHENYL-AMIDOXIM C(NOH)(NH₂),C=CH.C.CH:CH CH:CH.C.N=CH. [105°]. Formed

from C₀H₂N.Cy, hydroxylamine hydrochloride, Na₂CO₃, and dilute alcohol (Biedermann, B. 22, 2761). Yellowish needles, m. sol. hot water. Fehling's solution gives a greyish-green pp. FeCl₃ gives a deep-red colour. AgNO₃ gives a white pp. yielding a mirror.

Reactions.—1. Phthalic anhydride on warming forms C₉H₈N.C $\stackrel{\text{N.O}}{\sim}$ C.C₈H₄.CO₂H [208°].

2. Ac₂O yields $C_{\nu}H_{\nu}N.C < \stackrel{N.O}{N} > CMe$ [175°], crystallising in needles.—3. The hydrochloride is converted by potassium oyanate into $C_{\nu}H_{\nu}N.C(NOH).NH.CO.NH_2$ [165°].—4. Chloroformic ether forms $C_{\nu}H_{\nu}N.C(NH_2):NO.CO_2Et$ [97°], which is converted by NaOHAq into $C_{\nu}H_{\nu}N.C < \stackrel{N.O}{N} > O$ [155°].

Salts.—B'HCl: needles.—B'₂H₂PtCl₃.

Acetyl derivative C₂H₃N.C(NOAc).NH₄.

[115°]. Formed by means of AcCl in the cold.

Needles, insol. cold water, sl. sol. ether.

Ethyl ether C₂H₃N.C(NOEt).NH₂.

[85°].

Ethyl ether C.H.N.C(NOEt).NH. [857]. Formed from the amidoxim, NaOEt, and Etl.

Needles, v. sol. alcohol and hot water.

(Py. 3)-QUINOLYL-PROPIONIC ACID C₅H₄N.CH₂.CH₂.CO₂H. [116°]. Formed by reducing quinolyl-acrylic acid with sodium-amalgam (Carlier s. Einhorn, B. 23, 2896).

Ethyl ether EtA'. [116°]. Needles. Reference.—Oxy-quinolyl-propionic acid.

DI-(Py. 3)-QUINOLYL DISULPHIDE $(C_nH_nN)_2S_2$. [137°]. Formed by oxidising quinolyl-mercaptan with H_iO_2 in dilute alcohol (Roos, B. 21, 622). Plates (from alcohol), insolwater and alkalis.

QUINONE C₆H₄O₂ i.e. CH.CO.CH Bensoquinone. Mol. w. 108. [116°]. V.D. (H = 1) 53.4 (calc. 54) (Hofmann, B. 3, 583). H.C. 659,020. H.F. 45,200 (from diamond) (Berthelot a. Recoura, Bl. [2] 48, 699; A. Ch. [6] 13, 312, 335; C. R. 104, 1572).

Formation.—1. By oxidation of quinic acid with MnO₂ (4 pts.), H₂SO₄ (1 pt.), and water (\frac{1}{2} pt.) (Woskresensky, A. 27, 268).—2. By

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oxidation of hydroquinone (Wöhler, A. 51, 148).

8. By oxidation of benzidine, of aniline, and of p-phenylene-diamine by MnO₂ and H₂SO₄ (Hofmann, Pr. 18, 4).—4. By action of MnO₂ and H₂SO₄ on amido-benzene p-sulphonic acid and on phenol p-sulphonic acid (Schrader, B. 8,

759)

Preparation.—1. By adding conc. Na₂Cr₂O₃Aq to a cold solution of aniline (1 pt.) in H2SO. (8 pts.) and water (25 pts.), and extracting with ether (Nietzki, B. 19, 1467).—2. 20 g. of aniline are dissolved in 600 c.c. of water containing 160 g. of H₂SO₄; 20 g. of finely-powdered potassium bichromate are added by degrees, with constant agitation, keeping the temperature at 10°-15°. After standing over night 33 g. more potassium bichromate are added; the quinone is then extracted with ether, dried over CaCl2, and the ether removed by distillation. The yield is about 86 p.c. of the theoretical amount (Schniter, B. 20, 2283; cf. Seyda, B. 16, 687; Nietzki, B. 11, 1102; A. 215, 127).—3. From p-amidophenol. p-Amido-phenol sulphate is dissolved in water, some H₂SO₄ added, and then PbO₂ gradually, with shaking, till the violet colour first formed disappears. The quinone is extracted with ether. Yield theoretical (Schmitt a. Siepermann, J. pr. [2] 19, 317).

Properties.—Yellow monoclinic prisms (from water) or plates (from EtI) (Hesse, A. 114, 299), or needles (by sublimation). Its odour is characteristic. V. sl. sol. cold water, m. sol. alcohol and ether. The solutions turn the skin brown. Sol. HClAq and conc. HNO₃. Its alkaline solution turns brown in air (Laurent, C. R. 26, 35). An alcoholic solution of hydrocerulignone deposits, even in dilute solutions, crystals of cœrulignone with steel-blue reflex.

Reactions. -1. Chlorine forms tri - chloroquinone.—2. HCl and KClO₃ give tetra-chloro-quinone (Hofmann, A. 52, 55).—3. Conc. HClAq forms chloro-hydroquinone (Wöhler; Städeler, A. 69, 808).-4. HIAq forms quinhydrone and hydroquinone.—5. HBr passed into a solution in chloroform ppts. quinhydrone, and finally forms bromo-hydroquinone. HBrAq forms, on heating di-bromo-hydroquinone (Sarauw, A. 209, 99).— 6. Bromine. (1 mol.) in CHCl, is instantly absorbed. Excess of bromine (2 mols.) appears to form C.H.Br.O., which is split up by heat into HBr and two di-bromo-quinones C₈H₂Br₂O₂ [2:5:4:1] and [2:6:4:1] (Nef, J. pr. [2] 42, 167).—
7. HNO, on heating yields oxalic and picric acids and HCy (Nietzki, A. 215, 138).—8. Ammonia gas forms emerald-green quinonimide C.H.NO, sol. water, the solution soon turning black (Woskresensky). NH, in presence of chloroform produces hydroquinone, quinhydrone, and brown C.H., NO. (Zincke, B. 16, 1556).—9. An alcoholic solution exposed to sunlight yields hydroquinone and aldehyde (Ciamician, G. 16, 111). — 10. Hydrazine hydrate gives hydroquinone and a compound C_oH₄(OH)₂N₂H₄ (Curtius a. Thun, J. pr. [2] 44, 191).—11. Heated in a sealed tube alone at 160°, or with water at 100°, it yields hydroquinone and quinhydrone (Scheid, A. 218, 227).—12. PCl, forms a thick oily product, C₁₂H,O₄P₃Cl₈. Water acting upon this forms mono- and di-chloro-hydroquinone.—13. Heated with POCl, quinone gives mono- and di-chlorohydroquinone. An amorphous body CalHiaG,

is also formed.—14. Alcoholic H2S form C12H12SO4P a yellowish crystalline mass, melting below 100°.-15. AcCl forms in the cold the di-acetyl derivatives of chloro-hydroquinone. On warming it also forms C₆H₂Cl₂(OAc)₂ (Schulz, B. 15, 652). AcBr, in like manner, gives C₈H₄Br(OAc)₂ and C₈H₂Br₂(OAc)₂.—16. Ac₂O does not act below 200°, when it forms C₆H₄(OAc)₂ and a brown mass. 17. Cold dilute NaOAc converts it into hydroquinone (Hesse, A. 220, 365).—18. Aniline in boiling alcoholic solution forms di-phenyl-di-amidoquinone and hydroquinone 3C₆H₄O₂ + 2NH₂Ph = C₀H₂(NHPh)₂O₂+2C₀H₄(OH)₂. Other primary and secondary bases act in like manner.— 19. o- and p- Nitro-aniline form red crystals C₀H₄O₂C₀H₁(NO₂).NH₂ melting at 97° and 120° respectively (Hebebrand, B. 15, 1976). o-Nitroaniline in HOAc forms C₆H₂O₂(NH.C₆H₄.NO₂)₂ [1:4:2:5] [305°] crystallising from alcohol in red needles (Leicester, B. 23, 2794).-20. o-Amidophenol in hot alcohol forms a base (C2,H18N,O,?) which crystallises in violet needles [250°], sl. sol. alcohol (Zincke a. Hebebrand, A. 226, 61). In dilute acids it forms a red solution. Ac₂O yields $C_{24}H_{16}Ac_2N_4O_4$ [285°], while Bz_2O at 150° gives $C_{24}H_{16}Bz_2N_4O_4$ [265°]. Yields a crystalline nitrosoamine $C_{24}H_{16}(N)_2N_4O_4$. Forms the salts B"H₂Cl₂, crystallising in needles with green lustre, B"H₂SO₄, B"H₂PtCl₂, and a picrate crystallising in steel-blue needles [236°]. — 21. C₂H₄(NH₂)(OMe) [1:2] forms C₂₀H₁₈N₂O₄ or C₄H₂O₂(NH.C₄H₄OMe)₂ crystallising in reddishviolet needles [230°], forming a blue solution in H_2SO_4 .—22. p-Amido-phenol hydrochloride gives $C_6H_2O_2(NH.C_6H_4.OH)_2$ crystallising in violetbrown needles, not melted at 290°.—23. (1,3,2)-Amido-o-cresol forms a compound [285°], which gives a crystalline acetyl-derivative. -24. (2, 4, 1)-Nitro-toluidine forms C₈H₃O₂.NH.C,H₈NO₂, decomposing at 300°, and C₈H₃O₂.NH.C,H₈NO₂, (L.).—25. Acetoacetic ether and ZnCl₂ at 100° form C₁₆H₁₆O₆ [184°], crystallising in needles, insol. water, sol. hot alcohol. Conc. H₂SO₄ forms a deep-blue solution. Alcoholic potash forms $C_{14}H_{16}K_{2}O_{6}$ 2aq, the salt of an insoluble acid $C_{14}H_{12}O_{6}$ (Von Pechmann, B. 21, 3005). — 26. $C_{6}H_{4}(OH)(OMe)$ [1:4] in hot ligroin forms di-methyl-quinhydrone $C_{29}H_{20}O_{6}$, crystallising in lustrous greenish-black prisms, decomposed by warm water (Hesse, A. 200, 253).

Mono-oxim CO CH:CH C:N.OH or C(OH) CH:CH C.NO. p-Nitroso-phenol. Mol. w. 123.

Formation.—1. By boiling nitroso-di-methylaniline (or nitroso-di-phenyl-aniline) hydro-chloride with dilute NaOHAq (Baeyer a. Caro, B. 7, 809, 967; Ter Meer, B. 8, 622; Fischer, B. 19, 2995).—2. By adding KNO₂ and HOAc to an aqueous solution of phenol (B. a. C.; cf. Stenhouse a. Groves, A. 188, 360).—3. By adding hydroxylamine hydrochloride to a cold dilute aqueous solution of quinone (Goldschmidt, B. 17, 213).—4. Formed also from aqueous phenol, hydroxylamine hydrochloride and H₂O₂(Wurster, B. 20, 2631). Free hydroxylamine merely reduces quinone to hydroquinone.

Properties.—Nearly colourless needles or greenish trimetric plates; a:b:c= 593:1:2·469.

M. sol. water, forming a light-green solution

which turns brown on boiling. V. sol. ether, acetone, and aqueous alkalis, sl. sol. HOAc. Decomposes at 120°-130°. Decomposed by hot conc. HClAq. When dissolved in phenol it gives, on addition of H,SO₄, a cherry-red solution turned blue on addition of KOHAq. In this reaction the compound C₁₈H₁₅NO₈ is formed. Polyhydric phenols, containing two hydroxyls in the m-position, give by like treatment fluorescent 'dichroins'; thus resorcin gives C₁₈H₁₅NO₈, C₃₈H₂₈N₂O₁₉, and azoresorcin (Brunner a. Chuit, B. 21, 249).

Reactions.—1. Oxidised by alkaline K, FeCy, p-nitro-phenol .- 2. Reduced by tin and HClAq to p-amido-phenol.—3. Nitrous acid gas passed into an ethereal solution forms diazophenol nitrate (Jaeger, B. 8, 894).—4. Gaseous HCl passed into a cooled ethereal solution forms di-chloro-amido-phenol [175°] and tri-chloro-amido-phenol (J.; Hirsch, B. 13, 1908).—5. A solution of HCl in MeOH in presence of ice forms $C_eH_2Cl_2(NH_2)(OMe)$ [72°]; while EtOH and HCl give C₆H₂Cl₂(NH₂)(OEt) [46°] (275°). 6. Conc. KOHAq at 180° forms azophenol C₁₂H₁₀N₂O₂ [214°] (Jaeger). — 7. KClO₃ and HClAq form C₈H₄ClNO₂, crystallising in yellow needles, v. sol. alcohol, insol. water, and exploding at 70° (Möhlau, B. 19, 281).—8. ClO), Me added to its sodium salt forms C₆H, (NO).O.CO, Me [137°], while ClCO₂Et yields C₆H, (NO).O.CO₂Et [109°] (Walker, B. 17, 400). 9. BzCl added to the sodium salt in ether forms C₆H₄(NO).OBz, crystallising in yellowish needles [168°-175°], sol. hot alcohol (Walker).—10. Aniline forms azophenine C_eH₂(NPh)₂(NHPh)₂. p-Chloro-aniline forms, in like manner, tetra-chloro-azophenine [265°] (Fischer a. Hepp, B. 21, 677).--11. Bromine forms C₆H₂Br₂O(NOH), crystallising in small leaflets, decomposed by water (O. Fischer, B. 21, 674).—12. Phenyl cyanate gives C₆H₁O.NO.CO.NHPh, crystallising in short yellow prisms decomposed at 160° without melting. It is converted by alkalis into the oxim, aniline, and CO2 (Goldschmidt a. Strauss, B. 22, 3105).

Salts.—C₆H₄(NO)(ONa) 2aq. Red needles (from alcohol), v. sol. water, insol. ether. Decomposed by boiling water. Ppd. by NaOH from aqueous solution.—(C₆H₄(NO).O)₂Ba (dried at 100°). Red needles (from warm water).——C₆H₄(NO).OAg aq. Violet crystals with green reflex.

Dioxim C(NOH) CH:CH C(NOH). Formed by the action of hydroxylamine hydrochloride on the mono-oxim, on hydroxylamine hydrochloride on the mono-oxim (Nietzki, B. 20, 613; 21, 430; Fischer, B. 21, 685). Yellowish needles (from hot water). Decomposes at 240°. Reduced by SnCl₂ and HCl to p-phenylene-diamine. Oxidised by alkaline K₁FeCy₆ into insoluble di-nitrosobenzene. Ac₂O forms a crystalline di-acetyl derivative, sol. hot alcohol.

Quinone-chlorimide C_eH₄NCI. [85°].

Pormed by the action of bleaching powder on a solution of the hydrochloride of p-amido-phenol (R. Schmitt, J. pr. [2] 8, 1; 19, 316; Hirsch, B. 11, 1980; 13, 1903; Fogh, B. 21, 890; Andresen, J. pr. [2] 23, 167) or C_eH₄(NH₂).OEt. Purified by steam-distillation. Golden crystals (from HOAc).

Sl. sol. cold water, v. sol. alcohol and ether. Explodes above 86°. Smells like quinone. Stains the skin brown. Turned brown by alkalis. Reduced by tin and HCl to p-amidophenol. Water at 100° forms quinone. Aqueous SO₂ forms amido-phenol sulphonic acid. HCl gives chlorinated amido-phenols. Gives Liebermann's colour reaction with phenol and H₂SO₄. Quinone-di-chlorimide C₄H₁N₂Cl₂ i.e.

C₈H₄ NCl. Formed from p-phenylene-diamine hydrochloride and bleaching-powder solution (Krause, B. 12, 47). Needles (from water), decomposing at 124°. Almost insol. cold water. Its solution stains the skin brown. Reduced by SnCl₂ to p-phenylene-diamine. HCl forms tetrachloro-p-phenylene-diamine. Br gives di-chlorodi-bromo-quinone. (β)-Naphthylamine forms amido-naphthophenazine. (β)-Naphthol forms C₁₆H₁₆N₂O, which dyes silk crimson and yields B'HNO₃ (Nietzki a. Otto, B. 21, 1745).

Quinone-phenylimide C12H11NO i.e.

C_sH₄<^{NPh}_O [97°]. Formed by oxidising p-oxydi-phenyl-amine in benzene by HgO (Bandrowski, M. 9, 134, 415). Red crystals, v. solalcohol, ether, and chloroform. Decomposed by water and alkalis. Acids reproduce quinone. Reacts with aniline, forming p-oxy-di-phenyl-amine and di-phenyl-di-amido-quinone-phenyl-imide.

Quinone-p-tolylimide C₁₃H₁₁NO. [70°]. Formed in like manner from p-oxy-phenyl-p-tolyl-amine (B.). Dark-red scales (from ligroin), sol. most solvents.

Di-methyl-amido-phenylimide C,4H,4N2O i.e. C.H. \(\sigma_0^{N.C.H.\interpretation N.Me_2}\). Phenol-blue. Formed by the action of nitroso-di-methyl-aniline on an alkaline solution of phenol, or by oxidation of a mixture of phenol and phenylene-di-methyl-pdiamine (Mohlau, B. 16, 2851; 18, 2913). Formed also from quinone-chlorimide and di-methylaniline (Fogh, B. 21, 889). Steel-blue prisms (from water) (containing aq). Its solution in HClAq is blue. Conc. H₂SO₄ forms quinone and $C_6H_4(NH_2)(NMe_2)$. Boiling NaOHAq forms NMe₂H and C₆H₄.OH. The sulphonic acid C₁₄H₁₄N₂SO₄ ½aq, formed from p-amido-phenol-sulphonic acid in water by successive treatment with Cl and dimethylaniline, crystallises in bronzed needles, forming a blue solution in NaOHAq.

Quinone tetrshydride CO CH₂CH₂CO.

Diketohexamethylene. [78°]. Formed by heating the dihydride of di-oxy-terephthalic acid (succinyl-succinic acid) of 200° and distilling the product under reduced pressure (Hermann, A. 211, 322; Baeyer a. Noyes, B. 22, 2170). Prisms, v. sol. water. FeCl₃ does nct colour its solution. Its alkaline solution turns brown in the air. Bromine forms C₆Br₄O₂. Reduces Fehling's solution. HCy forms C₆H₅(OH)₂Cy₂ [180°]. Yields a di-oxim which when quickly heated melts at 200°; when slowly heated at 192°. The di-phenyl-di-hydrazide C₆H₅(N₂HPh)₂ is obtained as a white pp. [125°] crystallising from alcohol in yellow prisms [150°], yielding a crystalline hydrochloride B"H₂Cl₂.

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An isomeride of the tetrahydride C.H.O., [170°], is got by the action of NaOHAq on succinyl-succinic ether. It crystallises in prisms (containing aq). It changes into the preceding isomeride on distillation. It reduces Fehling's solution.

References .- AMIDO-, BROMO-, BROMO-NITBO-, CHLORO-, CHLORO-NITRO-, IODO-, NITRO-AMIDO-,

OXY-AMIDO-, and OXY-QUINONE.

TRIQUINONE v. BENZENE-TRI-QUINONE. QUINONE DICARBOXYLIC ACID dihy-

dride v. DI-OXY-TEREPHTHALIC ACID.

Tetrahydride v. Dihydride of DI-OXY-TEREPHTHALIC ACID.

Quinone tetracarboxylic acid. Methul ether C₆O₂(CO₂Me)₄. [207°]. Formed by oxidation of C₆(OH)₂(CO₂Me)₄ with HNO₂ (Net, 4. 258, Colourless prisms (containing 2MeOH)

(from MeOH) or yellow needles (by sublimation). Ethyl ether CoO2(CO2Et)4. [149°]. Formed by oxidising di-amido-pyromellitic ether with conc. HNO, (Nef, C. J. 53, 428; A. 237, 28). Golden needles (from alcohol). May be sublimed. On treatment with KOHAq or with H2SO4 in HOAc it yields hydroquinone tetracarboxylic ether. Zinc-dust and HOAc also reduce it to hydroquinone tetracarboxylic ether.

QUINOPHENOL v. OXY-QUINOLINE.

 $C_{\mathfrak{o}}H_{\mathfrak{c}} < \stackrel{N}{\sim} > C_{\mathfrak{o}}O_{\mathfrak{c}}.$ DIQUINO-PHENAZINE

Formed by the action of dilute HNO, on C₆H₄:N₂:C₆O₂(OH)₂, which is got from rhodizonic soid and o-phenylene-diamine (Nietzki a. Schmidt, B. 21, 1228). Yellowish needles (containing 3aq), sl. sol. water. Converted by o-phenylene-diamine sulphate into $C_{24}H_{12}N_{4}$, crystallising in greenish needles.

DIQUINOQUINONE CH CO.CO CH. Di-

The di-oxim C, H,O, (NÓH), quinoyl. is dinitrosoresorcin (q. v.) (Goldschmidt a. Strauss, B. 20, 1611; Kehrmann a. Messinger, B. 23, 2816). The dioxim is converted by hydroxylamine into the tetra-oxim, which, on heating with Ac.O, yields the anhydride C,H₂N,O₂ crystallising in needles [61°], sol. ether. The tetraoxim may be reduced by SnCl, and HCl to tetraamido-benzene.

Triquinoquinone v. Benzene TRIQUINONE.

Reference. — OXY-DIQUINOYL.

 $C_eH_*Me < N > C_eO_*$ DIQUINO-TOLAZINE

Formed from rhodizonic acid and tolylene-odiamine, the product being oxidised by HNO₃ (Nietzki a. Kehrmann, B. 20, 324). Yellow needles (containing 2aq). On warming with an aqueous solution of a salt of tolylene-o-diamine it is converted into C_2 , $H_{18}O_9$, which crystallises from chloroform in needles (containing CHCl.).

QUINOVATANNIC ACID C14H18O8? Occurs in the bark of Cinchona nova (Quina nova) (Hlasiwetz, A. 79, 129). Translucent yellow mass, sol. water and alcohol, insol. ether. FeCl₂ gives a dark-green colour. NH₂Aq colours its solution brown. Its alkaline solution absorbs oxygen from the air. Boiling dilute H₂SO₄ splits it up into sugar and 'quinova red' C₂₈H₂₆O₁₂ (Rembold, A. 148, 273), which also occurs in the bark. Quinova red is a nearly black resin, nearly insol. water, v. sel alcohol, ether, and alkalis,

gives no colour with FeCl_s, and yields proto-catechnic acid on fusion with potash.

QUINOVIN $C_{10}H_{40}O_{8}$ (Hlasiwetz) or $C_{10}H_{44}O_{11}$ (Oudemans, jun., R. T. C. 2, 160). Quinovabitter. Quinovic acid. [a]_D = +59°. Occurrent false einchona bark from Cinchona nova (Pelletier a. Caventou, J. Ph. 7, 112; Winckler, Rep. Pharm. 51, 193; Buchner, jun., A. 17, 161; Peterson, A. 17, 165; Schnedermann, A. 45, 277; Rochleder a. Hlasiwetz, A. 79, 129; 111, 182). Occurs also in true cinchona bark (Schwarz, A. 80, 830; De Vrij, J. Ph. [3] 37, 255), in all parts of Cinchona Calisaya (from Java), and in tormentilla root (Rembold, A. 145, 9).

Preparation.-The bark is extracted with boiling milk of lime, the extract ppd. by HCl, and the pp. repeatedly dissolved in alcohol

and thrown down by water.

Properties.-Needles (from alcohol), almost insol. hot water, v. sl. sol. ether, v. sol. dilute alcohol. Tastes bitter. Dextrorotatory. Sol. aqueous alkalis. Resolved by acids into quinovic acid and quinovite. Does not reduce Fehling's solution.

Salts.—PbC₃₀H₄₆O₈aq.—(CuO)₃4C₃₀H₄₈O₈. (β)-Quinovin. [c. 235°]. [α]_D = +28° in a 2·7 p.c. alcoholic solution. Occurs in cuprea bark (from Remijia) (Liebermann a. Giesel, B. 16, 928). Scales (from dilute alcohol). Insol. ether, v. e. sol. alcohol. Dextrorotatory. Forms with alcohol the compound C_{ss}H_{s2}O₁₁5EtOH, crystallising in large prisms. Split up by dilute acids into quinovic acid and quinovite.

Quinovite $C_aH_{12}O_4$ i.e. $C_aH_aO(OH)_a$. Quinovasugar. (c. 300°). $[a]_D=60°$ (O.); = +78° (L. a. G.). Formed by the action of HCl on a hot alcoholic solution of quinovin; quinovic acid crystallises out, and the mother-liquor is neutralised by BaCO, filtered, and evaporated. Hygroscopic mass, sol. ether. Has a bittersweet taste. Does not ferment with yeast. Reduces Fehling's solution.

CaHaO(OAc)3. Tri-acetyl derivativ**e** [47°]. (c. 303°). White needles (Liebermann,

B. 17, 872).

Quinovic acid C₃₃H₃₀O₅ (Oudemans, R. T. C. 2, 160); $C_{21}H_{35}O_4$ (Hlasiwetz a. Gilm, A. 111, 182); $C_{22}H_{48}O_6$ (Liebermann). $[\alpha]_D = +86^{\circ}$. Occurs in tormentilla root (Rembold, A. 145, 6). Formed from quinovin as above. White sandy powder composed of minute trimetric six-sided laminæ, insol. water, v. sol. hot alcohol, sl. sol. Tasteless. Ppd. from alkaline solutions by soids in a gelatinous form, gradually becoming pulverulent. Dextrorotatory. H2SO. gives off CO and forms novic acid, quinochromin $C_{20}H_{40}O_2$ (crystallising in needles), quinovene $C_{20}H_{12}$ (?), and apoquinovic acid $C_{16}H_{28}O_4$ (crystallising in needles; whence $C_{16}H_{28}NaO_4$ 3 $\frac{1}{2}$ aq). Salts.— $K_2C_{24}H_{26}O_4$ 1 $\frac{1}{2}$ aq: bulky pp. — $CuA''Cu_8(OH)_8$ 5aq: light-blue pp. — Ag_2A'' :

bulky pp. Ethyl ether Cs2H46Et2Oc. [127°-130°].

Crystals, v. sol. alcohol and ether.

Pyroquinovic acid C_s, H_{ss}O_s. (above 360°).

Formed by heating quinovic acid (Liebermann a. Giesel, B. 16, 936). Needles, insol. water, v. sol. alcohol and ether. Its alkaline solution is lævorotatory. On distillation, the distillate solidifies to a clear glassy mass, easily soluble in ether, and resembling gum copal in its properties. When this substance is heated with HI and P it yields a terpene (quinoterpene) which probably has the formula C_{so}H₄₈. This terpene is also formed by direct reduction of pyroquinovic acid with HI and P (Liebermann, B. 17,

Salts.—KA' (dried at 110°).—BaA'₂. Quinoterpene C₃₀H₄₈. (above 360°). Formed

as above. Dextrorotatory.

Oxy-quinoterpene C₂₀H₄₈O₂ or C₂₀H₃₄O. Cholestol. [139°]. (above 360°). Accompanies quinovin in false cinchona bark (Liebermann, B. 17, 871; 18, 1803; Hesse, A. 234, 877). Needles (from alcohol). Reduced by HI and P to quinoterpene. Yields an acetyl derivative [126°] and a benzoyl derivative [144°].

QUINOXALINE C.H., N. i.e. C.H. < N:CH

[27°]. (222° uncor.). Formed by the action of o-phenylene-diamine on glyoxal in aqueous solution (Hinsberg, B. 17, 320; A. 237, 334). White crystals, smelling like quinoline and piperidine, miscible with cold water, alcohol, ether, and Partially separated from aqueous benzene. solution by warming or by adding KOHAq. Its aqueous solution gives white pps. with HgCl, and AgNO. Yields a sparingly soluble oxalate. Little attacked by oxidising agents. Sodium reduces it, in alcoholic solution, to phenylene-ethylene-diamine (Merz a. Ris, B. 20, 1190).—B'HCl: needles, v. sol. water. Decomposes at 184°.—B'H₂SO₄. [187°]. Silvery plates, v. sol. water.—B'2H2PtCl (dried at 100°). Needles.

References .- AMIDO- and OXY-QUINOXALINE.

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RACEMIC ACID v. TARTARIO ACID. RACEMIC CAMPHOR v. CAMPHOR.

RADICLE. Lavoisier and his followers regarded an acid as a substance formed by the combination of much oxygen with another body; the body with which the oxygen was combined was called by de Morveau the radicle of the acid, and the name was used in this sense by Lavoisier. 'Il faut donc distinguer dans tout acide la base acidifiable, à laquelle M. de Morveau a donné le nom de radical, et le principe acidifiant, c'est-à-dire, l'oxigène ' (Traité élé-mentaire de Chimie [1789] 1, 69). That the radicle of an acid might be either a simple or a compound substance is evident from de Morveau's use of the term; in speaking of the nomenclature of acids of unknown composition, he says: 'Nous nous sommes contentés de désigner l'être simple qui y modifie l'oxigène par l'expression de base acidifiable, ou, pour abréger, de radicle, de têt acide' (Mémoire sur le Développement des Principes de la Nomencla-ture Méthodique [1787]). Lavoisier, in 1789, extended the meaning of the term radicle when he said (Traité, 1, 209) that most of the oxidisable and acidifiable radicles of mineral compounds were simple bodies, whereas the radicles of compounds of vegetable and animal origin were generally composed of at least two simple bodies. The notion of radicles was here, and elsewhere, extended by Lavoisier to mean the bases or foundations whereon more complicated bodies were built up, generally by the addition of oxygen.

As organic chemistry advanced, and many compounds were obtained from animal and vegetable sources, attempts were naturally made to find some reason for the existence of so vast a number of compounds all composed of but three or four elements. Lavoisier's assertion that the radicles of organic compounds are themselves composed of two or more elements, was revived, adopted, and developed. Berzelius, Liebig, Dumas, and other earlier workers in organic chemistry were struck by the enormous number of compounds produced by the union

of the four elements carbon, hydrogen, oxygen, and nitrogen. They said the true elements of organic chemistry are the radicles cyanogen, amidogen, benzoyl, ethyl, &c., &c. Dumas (C.R.5, 300) said: 'In mineral chemistry the radicles are simple, in organic chemistry they are compound; therein consists the whole difference. The laws whereby the compounds are formed and their reactions are regulated are the same in both.' Liebig said: 'Organic chemistry is the chemistry of compound radicles.' In 1832 Liebig and Wöhler made apparent the meaning and wide applicability of the dictum that 'organic chemistry is the chemistry of com-pound radicles' by their researches into the constitution of the compounds obtained from oil of bitter almonds. By regarding these bodies as compounds of the radicle benzoyl, C,H,O, they became comparable with the compounds of the radicles potassium and other metals. From this time the conception of the radicle was firmly established.

The elements are the simple radicles, on which, and by the combinations of which, compounds are built up. The reactions of a series of compounds often show such similarities that we are obliged to conclude that the compositions of these compounds are also similar; in many cases this similarity of composition can be made apparent only by supposing that a certain group, or collocation, of elements enters into the composition of all the compounds. Such a group of elements, playing the part of a simple body throughout a series of reactions, but nevertheless separable into two or more elements, is called a compound radicle.

For an example of the working out of the conception of the radicle v. Ammonium com-POUNDS, vol. i. pp. 200-201.

M. M. P. M. RADI OIL. The product of the distillation of the wood of a juniper. Contains a sesquiterpene C₁₃H₂₄, whose hydrochloride C₁₃H₂₄2HCl melts at 118° (Wallach, A. 238, 82).

RAFFINOSE C₁₈H₂₂O₁₆5aq (Morris a. Brown, C. J. 53, 619; De Vries, R. T. C. 8, 326; C. R.

106, 751; Tollens a. Mayer, B. 21, 1569). Melitose. Gossypose. Melitriose. Mol. w. 528 to 644 (by Raoult's method) (cal. 594); 596 (by rate of osmosis in leaves of Tradescantia). [c. 87°]. S. 17 at 16° (R.); 14 at 20° (Loiseau). S. (80 p.c. alcohol) 1.4 at 70°. [a]_D = 104.6 (M. a. B.). [a]_J = 116.6. H.C. 2,019,700 (Stohmann, J. pr. [2] 45, 320; cf. Berthelot a. Matignon, C. R. 111, 13). Discovered by Johnston (C. S. Mem. 1, 159) in a manna, which drops from various kinds of eucalyptus in Tasmania. Further examined by Berthelot (C. R. 41, 392) and Rischbiet a. Tollens (B. 18, 2611). Occurs also in cotton-seed (Ritthausen, J. pr. [2] 29, 351; Böhm, J. pr. [2] 30, 37), in beet-root molasses (Loiseau, Bl. [2] 26, 365; Tollens, B. 18, 26; A. 232, 201; Lipmann, B. 18, 3087; Lindet, C. R. 110, 795; Bl. [3] 3, 682).

Preparation.—Cotton-seed cake is extracted with spirit (S.G. 848). The extract is boiled down to a small bulk, and then shaken with ether (to remove colouring matters). After a time lumps of sugar separate; these are dissolved in 80 p.c. alcohol at 70°, boiled with animal charcoal, and allowed to stand. In a week glittering needles, arranged in hemispherical masses, separate (H. Ritthausen, J. pr. [2] 29, 851).

Properties.—Groups of needles, v. e. sol. hot water, v. sl. sol. alcohol. Has a slightly sweet Does not reduce Fehling's solution. Not turned brown by boiling KOHAq. At 108° it gives off its water of crystallisation without melting; the anhydrous raffinose is not very hygroscopic. A second hydrate (containing 6aq) may be got as lamellæ by crystallisation from dilute alcohol (Berthelot, C. R. 109, 548; Bl. [3] 2, 656). Readily ferments, giving with good yeast approximately as much alcohol as cane-sugar, while with feeble yeast only onethird of that quantity is formed. Raffinose can be assimilated by young plants, being converted into starch (Brown a. Morris, C. J. 57, 486).

Reactions.—1. Boiling dilute H₂SO₄ splits it up into lævulose and 'melibiose' C₁₂H₂₂O₁₁, which yields with phenyl-hydrazine the osazone $C_{2i}H_{32}N_4O_9$. The mixture of lævulose and melibiose shows $[\alpha]_D=50^\circ$. On further boiling with dilute H2SO, the melibiose is hydrolysed, the product consisting of galactose (1 mol.), lævulose (1 mol.), and dextrose (1 mol.) (Scheibler a. Mittelmeier, B. 22, 1678, 3118; Tollens, A. 238, 308; 249, 227). Melibiose is identical with eucalyn, and may be reduced by sodiumamalgam to melibiotite C₁₂H₂₄O₁₁, a syrup which does not reduce Fehling's solution, but yields galactose on boiling with dilute acids. Invertin also splits up raffinose first into lævulose (which may be fermented by yeast) and melibiose, and then this melibiose may be split up at 36° by a conc. solution of invertin into dextrose and galactose. The Pneumococcus of Friedländer sets up fermentation in suitable solutions of raffinose (Percy Frankland, C. J. 59, 270).-2. NaOEt gives a compound containing 6 to 7 p.c. sodium.—3. Boiling HNO, (S.G. 1·15) gives 23 p.c. of mucic acid. Saccharic acid is also formed.

Salts (Beythien a. Tollens, A. 255, 195) .- $\begin{array}{lll} \operatorname{NaC_{1s}H_{3i}O_{16}}. & - \operatorname{Na_{2}C_{18}H_{30}O_{16}} \text{ aq.} & \operatorname{Crystals.} - \\ \operatorname{C_{1s}H_{2c}O_{16}(SrO)_{2}aq.} & \operatorname{Sl. sol. water.} - \operatorname{C_{18}H_{2c}O_{16}SrO.} \end{array}$ $-C_{18}H_{12}O_{16}BaO.-C_{18}H_{22}O_{16}2BaO.$ $C_{18}H_{22}O_{16}(CaO)_{s}2aq.-C_{18}H_{22}O_{16}3PbO.$ Crystals .--

Estimation.—In a mixture of cane-sugar and raffinose, the amount of raffinose may be determined by observing the change of rotatory power after hydrolysis (Creydt, B. 19, 3115; Gunning, Fr. 28, 45). The raffinose may also be ppd. by lead acetate from a solution in methyl alcohol, and the effect on the polarising action of the liquid observed (Lotman, Chem. Zeit. 12, 391; Gunning).

Cours, together with atranoric acid, in the lichen Cladonia rangiformis (Paterno, G. 12, 259). Plates (from henzeno) 259). Plates (from benzene), sol. ether. Its ammoniacal solution gives a floculent pp. of

AgA' on adding AgNOs.

RAPIC ACID C18H34O3. Occurs as glyceride, together with the glycerides of erucic and behenic acids, in rape-seed oil (Reimer a. Will, B. 20, 2387). Oil. Yields stearic acid on fusion with potash. The zinc salt melts at 78°. The Na salt is gelatinous, v. sol. water, sl. sol. alcohol.

REDUCINE $C_{12}\dot{H}_{2c}N_cO_p$ or $\dot{C}_oH_{11}N_3O_4$. An alkaloid occurring, as well as para-reducine $C_oH_oN_3O$, according to Thudichum (C. R. 106, 1803) in urine. It reduces ferric, cupric, mercuric, and silver salts, and forms an insoluble barium compound.

REDUCTION. This term is used as synonymous with deoxidation in its widest sense. v. DEOXIDATION, vol. ii. p. 377; and cf. OXIDA-TION, vol. iii. p. 657.

REGIANÍN v. JUGLONE.

RENNET v. MILK and PROTEÏDS. RESACETIC ACID v. vol. i. p. 18.

RESACETOPHENONE DI-OXY-ACETOv. PHENONE.

RESINS. Amorphous substances, occurring in all parts of plants, but especially in the bark (Wiesner, Sitz. W. 52 [2] 118). Frequently associated with essential oils. Resins are also formed from various oils by atmospheric oxidation or by the action of alcoholic potash. Boiling aqueous potash resinifies aldehydes. P2O2 converts various aldehydes (e.g. benzoic aldehyde) into resins. These resins are solid, translucent, with conchoidal fracture, insol. water, wholly or partially sol. alcohol and aqueous alkalis. The resins, therefore, contain acids. The resins soften when heated, but are decomposed by distillation. By potash-fusion protocatechnic acid is obtained from guaiacum, benzoïn, dragon's blood, asafœtida, esparto resin, myrrh, acaroid resin, and opopanax. By potash-fusion p-oxy-benzoic acid is got from benzoïn, dragon's blood, aloes, and acaroid resin; phloroglucin from dragon's blood, esparto resinand gamboge; and resorcin from galbanum, asafœtida, gum ammoniac, sagapenum, and acaroid resin (Hlasiwetz a. Barth, A. 134, 265; 138, 61; 139, 83). Dammar, sandarac, mastic, and incense-resin are not attacked by fused potash. Resins containing gum or mucilage, soluble in water, are called gum-resins. Resins mixed with essential oils are termed balsams. The following resins are completely soluble in alcohol of 95 p.c.: benzoïn, caranna, resins and balsams from conifers (colophony), dragon's blood, guaiacum, mani-resin, mastic from Alex-andria and from Bombay, black balsam of Peru,

podocarpus resin, sandarac, balsam of Tolu, xanthorrhœa resin. The following resins are only partially dissolved by alcohol of 95 p.c.: ammoniac, asafœtida, bdellium, Canada balsam, ceradia resin, copaiba balsam, copal, dammar, elemi, euphorbium, euryops resin, galbanum, gamboge, liquidambar, common mastic, Mecca balsam, myrrh, olibanum, opopanax, white balsam of Peru, sagapenum, sonora-lac, and liquid storax (Hirschsohn, Ar. Ph. [3] 10, 481; 11, 54, 152, 247, 312, 434). Ether dissolves completely: caranna, Canada balsam, conifer resins, copaiba balsam, dragon's blood, elemi, guaiac-resin, mani-resin, mastic, podocarpus resin, and sandarac. Most other resins are partially soluble in ether. FeCl, gives in alcoholic solution a blue colour with guaiacum and caranna, a dark-green colour with benzoïn and some sorts of asafætida, and a black colour with gamboge, balsam of Peru, opopanax, storax, sagapenum, shellac, and xanthorrhea resin. FeCl₂ gives in an alcoholic solution of Canada balsam and of some sorts of dammar, a pp. which disappears on heating, and with copal and sonora-lac a pp. which does not disappear on heating. Alcohol containing HCl is coloured brick-red by white balsam of Peru and ceradiaresin; red to violet by common myrrh and euryops-resin; blue to violet by some sorts of elemi; yellowish-brown to green by guaiacresin; yellow, changing through brown to cherryred, by benzoin and balsam of Tolu; crimson by xanthorrhœa resin; greenish, changing to dingy violet, by asafætida; yellow by gamboge and caranna; light rose-coloured by podocarpus resin; and brown by other resins. Conc. H₂SO₄ forms a cherry-red solution with benzoin from Siam and with balsam of Tolu; a yellow solution with gamboge; a solution with yellowishbrown fluorescence with asafætida; and brown with other resins. A drop of H2SO, added to a solution of pine-wood resin in HOAc gives a red or violet colour as the liquids mix (Morawski, C. C. 1888, 1630). Fossil resins (e.g. amber) are often found in beds of coal and lignite, being clearly derived from plants. Schmidt a. Erban (M. 7, 655; cf. Kremel, Fr. 26, 262) have determined, for a great variety of commercial resins, the quantity of alkali necessary to neutralise an alcoholic solution, and the amount required for saponification, and also the amount of iodine the resins can take up. They also base a method of separation upon the relative solubilities of resins. A classification of red resins according to their solubility in chloroform, benzene, and CS2 is given by Dobbie a. Henderson (Tr. E. 30, 624). Colophony softens under boiling water, while powdered shellac, mastic, elemi, and dammar agglomerate, and sandarac, copal, and amber remain unchanged (Kliebhau, C. C. 1888, 87). The products of distillation of colophony have been examined by many chemists (Fremy, A. Ch. [2] 59, 13; A. 15, 284; Pelletier a. Walter, A. Ch. [2] 67, 267; Thénard, Robiquet, a. Dumas, C. R. 1838, i. 460; Schiel, A. 115, 96; Couerbe, J. pr. 18, 165; Curie, C. N. 30, 189; Kelbe, B. 13, 1157; B. 14, 1240; A. 210, 1; B. 15, 308; Renard, C. R. 91, 416; B. 13, 2000; Bl. [2] 36, 215; Tilden, B. 13, 1604; Anderson, C. N. 20, 76; Mills, 'Destructive Distillation,' 31; Tichborne, Ph. [8] i. 302; Morris, C. J. 41,

167) by whom the following substances have been described as constituents: water, retinaphtha C,H_e (108°) (P. a. W.), retinyl C₂H_{1,2} (150°) (P. a. W.), retinol C_aH_e [150°) (236°-246°) (P. a. W.), retisterene [67°] (325°) (T. R. a. D.), carbonic acid, carbonic oxide, ethylene (S.), propylene (S.), heptane (97°) (T.), octane (S.), a pytene (S.), neptane (97) (1.1, octane (S.), a valerylene (50°) (C.), cymenes (170°–178°) (K.), heptinene C_{H_2} (104°) (R.), colophenone $C_{11}H_{18}O_2$ (S.) (97°), a terpene (160°) (S.), abietic acid (K.), isobutyric and methyl-propyl-acetic acids (K.), hydrocarbons $C_{11}H_{18}$ (190°–200°) (K.), and iso-butyric aldehyde (T.). Renard (4. Ch. [6] 1, 223) found among the products of distillation of colophony (rosin oil), pentane, amylene, hexane, hexylene, toluene, toluene tetrahydride, toluene hexahydride, xylene, xylene tetrahydride, xylene hexahydride, m-ethyl-propyl-benzene, terpenes, isobutyric aldehyde and acid, and valeric aldehyde and acid. Lwoff (B. 20, 1017) found, in resin oil, valeric, heptoic, ennoic, and hendecoic acids. Resin oil is coloured violet by

H₂SO, of S.G. 1.53 (Holde, C. C. 1888, 952). Resin of Pinus Laricio (Poir). [c. 100°]. V. sol. alcohol, ether, and oil of turpentine; insol. ligroin. Contains much methoxyl (Bamberger, M. 12, 441). Turned red by air, and finally brown. It contains a little free caffere acid $C_0H_3(OH)_2CH:CH.CO_2H$ [195°] and ferulic acid [4:3:1] $C_eH_s(OH)(OMe).CH:CH.CO_uH$ [169°]. 4 p.c. of caffeic and 1 p.c. of ferulic acid may be extracted by boiling with water. The resin also contains some vanillin C.H. (OH)(OMe).CHO. The resin yields pyrocatechin and protocatechuic

acid on fusion with potash.

Resin of Picea vulgaris (Link). [c. 100°]. Contains methoxyl (Bamberger, M. 12, 456). Contains p-coumaric acid and vanillin. Potash. fusion gives protocatechuic and p-oxybenzoic

Resin of Pinus sylvestris contains an acid $C_{40}H_{38}O_5$ [143°], insol. water, v. sol. alcohol. ether, and HOAc. $[a]_1 = -74^\circ$. It yields the salts $C_{20}H_{20}AgO_3$, $(C_{20}H_{20}O_3)_2Ba 2aq$, $(C_{20}H_{20}O_3)_2Ca$. and $(C_{20}H_{.9}O_{.9})_{.2}Cu$, and oily $C_{20}H_{.9}$ EtO_{.3}, which on distillation forms oily $C_{20}H_{.9}$ EtO_{.2}. Alcoholic HCl converts the acid into an isomeride [160°]. $[a]_j = -93^{\circ}$ (Shkateloff, J. R. 20, 477).

Resin from Ficus rubiginosa contains 'sycoceryl acetate ' C24H36O2 [121°] which on saponification gives acetic acid and sycoceryl alcohol-C. H. O [114°] (De la Rue a. Müller, Tr. 1860.

43; Rennie a. Goyder, C. J. 61, 916).

References .- ABIETIC ACID, ALDEHYDE RESIN. AMBER, ARBOL-A-BREA RESIN, ASAFŒTIDA, AS-PHALT, BENZOIN (gum), CANADA BALSAM, COLO-PHONY, COPAIBA BALSAM, COPAL, DAMMARA BESIN. DRAGON'S BLOOD, ELEMI, EUPHORBIUM, GAL-BANUM, GUALACUM, GUM AMMONIAC, GUTTA PERCHA. Jalap, Lac, Larch fungus, Larrea resin, Masopin, MASTIC, MAYNAS RESIN, MECCA BALSAM, MYRBB. OLIBANUM, OPOPANAX, PALISANDER RESIN, PODO-CARPIC ACID, PODOPHYLLIN, SAGAPENUM, SAN-DARAC, SCAMMONY, and STORAX.

RESORGIN C_eH_eO₂ i.e. C_eH_e(OH)₂ [1:8]. [112°]. (267°) (Kopp). S. 86°4 at 0°; 147°3 at 12°5°; 228°6 at 30° (Calderon, Bl. [2] 29, 234). V.D. 3°85 (calc. 3°81) (Troost, C. R. 89, 351). H.C.v. 683,100. H.C.p. 683,400. H.F. 87,600 (Stohmann, J. pr. [2] 45, 335). S.V. 103 (Lossen, A. 254, 59). S.V.S. 93.05.

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Formation.—1. By potash-fusion from galbanum (Hlasiwetz a. Barth, A. 180, 354), m-iodo-phenol (Körner, Bl. [2] 7, 261), phenol-p-sulphonic acid (Glutz, Bl. [2] 8, 861), p-chlorobenzene sulphonic acid (Oppenheim a. Vogt, A. Suppl. 6, 376), from umbelliferone (Hlasiwetz a. Grabowski, A. 139, 99), from asafetida, from gum ammoniac, from sagapenum, from acaroid resin (Hlasiwetz, A. 130, 354; 138, 63; 139, 78), from o-, m-, and p- bromo-phenol (Fittig a. Mager, B. 8, 362), from p-chloro-phenol (Faust, B. 6, 1022), from p-iodo-phenol above 165° (Nölting a. Wrzesinsky, B. 8, 820), from phenol (Barth a. Schreder, B. 12, 420), and from benzene m- or p- disulphonic acid (Barth a. Senhofer, B. 8, 1483; Fahlberg, Am. 2, 195).—2. By dry distillation of brazilin (E. Kopp, B. 6, 446).—3. By the action of nitrous acid on m-amido-phenol (Bantlin, B. 11, 2101).

Preparation.—By fusing benzene m-disulphonic acid with NaOH (Mühlhäuser, D. P. J.

263, 154).

Properties.—Colourless trimetric tables (from water, alcohol, or ether); a:b:c=:912:1:1059 (Calderon, C. R. 84, 779) or needles (from benzene). V. sol. water, alcohol, and ether, insol. chloroform and CS. Acquires a reddish tint on exposure to air. Neutral to litmus. Has a sweet taste. FeCl, colours its aqueous solution dark violet. Its ammoniacal solution, exposed to the air, becomes rose-red and finally brown. The ammoniacal solution leaves on evaporation a dark-blue mass, which forms a blue solution, turned red by acids. Bleaching-powder gives a transient violet colour. A drop of a solution of NaOCl gives a violet colour, quickly changing to yellow, and on heating to dark-red or brown (Stark, Ph. [3] 21, 848; Boddé, Ar. Ph. [3] 27, Resorcin reduces boiling ammoniacal AgNO, and boiling Fehling's solution. On heating resorcin with a few drops of nitro-benzene and conc. H2SO, a blue mass is got which, when poured into water and made alkaline, gives a solution with vermilion fluorescence. If resorcin is added to conc. H₂SO, and a little NaNO₂, and the mixture is heated to 100°, a product is got which, when mixed with water and rendered alkaline by NH₃, imparts to fusel oil a crimson colour with vermilion fluorescence (Bindschedler, M. 5, 168). Resorcin fused with phthalic anhydride forms fluorescein, which dissolves in squeous alkalis with strong green fluorescence Baeyer, A. 183, 8). Resorcin heated with alco-10l, beet-sugar, and HClAq gives a reddishriolet colour (Ihl, Chem. Zeit. 13, 264). A soluion of resorcin (1 pt.) in absolute alcohol 2 pts.), mixed with an aldehyde and a few drops of conc. HClAq, gives, after standing for some 10urs and then pouring into water, a resinous or crystalline pp. (Michael a. Ryder, Am. 9, 134). A mixture of resorcin and furfuraldehyde touched with a drop of HCl gives an indigo-blue subtance, which dissolves with green colour in vater, and is ppd. by HCl in blue flakes (Baeyer, B. 5, 25). On warming a liquid containing hloral or chloroform with resorcin and NaOH yellowish-red colour with green fluorescence s got (Schwarz, Fr. 27, 668). Cupric sulphate and excess of ammonia form a deep-black liquid, vhich dyes wool black (Wagner, D. P. J. 220, 16).

Titration.—1. Bromine-water is run in until all the resorcin is converted into tribromo-resorcin, which is ppd.: $C_8H_4(OH)_2 + 3BR_2 = C_8HBr_8(OH)_2 + 3HBr$. The excess is determined by adding KI and titrating with hyposulphite (Degener, J. pr. [2] 20, 322).—2. Potash and potassium iodide are added to the solution, and then a neutral solution of bleaching-powder is run in. On adding HCl a pp. of tri-iodo-resorcin is formed, and the excess of iodine is

titrated by hyposulphite (Degener).

Reactions.—1. Soda-fusion gives phloro-glucin (65 p.c.), some pyrocatechin (3 p.c.), and tetra-oxy-diphenyl (1½ p.c.) (Barth a. Schreder, B. 12, 503).—2. Exhaustive chlorination in presence of I forms CCl, and CO2 (Ruoff, B. 9, 1483).-3. ICl forms tri-iodo-resorcin (Michael a. Norton, B. 9, 1752).—4. An alkaline solution gives with KI a violet-red pp. of C₄H₂I₃KO₂ (Messinger a. Vortmann, B. 22, 2320).—5. A solution of resorcin (10 g.) in water (100 c.c.) gives a purple colour with 5 c.c. of very dilute solutions of nitrates, to which a drop of 15 p.c. HClAq and 2 c.c. of H₂SO₄ have been added (Lindo, C. N. 58, 176).—6. Resorcin (15 g.) dissolved in water (70 c.c.) and warmed with a mixture of (60 c.c. of) HBrAq (8.G. 1.47) and (20 c.c. of) HNO₃ (S.G. 1.39) gives C₄₈H₃₅BrN₂O₁₃ as a lustrous violet mass, which forms a red solution in alcohol, changed by alkalis to a blue liquid (Brunner a. Krämer, B. 17, 1873; 21, 2481). It yields C₄₉H₂₉Ac₆BrN₂O₁₂ as an orangered amorphous powder [120°], v. sol. hot alcohol. 7. Resorcin (15 g.) heated with water (60 g.), HNO₂ (20 c.c. of S.G. 1·39), and HCl (60 c.c. of S.G. 1·2) for half an hour on a water-bath forms a violet mass $C_{ae}H_{4g}ClN_sO_{1g}$, insol. chloroform, sol. alcohol and ether. It yields yellowish-brown flocculent $C_{4e}H_{4o}Ac_gClN_sO_{1g}$ (Brunner, B. 21, 2479).—8. A blue colouring matter (lacmoid) is got by heating resorcin (55 pts.) with NaNO, (18 pts.) at 130°, dissolving in NH, Aq and reppg. by HCl (Benedikt a. Julius, M. 5, 534).—9. Bromine in CS₂ forms C₀H₂Br₂(OH)₂ [111°] (Zehenter, M. 8, 293).—10. With Al₂Cl₂ it gives the compound (C₀H₄O₂)Al₂Cl₄, which is v. sol. hot CS₂, less sol. cold CS₂; decomposed at once by water into resorcin, Al₂O₃, and HCl (Claus a. Merklin, B. 18, 2934).—11. Dry ammonia passed into a solution of resorcin in dry ether forms (C,H,O,)NH, which separates as a liquid, solidifying as colourless deliquescent crystals, turning green and afterwards blue in the air (Malin, A. 138, 80) .-12. H₂O₂ and NH₂Aq give, after acidification, a brown pp. ('lacmoid') which forms an indigo blue solution in alkalis (Zulkowsky a. Peters, M. 11, 243; cf. Wurster, B. 20, 2984).—13. Anmonium carbonate solution at 125° forms (4,2,1)and (6,2,1)- di-oxy-benzoic acid (Senhofer a. Brunner, Sits. W. [2] 80,504).—14. Ammoniacal CaCl₂ at 300° forms a brownish-red colouring matter [72°], insol. water and alkalis, sol. alcohol (Seyewitz, C. R. 109, 946).-15. Sulphur and NaOHAq form, on boiling, C.H.O.S., a yellow powder, sol. alkalis, insol. water, and decomposing before fusion (Lange, B. 21, 263).— 16. K₂S₂O, acting on potassium resorain forms C.H.(ÔH)(O.SO.K) and C.H.(O.SO.K). The latter salt is converted, by heating in the dry state at 160°, into a salt of resorein disulphonic acid (Baumann, B. 11, 1911).--17. COCL, forms

C.H.: CO., a red amorphous substance, insol. water, v. sol. alcohol (Birnbaum, B. 14, 1753).-18. An alcoholic solution of potassium xanthate forms C₆H₂(OH)₂.CS₂H, crystallising in yellow needles [131°], sol. hot water (Lippmann a. Fleissner, M. 9, 296).—19. Acetone ducarboxylic acid and conc. H2SO, form(B)-umbelliferone-acetic C(OH):CH.C.O.CO $CH = CH.C.C(CH_2.CO_2H)$ > CH [202], crystallising in needles (containing aq), insole ther (Michael, J. pr. [2] 37, 469).—20. Benzene sulphonic chloride added to a solution kept slightly alkaline forms C₆H₄(O.SO₂C₆H₅)₂, crystallising from hot alcohol in needles [70°] (Georgesen, B. 24, 417).-21. Hydrogen ammonium o-sulphobenzoic acid forms C13H13NSO62aq as pale-yellow crystals, v. sol. water and alcohol. Its alkaline solutions are slightly fluorescent. Conc. HClAq at 220° reconverts it into the parent substances (Fahlberg a. Barge, B. 22, 754). Resorcin (4 mols.) heated with o-sulphobenzoic acid (1 mol.) forms C₃₁H₂₀NSO₈ 4aq. 22. Chloro-acetic acid forms C₄H₄(O.CH₂.CO₂H)₂ [194°], which yields a di-bromo-derivative (Gabriel, B. 12, 1640).—28. ClCO₂Et acting on C_eH₄(OK)₂ forms C_eH₄(O.CO₂Et)₂, a thick oil (300°) (M. Wallach, A. 226, 84).—24. Dry oxalic acid (1 mol.) in a sealed tube at 200° forms 'resorcin-oxaleïn' $C_{20}H_{14}O_{7}$, a hygroscopic red powder, sol. alcohol and ether (Claus, B. 10, 1305; 14, 2563). At 100° it becomes $C_{20}H_{12}O_6$, which is less sol. alcohol. Dilute alkaline solutions are yellow, with dark-green fluorescence. Bromine gives $C_{20}H_{1}Br_{5}O_{5}$. HNO_{2} forms $C_{20}H_{4}(NO_{2})_{4}O_{6}$. $H_{2}SO_{4}$ yields $C_{20}H_{6}(SO_{2}H)_{3}O_{6}$. $Ac_{2}O$ forms red $C_{20}H_{10}Ac_{2}O_{6}$ and colourless $C_{20}H_{4}Ac_{5}O_{6}$.—25. Crystallised oxalic acid (1 mol.) heated with resorcin (2 mols.) at 120° yields two compounds C, H,O, one of which fluoresces green in alkaline solutions (Gukassianz, B. 11, 1184). 26. HOAc and ZnCl₂ at 145° form di-oxy-acetophenone. Resorcin (100 g.) boiled with HOAc (200 g.) and ZnCl₂ (300 g.) for two hours forms 'acetfluorescein' $C_{24}H_{18}O_{5}$ and 'resacetein' $C_{16}H_{12}O_{4}$. Resacetein crystallises by spontaneous evaporation of its ammoniacal solution in red needles. Its solution in KOHAq is red, but soon It yields the salts B'HCl 2aq and B'H2SO, (dried at 110°) and a triacetyl derivative C₁₆H₀Ac₂O₄ [229°]. Acetfluorescein forms minute brownish-red crystals. Its dilute alkaline solutions exhibit green fluorescence (Nencki a. Sieber, J. pr. [2] 23, 540; Rasiński, J. pr. [2] 26, 58).—27. Resorcin (20 pts.) heated with formic acid (10 pts.) and ZnCl2 (20 pts.) for 80 minutes at 140° forms 'resaurin' $C_{10}H_{14}O_6$ or $(C_6H_2(OH)_2)_2C < \stackrel{\vee}{C_6H_2(OH)}$. Resaurin is a brick-red hygroscopic powder, forming a red solution in alkalis, sol. alcohol, insol. ether and acids (Nencki a. Schmid, J. pr. [2] 23, 547).—28. On heating with ZnCl, and acetoacetic ether or citric acid 'resocyanin acetoacetic etner or office and resources is formed (Wittenberg, J. pr. [2] 24, 125; 26, 74; Schmid, J. pr. [2] 25, 81). Resocyanin can be prepared by heating dry citric acid (60 g.) with resorcin (60 g.) and H₂SO₄ (150 g.) for an hour at 180°. Resocyanin C₂₁H₁₈O₄ (150 g.) there is not a contract of solutions of the property and the contract of solutions are solutions. [185°] is insol. cold water, sl. sol. ether, v. sol. alcohol. Its alkaline solutions are colourless with blue fluorescence. Its solutions are not

coloured by FeCl₂. It yields $\mathbf{C}_{21}\mathbf{H}_{12}\mathbf{Br}_{\bullet}\mathbf{O}_{\bullet}$ [250°] and $\mathbf{C}_{21}\mathbf{H}_{12}\mathbf{Ac}_{\bullet}\mathbf{O}_{\bullet}$ [150°]. Resocyanin yields resorcin when fused with potash. It may be reduced to a hydride C21H20Oc [259°] which yields C₂₁H₁₅Ac₂O₆ [222°]. NaOMe and MeI give C₂₁H₁₆Me₂O₆ [159°].—29. A cold alcoholic solution of sodium malonic ether forms C₁₁H₂O₅ [191°], which is v. sol. hot alcohol, insol. cold water. Its alkaline solutions are fluorescent. It splits up when heated above 191° into CO2 and methyl-umbelliferone (Michael, Am. 5, 434; J. pr. [2] 35, 455; 37, 469).—80. Resorcin (7 pts.) heated with phthalic anhydride at 200° forms FLUORESCEÏN (vol. ii. p. 557).-31. Resorcin-(2 mols.) heated with phthalimide (1 mol.) and H₂SO, at 100° forms C₂₀H₁₈NSO, a light-yellow powder, insol. benzene and ether, v. sol. alcohol. It forms $C_{20}H_{12}NaNSO_7$ 7aq and $C_{20}H_{11}Ac_2NSO_7$, a yellowish-green crystalline powder (Ostersetzer, M. 11, 425).-32. o-Benzoyl-benzoic acid forms, on heating, di-oxy-tri-phenyl-carbinol carboxylic anhydride (v. vol. iii. p. 738).-83. Maleic anhydride forms, on heating, maleïc-fluorescein CH:CH>C<C₆H₄(OH)>O, which exhibits deep green fluorescence in alkaline solution and gives crystalline C₁₆H₈Et₂O₅ and C₁₆H₈Ac₂O₅ [157°] (Burckhardt, B. 18, 2864).—34. Succinic acid 13 g.) heated with resorcin (20 g.) and H_2SO_4 (40 g.) at 195° forms 'succinyl-fluorescein $C_{16}H_{12}O_{5}$ or CH_{2} : CH_{2} $C < C_{6}H_{3}(OH) > 0$, which is crystalline (containing 3aq) and fluoresces in alkaline solution. It yields C₁₀H₃Br₄O₅ (Nencki a. Sieber, J. pr. [2] 23, 153).—35. Tartaric acid (1 mol.) heated with resorcin (2 mols.) and H₂SO₄ (1 p.c.) at 165° forms resorcin-tartreïn, a dark olive-green powder, which fluoresces in alkaline solution (Fraude, B. 14, 2558).—36. Na₂CO₃ added to a solution of resorcin and quinone forms a deep-green solution changing through yellow to brownish-red on shaking with air (Wurster, B. 20, 2934).—37. Heated with aniline and CaCl₂ it yields m-oxy-di-phenylamine. With aniline and ZnCl2 it yields diphenyl-m-phenylene-di-amine (Calm, B. 16, 2786; cf. Merz a. Weith, B. 14, 2345).—38. Aldehyde in presence of weak acids forms C14H14O4 or CH₃.CH(O.C₆H₄.OH)₂, yellow crystals, insol. water and ether, sol. alcohol. At 120° this body is converted into brown crystalline C28H26O7. By heating the compound C₁₄H₁₄O₄ with zino-dust in a current of hydrogen at 300°, resorcin is produced. Ac₂O at 140° gives C₁₄H₁₂Ac₂O₄ [282°] (Causse, Bl. [2] 47, 89; J. Ph. [5] 13, 354).—39. Chloral hydrate in aqueous solution containing NaHSO₄ forms silky needles of C₁₄H₁₂O₆, insol. water and benzene, sol. ether and alcohol. Its alkaline solutions are fluorescent. It yields a diacetyl derivative [252°] (Causse, Bl. [3] 3, 861). Resorcin and glyoxylic acid yield the same C14H12Os. By boiling chloral hydrate (5 pts.) with resorcin (10 pts.) and water (40 pts.) there is formed C₈H₆O₃ crystallising from dilute alcohol in yellowish needles, and yielding C_uH₁Ac₂O_s [159°] and C_uH₁Bc₂O_s [165°] (Michael a. Comey, Am. 5, 350).—40. Bensoic aldehyde, alcohol, and a little HCl give C₂₈H₂₀O₄ xaq [above 330°], a colourless resin, insol. water, v. e. sol. alcohol. Its alkaline solution absorbs oxygen from the air. Ac,O and NaOAc form C,H,Ac,O,. HCl.

converts it into the isomeric C28H20O4 4aq crystallising from alcohol in dimetric tables, reduced by sodium-amalgam to C26H22O4 crystallising from alcohol in prisms, and converted by Ac.O and NaOAc into C₂₀H₁₆Ac₁O₄ crystallising from xylene in prisms (Michael, Am. 5, 340).—41. CH₃.CCl₃ and caustic soda form CH₃.C(O.C,H,OH), [159°], v. sol. alcohol, sl. sol. hot water (Heiber, B. 24, 3684).-42. Camphor forms the compounds $C_0H_0O_2(C_{10}H_{10}O)$ [29°], $\lfloor a \rfloor_D = 22 \cdot 5^\circ$, crystallising in hygroscopic rectangular plates, and $C_0H_0O_2(C_{10}H_{10}O)_2$, a syrupy liquid, S.G. 12 1·037; $\lfloor a \rfloor_D = 25 \cdot 9^\circ$ (Léger, C. R. 111, 110). 43. Phenyl-hydrazine (2 mols.) rubbed with powdered resorcin (1 mol.) forms C₈H₈O₂2N₂H₃Ph, crystallising in unstable satiny needles [76°], v. sol. benzene alcohol, and ether. It is decomposed by water and alkalis (Baeyer a. Kochendörfer, B. 22, 2195).-44. Quinone (1 mol.) added to a solution of resorcin (1 mol.) in hot benzene forms C₁₂H₁₀O₄ [c. 90°] crystallising in dark-red needles with green reflex, m. sol. water (Nietzki, A. 215, 136).—45. Cyanic acid passed into an ethereal solution of resorcin forms C₈H₈N₂O₄ [120°], sl. sol. ether, m. sol. hot water (Traube, B. 22, 1579).—46. Phenyl cyanate forms at 100° C₈H₄(O.CO.NHPh)₂ [164°] (Snape, C. J. 47, 772).—47. Urea (2 pts.) in a current of CO₂ at 250° forms C₃₀H₂₀N₆O₈ 6aq, an olive-brown amorphous powder, melting above 360°, v. sl. sol. hot HOAc, sol. alkalis (Birnbaum, B. 13, 1619).-48. Heated with aqueous KHCO3 or ammonium carbonate it yields di-oxy-benzoic acid.—49. Resorcin (1 pt.) mixed with acetone (2 pts.) and fuming HClAq (1 pt.) forms C1.3H1.6O4 or CMe₂(O.C, H₄.OH)₂, which crystallises in small prisms [213°], insol. water and ether, sol. KOHAq and Na₂CO₂Aq. It is decomposed by heat into acetone and resorcin. It forms a crystalline hydrate C_{1,}H₁₀O₄ aq. Ac₂O yields C_{1,}H_{1,4}Ac₂O₄ [126°] while BzCl gives C_{1,}H_{1,4}Bz₂O₄ [115°] (Causse, Bl. [3] 7, 563).—50. Resorcin (20 g.) heated with K₂CS₂ under pressure at 100° forms C₇H₆S₂O₂ [150⁵-155°], sl. sol. CS₂, converted by potash-fusion into resorcin and (4,2,1)-di-oxybenzoic acid [205°] (Pribram a. Glücksman, M. 13, 626).

Mono-methyl ether C₅H₄(OH)(OMe). (244°). V.D. (H=1) 62·2 (obs.). Formed by heating resorcin (1 mol.) with KOH (1 mol.) and KMeSO₄ at 160° (Habermann, B. 10, 868). Formed also from resorcin, NaOMe, and MeI (Tiemann, B. 13, 2362; 14, 2019). Prepared by heating resorcin with MeOH and KHSO₄ for 10 hours at 180° (Wallach, B. 16, 151). Liquid, sol. hot water, alcohol, ether, benzene, and NaOHAq. Slightly volatile with steam. FeCl₄ colours its solution violet. Acetic anhydride forms C₆H₄(OAc)(OMe) (255°). KOH and K₂S₂O₇ form C₆H₄(OAc)(OMe), crystallising in plates, sol. water and hot alcohol.

Di-methyl ether C_cH₄(OMe)₂. (215°) (R. Schiff, B. 19, 562); (224°) (Stohmann). V.D. 68·8 (obs.). S.V. 157·13. S.G. 2 1·075. H.F.p. 74,034 [C,O₂=94,000; H₂₁O=69,000] (Stohmann, J. pr. [2] 35, 27). S.V. 157·1. Prepared by heating resorcin (1 pt.) with MeOH, KOH (1·5 pts.), and MeI (3 pts.) for 6 hours at 250° (Oechsner de Coninck, Bl. [2] 34, 149). Oil, v. sol. alcohol and ether. Volatile with

steam. Not coloured by FeCl.

Methyl ethyl ether C₄H₄(OMe)(OEt). (216°). Formed from C₄H₄(OH)(OMe), KOH, and KEtSO₄ at 165°; the product being distilled with steam (Spitz, M. 5, 488). Liquid.

Methyl propyl ether C.H. (OMe) (OPr).

(226°). Colourless liquid.

Methyl isobutyl ether C.H.(OMe)(OCH.Pr). (234°). Liquid. Methyl isoamyl ether

C₆H₄(OMe)(OC₅H₁₁). (236°). Liquid.

Mono-ethyl ether C₀H₄(OH)(OEt). Liquid. HNO₃ saturated with nitrous acid added to its ethereal solution at 0° forms C₀H₃(NO₂)(OH)(OEt) and two colouring matters: C₂H₂₀N₂O₄ [230°] crystallising in red needles, insol. water, sl. sol. boiling alcohol, forming a purple solution in H₂SO₄; and C₁H₁₁NO₃ [228°] crystallising in orange-red needles, forming a bluish-violet solution in H₂SO₄ (Weselsky a. Benedikt, M. 1, 891).

Di-ethyl ether C₆H₄(OEt), [12·4°]. (229°) (Pukall, B. 20, 1140); (235°) (Herzig a. Zeisel, M. 11, 300). Formed from resorcin, KOH, and Etl. Colourless prisms, volatile with steam. On adding one drop of KNO, solution followed by HClAq to its solution in HOAc, an intense emerald green colour is produced. Yields two di-bromo-derivatives [101°] and [77°]. Converted by dissolving in HOAc, adding NaNO, and passing in HCl into C₆H₃(NO)(OEt)₂ [123°] and C₆H₃(NO)(OH)(OEt), whence BzCl forms yellow crystals of C₆H₃(NO)(OEt)(OBz) [155°] (Kraus, M. 12, 374).

Hexa-chioro-di-vinyl ether C₄H₄(OC₂Cl₃)₂. [54°]. Formed by heating C₅H₄(OAc)₂ with PCl, at 100° (Michael, Am. 9, 210). Long prisms, insol. hot water.

Di-propyl ether $C_eH_4(OPr)_2$. (251°). V.D. 7.02 (obs.). Liquid, m. sol. hot water, sol. alcohol, ether, and ligroin (Kariof, M. 1, 258; B. 13, 1677). Br forms $C_eH_2Br(OPr)_2$ [70°].

Mono-benzyl ether $C_8H_4(OH)(OC,H_7)$. Formed from resorcin, KOH, alcohol, and benzyl bromide (Schiff a. Pellizzari, A. 221, 376; G. 13, 504).

Di-benzyl ether C_eH₄(OC,H₇)₂. [76°].

Glittering tables (from alcohol).

Tetra-nitro-di-phenyl ether $C_6H_4(O.C_6H_2(NO_2)_2)_2$. [184°]. Formed from resorcin, NaOEt, and (1,2,4)-chloro-di-nitro-benzene in alcohol (Nietzki a. Schündelen, B. 24, 3586). Colourless plates. Converted by HNO₃ into a penta-nitro-derivative [68°], and by H_2SO_4 and HNO_3 into the hexa-nitro-di-phenyl ether [220°].

Di-acetyl derivative C.H. (OAc). (273°) (Nencki, J. pr. [2] 23, 147); (278° i.V.) (Typke, B. 16, 552). Formed from resorcin

and AcCl (Malin, A. 138, 78). Oil.

Di-benzoyl derivative C₆H₄(OBz)_x [117°]. H.F. 124,598 (Stohmann, J. pr. [2] 36, 10). From resorcin (5 g.), benzoic acid (11 g.) and POCl₂ (13 g.), the latter being added slowly (Rasiński, J. pr. [2] 26, 64). Formed also by shaking resorcin with BzCl and NaOHAq (Hinsberg, A. 254, 254). Plates (from ether).

Anhydride C₁₂H₁₀O₂ i.e. O(C₀H₄·OH)₂. Resorcinyl oxide. Resorcin ether. Formed by heating resorcin with NaOH and CO₂ (Böttinger, B. 9, 182), with Na and CO₂ (Barth, B. 9, 308),

with HClAq under pressure (Barth a. Weider, B. 10, 1464), or with H2SO4 (Barth, A. 164, 122; Kopp, B. 6, 447; Annaheim, B. 10, 976). Formed also by heating resorcin at 195° with the disulphonic acid of resorcin or of phenanthrene (Hazura a. Julius, M. 5, 191). Brownishred amorphous powder, acquiring by pressure a green metallic lustre. Nearly insol. water, v. sl. sol. cold alcohol and ether. KOHAq forms a dark red solution with green fluorescence. Potash-fusion reconverts it into resorcin. Oxidised by nitric acid to isophthalic acid. Yields $C_{12}H_8Ac_2O_3$, which forms a reddish-violet solution in alkalis, and C12H6Br4Q6.

Anhydride C24H18O5. Formed, together with C12H10O3, by heating resorcin with HClAq at 180°. Brick-red powder, v. sol. alcohol and ether. Its alkaline solution is brownish-yellow with violet-blue fluorescence. Yields C24H16Ac2O4 and C24H12Br6O5. Oxidised by nitric acid to

isophthalic acid.

Resazurin C₁₂H₉NO₄ (B. a. K.), or C₁₂H₇NO₄ (Nietzki, B. 22, 3021; 24, 3366). Diazoresorcin. Azoresorcin. Resazoin. Formed by the action of nitrous acid on an ethereal solution of resorcin (Weselsky, B. 4, 613; M. 1, 889; 5, 607). Formed also by the action of MnO2 and H2SO4 on an alcoholic solution of resorcin and nitroso-resorcin. Prepared by adding fuming HNO, (6 c.c.) to resorcin (10 g.) dissolved in ether (500 c.c) at -7° . Dark-red prisms with green reflex. Insol. water and ether, v. sl. sol. cold alcohol and HOAc. Its alkaline solutions are bluish-violet. Conc. H2SO4 forms a red solution. HCl forms a crystalline hydrochloride. Yields Ba(C12H8NO4)2 crystallising in brown needles, and a sodium salt, which forms greenish needles, v. sol. water, sl. sol. NaOHAq, fluorescing brick red in dilute alcoholic solution. Resazurin forms C₁₂H₃Br₄NO₄ on bromination.

Reactions.-1. AcCl in a sealed tube at 100° forms C₁₂H₉Cl₂NO₃ (?) crystallising from HOAc in golden plates and amorphous C₂₆H₂₁Cl₃N₂O₆ (?) (Weselsky, A. 162, 288; Brunner a. Krämer, B. 17, 1854).—2. Conc. H₂SO₄ at 210° and conc. HClAq at 100° form resorufin.—3. On heating with tin and conc. HClAq an emerald-green solution is got, from which on cooling 'hydrodiazo-resorufin hydrochloride' separates as colourless leaflets or needles, which, when exposed to air, acquire a coppery lustre, and when heated in a current of air produce resorufin. Hydrodiazo-resorufin is dioxyphenoxazine, the formula being $C_eH_3(OH) < \stackrel{O}{NH} > C_eH_3OH$ (Nietzki, B. 22, 3020). 4. Hot conc. HNO, forms 'tetrazoresorcin nitrate' C18H6N7O15 (?), crystallising in lustrous garnetred needles, sol. water, alcohol, and ether with indigo-blue colour. According to Brunner a. Krämer (B. 17, 1864; 18, 587) these crystals are tri-nitro-resazurin $C_{12}H_s(NO_2)_sNO_4$. By heating them with conc. HNO₃ Weselsky obtained tetrazoresorufin nitrate' $C_{89}H_sN_{14}O_{27}(?)$, orystallising from NHO, in dark-red needles, and from wet ether in purple needles (containing 11 aq). Tin and HClAq acting upon either of these bodies form a red body, probably tri-amidoresorufin hydrochloride, and a colourless body, probably tri-amido-di-oxy-phenoxazine. By passing air through an ammoniacal solution of the colourless body there are formed lustrous green crystals of 'hydroimidotetrazoresorufin' $C_{16}H_{29}N_{14}O_{9}$ aq (W.) or $C_{12}H_{6}(NH_{2})_{3}NO_{4}$ (B.) (possibly tri-amido-resorufin). — 5. Bromine added to its solution in NaOHAq forms (C12H7Br2NO4)HBr as a lustrous green mass, forming a blue alcoholic solution with red fluorescence (Brunner a. Krämer, B. 17, 1862). Nietzki obtained $C_{12}H_1Br_1NO_4$, which gave $C_{12}H_2NaBr_4NO_4$ 2aq, crystallising from dilute alcohol in lustrous green prisms.—6. Oxidised in KOH solution by H_2O_2 to oxyresazurin C₁₈H₁₂N₂O₇ (?), which forms almost colourless crystals giving a reddish-yellow solution in alkalis. This compound is also formed by alkaline KMnO. It is reduced by zinc-dust to C₁₈H₁₆N₂O₇ (?), crystallising in colourless needles (Ehrlich, M. 8, 425).

Acetyl derivative C12HACNO4. [222°]. Formed by heating the sodium compound with NaOAc and Ac₂O at 100° (Nietzki, B. 22, 3024).

Ruby-red needles.

Ethyl ether C₁₂H₅EtNO₄. [215°]. Formed from the Ag salt, alcohol, and EtI. Dark-red needles. According to Weselsky the formula of the ether [202°] got from resazurin, alcohol, and HCl at 100° is C_{1a}H_aEt₂N₂O₆.

Resorufin C₁₂H₇NO₃ i.e.

CO.CH:C.O.C.CH:C.OH Azoresorufin. Diazo-CH:CH.C:N.C.CH:CH

resorufin.

Formation.—1. By heating resazurin with conc. H₂SO₄ at 210°.—2. By heating resorcin with a solution of nitrous acid in H2SO4 (Brunner a. Krämer, B.17, 1847).—3. By warming nitrosoresorcin with resorcin and H2SO4 (Fevre, Bl. [2] 39, 593).-4. By heating resorcin with nitrobenzene and H₂SO₄ at 170°.-5. By the action of zinc-dust on an ammoniacal solution of resazurin (Weselsky a. Benedikt, M. 5, 608) .-6. By boiling resazurin (1 pt.) with FeCl₂ (2 pts.) and fuming HClAq (10 pts.) and ppg. with water (W. a. B.).-7. By action of nitroso-phenol on resorcin or of nitroso-resorcin on phenol in presence of H₂SO₄ (Nietzki, B. 22, 3020; 23, 718).—8. By adding MnO₂ to a solution of pamido-phenol and resorcin in conc. H2SO4.

Properties. - Small dark-red grains (from dilute HClAq), insol. water and ether, sl. sol. alcohol. Forms a bluish-violet solution in H.SO. Alkalis form a crimson solution with scarlet

fluorescence.

Reactions.—1. Reduced by tin and HClAq or by zinc and HClAq to dioxy-phenoxazine $C_{12}H_1NO_2$, crystallising in nearly colourless needles, which soon become green in the air, and yielding C₁₂H_sAc₃NO₃ [216°] crystallising in colourless needles, sl. sol. hot alcohol.—2. Bromine added to the alkaline solution yields the sodium salt C₁₂H₂Br₄NO₃Na 2aq, crystallising in lustrous green needles. - 3. Nitric acid (S.G. 1.37) forms a body crystallising in green needles, forming a purple solution in water, alcohol. and ether. This body is probably tri-nitro-resorufin. 4. Fuming HCl at 100° forms 'azoresorufyl chloride' C₂₄H₁₈Cl₄N₂O₅ (?) crystallising in red plates, sol. ether (Brunner a. Krämer, B. 17, 1857). alkaline solution exhibits red Its fluorescence.

Acetyl derivative C12HAAcNO. [223°]. Formed by heating resorufin with Ac, O and NaOAc. Orange scales, v. sol. acetone, sl. sol.

alcohol ar i ether.

Ethyl ether C,2H,EtNO, [225°]. Orangered needles (Nietzki).

Nitroso-resorcin v. vol. iii. p. 619. Diresorcin v. Tetra-oxy-diphenyl.

References.—Amido-, Amido-di-imido, Bromo-, DI-BROMO-AMIDO, BROMO-NITRO-, CHLORO-, IODO-, DI-IODO-NITRO-, NITRO-AMIDO-, and NITROSO-RESORCIN.

RESORCIN-AZO- v. Azo- compounds and DISAZO- COMPOUNDS.

RESORCIN-BENZEÏN TETRA-OXY-TRI-12. PHENYL-CARBINOL.

RESORCIN CARBOXYLIC ACID v. Dr-oxy-BENZOIC ACID.

Resorcin (a)-dicarboxylic acid

C_dH₂(OH)₂(CO₂H)₂. [276°]. Formed, together with di-oxy-benzoic acid, by heating resorcin (1 pt.) with ammonium carbonate (4 pts.) and water (5 pts.) in a sealed tube (Senhofer a. Brunner, Bn. 2, 1266). Minute tables (from water), v. sl. sol. hot water. FeCl, colours its aqueous solution red.—KHA"aq: needles.—K_A"3aq: needles, v. e. sol. water.—BaA"5;aq: needles.—CuA" 51aq.—Ag2A": amorphous pp.

Resorcin (β) -dicarboxylic acid $C_0H_2(OH)_2(CO_2H)_2$. [250°]. Formed by heating (1,3,5)-di-oxy-benzoic acid with ammonium carbonate and water (S. a. B.). Four-sided prisms (containing aq), sl. sol. cold water. FeCl, colours its solution violet.— K,A".—Ba(HA"),7aq.—
BaA"4aq: needles.—Ba,C,H,O,2aq (dried at 160°).—PbC,H,O,1}aq.—CuA"33aq.

Resorcin dicarboxylic acid $C_0H_2(OH)_2(CO_2H)_2[4:2:5:1]$ (?). [192°]. Formed by oxidising the corresponding aldehyde [127°]

(Tiemann a. Lewy, B. 10, 2212). Slender needles. Diresorcin dicarboxylic acid v. Tetra-oxy-DIPHENYL DICARBOXYLIC ACID.

RESORCIN ALDEHYDE v. DI-OXY-BENZOIC

ALDEHYDE.

Resorcin dicarboxylic aldehyde v. Dr-oxy-ISOPHTHALIC ALDEHYDE.

RESORCIN-INDOPHANE C,H,N,O,. Ppd., as K salt, by warming potassium tri-nitro-resorcin with aqueous KCy (Schreder, A. 163, 297). Small lustrous needles, forming a bluishviolet aqueous solution.—Na₂C₂H₂N₄O₄ aq. K2C2H2N4O6 aq: dark-brown lustrous crystals,

exploding when heated.—BaO.H.N.O. aq.

DIRESORCIN-PHTHALEIN C.,H.2O. 53aq
(Link, B. 13, 1654) or C. H.1O. (Benedikt a. Julius, M. 5, 182). [245°]. Formed by heating diresorcin (tetra-oxy-diphenyl) with phthalic anhydride and SnCl, or H₂SO, at 115°. Silvery plates or needles, forming an indigo-blue solution in alkalis. Converted by heating with zincdust and caustic soda into diresorcin-phthalin $C_{52}H_{24}O_{10}\,8\frac{1}{9}aq$ (L.) or $C_{50}H_{14}O_{6}\,2aq$ (B. a. J.), crystallising from water in colourless plates [238°], forming a colourless solution in alkalis.

RESORCIN SULPHONIC ACID

C_sH_s(OH)₂(SO_sH). Formed by potash-fusion from the disulphonic acid (H. Fischer, M. 2,

887).—KA' 2aq : crystals. Resorcin disulphonic acid C,H2(OH)2(SO,H)2. Prepared by sulphonation of resorcin (Piccard a. Humbert, B. 9, 1479; Tedeschi, B. 12, 1267). Deliquescent needles (containing 2aq), sol. water Formed by adding Na to a and alcohol, insol. ether. FeCl. gives a red isoamyl alcohol (Bamberg colour. Gives phloroglucin on fusion with 3076). Pale-yellow liquid.

potash. Bromine forms tri-bromo-resorcin even in the cold.—K,A" aq : needles.—K,A" 4aq : deliquescent needles. — Na,A" aq. — BaA" 3aq. — BaA" 3aq. — BaA" 3aq. — Ba,C, H,S,O, 4aq.—CuA" 10aq: triclinio crystals.—Ba,C,H,S,O, 5aq: crystals.— Pb,C,H,S,O, 4aq.

Resorcin disulphonic acid

C_eH₂(OH)₂(SO₂H)₂. Formed from di-amido-benzene disulphonic acid by the diazo- reaction (Limpricht, B. 8, 290). Long four-sided needles. BaA" 2aq : crystalline pp., got by adding alcohol to its aqueous solution.

Resoroin trisulphonic acid C₈H(OH)₂(SO₂H)₃. Formed by heating the disulphonic acid with fuming H₂SO₄ at 200° (Piccard a. Humbert, B. 10, 182). In neutral solution it gives a violet colour with FeCl, Ba, A''', 3 aq. Insol. water and HClAq.

References .- Iodo-, Nitro-, and Nitroso-RESORCIN SULPHONIC ACID.

RESORCYLIC ACID v. DI-OXY-BENZOIC ACID. RESORCYLIC ALDEHYDE v. DI-OXY-BENZOIC ALDEHYDE.

Resorcylic dialdehyde v. DI-OXY-ISOPHTHALIC ALDEHYDE.

RETENE C₁₈H₁₈ i.e. CH.C₈H₄ [1:2] CH.C₆H₂MePr[1:2:6:3] Mol. w. 234. [98.5°]. (394°) (Schweizer, A. 264, 195). V.D. 8·3 (calc. 8·1) (Knecht, B. 10, 2074). S. (95 p.c. alcohol) 3 in the cold; 69 at 78°. S.G. (solid) 1.13 at 16°. H.C.v. 2,323,600. H.C.p. 2,326,100. H.F. -13,100 (Berthelot a. Vieille, A. Ch. [6] 10, 447). Occurs in scales in fossil pine-stems, accompanying fichtelite in peat bogs, and is a product of the distillation of wood (Fritsche, J. pr. 75, 281; Fehling, A. 106, 388; Wahlforss, Z. [2] 5, 73; Krauss, A. 106, 391; Ekstrand, Bl. [2] 24, 53; A. 185, 75; Bamberger a. Hooker, A. 229, 115). Produced by passing acetylene through a red-hot tube (Berthelot, J. 1866, 516). White plates, sol. alcohol, v. sol. hot HOAc, ether, and CS2. Not attacked by potash-fusion or by alkaline KMnO. Dissolved by fuming HNO₃. It is not attacked

by sodium-amalgam or HIAq at 200°.

Reactions.—1. Absorbs chlorine, forming C₁₈H₁₈Cl₂, which splits up on heating into HCl and chloro-retene C₁₈H₁₇Cl, a crystalline body.— 2. Bromine and water form di-bromo-retene C₁₈H₁₆Br₂ [180°] crystallising from CS₂ in colourless tables. Excess of Br at 100° forms viscid $C_{18}H_{16}Br_{s}$ and crystalline $C_{18}H_{14}Br_{4}$ [212°].—8. Oxidised by chromic acid mixture to retenequinone, phthalic acid, and HOAc. CrO, in HOAc forms retenequinone, retenic acid C, H, O, and an acid C₁₆H₁₆O₂ crystallising from hot alcohol in plates [139°], forming NaA' and BaA'₂, both crystallising in plates.

Compound with picric acid C₁₈H₁₈C₆H̄₂(NO₂)₂OH. [124°]. S. (95 p.c. alcohol) 2.3 at 10°; 20 at 78°. Orange-yellow needles (from alcohol). Decomposed by water. Crystallises as $(C_{1s}H_{1s})(C_{s}H_{s})C_{s}H_{2}N_{s}O_{7}$ from benzene.

Compound with di-nitro-anthraquinons. Dark orange-red needles (from HOAc).

Tetra-hydride $C_{18}H_{22}$. (280° at 50 mm.). Formed by adding Na to a solution of retene in isoamyl alcohol (Bamberger a. Lodler, B. 20,

Dodeca-hydride CisH30. (336° uncor.). Formed by heating retene with 260° (Liebermann a. Spiegel, B. 22, 780). Formed by heating retene with HI and P at

no retenequinone when oxidised by HNO₃.

**Perhydride C₁₈H₃₂. [48°]. (335° i.V.) at 719 mm. V.D. 8·69 (calc. 8·58). This is probably the constitution of fichtelite (Bamberger a. Strasser, B. 22, 3361; Spiegel, B. 22, 3369; cf. vol. ii. p. 548). By heating with iodine it is converted into C₁₈H₃₀ (346° cor.) at 714 mm. RETENE-FLUORENE v. METHYL-ISOPROPYL-

RETENE-GLYCOLLIC ACID C18H18O3 i.e. C₁₆H₁₆:C(OH).CO₂H. Formed by boiling retenequinone with NaOHAq (Bamberger a. Hooker, A. 229, 132). White crystals, sl. sol. hot water. -AgA': flocculent pp., sl. sol. hot water.

RETENE KETONE v. METHYL-ISOPROPYL-

DIPHENYLENE KETONE.

RETENEQUINONE C18H18O2 i.c.

ÇO.Ç₆H₄ Retistenequinone. Dioxy-CO.C.H.MePr retistene. [192°]. S. (95 p.c. alcohol) ·15 at 0°; 2·2 at 78°. Formed by oxidising retene with CrO₃ in HOAc (Wahlforss, Z. [2] 5, 73; Ekstrand, A. 185, 75; Bamberger a. Hooker, B. 18, 1024; A. 229, 117). Orange-red prisms, v. sl. sol. cold alcohol and ether, sl. sol. aniline and HOAc, sol. benzene and chloroform. Not attacked by Cl in the cold. Not affected by hot HNO₃. Conc. H₂SO₄ forms a green solution. Alcoholic potash imparts to its alcoholic solution a claret colour which disappears on shaking with air, but reappears on warming in absence of air.

Reactions.-1. Bromine forms C18H14Br2O2 [252°], crystallising from HOAc in orange prisms. 2. Cold NaOHAq does not dissolve it, but on boiling converts it into retene-glycollic acid.-3. On distillation with baryta it yields reteneketone and oily C₂₁H₂₄ (215-220°).—4. Distillation with sinc-dust forms retene.—5. HI and P reduce it to retene. -6. In boiling alcoholic solution it is reduced by sodium-amalgam to retene diphenic acid C₁₆H₁₆(CO₂H)₂, an unstable resin yielding Ag₂A".—7. KMnO₄ forms oxy isopropyldiphenylene ketone carboxylic acid.—8. Phenylene-o-diamine forms $C_{16}H_{16} < \stackrel{C:N}{C:N} > C_6H_4$, crystal-

lising in needles [164°], insol. water, sl. sol. alcohol, v. sol. ether, resinified by strong acids.-9. Alcoholic ammonia added to a solution of the quinone in chloroform slowly forms the imide C₁₈H₁₈O(NH), crystallising in unstable golden needles.—10. Aqueous SO, forms hydroretenequinone $C_{18}H_{16} < \stackrel{C.OH}{C_{OH}}$, crystallising in white

plates, sol. alcohol and alkalis. The same body is got by reducing the quinone with Zn and NaOHAq. It is oxidised by air to retenequinone.

Oxim C1. H1. O(NOH). [128.5°]. Golden needles (from alcohol), decomposed by acids into the quinone and hydroxylamine. Forms green compounds with iron mordants (Von Kostanecki, B. 22, 1347). RETENE DISULPHONIC ACID

C₁₈H₁₈(SO₂H)₂. S. c. 40 in the cold. Formed by sulphonation (Ekstrand, A. 185, 86). Needles (containing 10 aq), v. sol. water and alcohol. Vor. IV.

 $H_zA''5H_zSO_4$, crystallising in hair-like needles.— $K_zA''\frac{1}{2}aq$ (dried at 100'). S. c. 18. Small silky needles.—Na,A'' $\frac{1}{2}aq$ (dried at 100°). S. c. 40 in the cold.—BaA'' faq. S. 1'6.—SrA'' $\frac{1}{2}aq$. S. 4.— CaA"8aq. S.5.—MgA"2aq (dried at 100°). S.4.— CuA"5aq: long needles. S. c. 30.—PbA"aq (dried at 100°). S. 2.

Chloride C18H16(SO2Cl)2. [175°]. Prisms

(from HOAc).

Retene trisulphonic acid C18H15(SO2H)2. Got by heating retene with fuming H,SO, at 100°. Crystalline mass, v. sol. water, alcohol, and ether; not ppd. by H₂SO₄.—Ba₂A"₂18aq.

Needles. S. 7.—Pb,A''', 18aq: slender needles.

RETENIC ACID C₁₈H₁₈O₂. [222°]. A product of oxidation of retene (Ekstrand, A. 185, 111). Needles (from alcohol), v. sol. alcohol, ether, and HOAc. May be sublimed.—NaA: plates, m. sol. water.

RETINAPHTHA is TOLUENE.

RETINDOLE v. INDOLE. RETISTENE is RETENE.

RHAMNETIN is the Methyl ether of QUERCETIN (q. v.).

RHAMNITE C.H.O. i.e. CH₁.CH(OH).CH(OH).CH(OH).CH₂OH. [121°]. [α]_D = 10·7°. Formed by reducing isodulcite with sodium-amalgam in a solution kept nearly neutral by H_2SO_4 (E. Fischer a. Piloty, B. 23, 3104). Triclinic crystals, with sweet taste, v. sol. water and alcohol, sl. sol. chloroform, v. sl. sol. ether. Dextrorotatory. Does not reduce Fehling's solution.

RHAMNOHEPTOSE C,H,O, i.e. ÇH(OH).CH(OH).CH(OH).CH, CH(OH).CH(OH).CH(OH).CHO [α]_D = 8°.

Formed by the action of sodium-amalgam on the lactone, [c. 160°], $[a]_{D} = 55.6^{\circ}$, of hexa-oxy-octoic acid, which is formed from rhamnohexose by successive treatment with HCy and baryta (Fischer a. Piloty, B. 23, 3106). Sweet syrup. Dextrorotatory. Yields C₄H₁₀O₄(N₂HPh) [200°] and C₈H₁₄O₅(N₂HPh)₂ [c. 200°]. Converted by treatment with HCy, followed by saponification, into hepta-oxy-ennoic СН(ОН).СН(ОН).СН(ОН).СН ĊĦ(OĦ).CĦ(OĦ).CĦ(OĦ).CĦ(OĦ).CO₂Ħ¹ lactone of which [c. 172°] is lavorotatory $[a]_{D} = -51^{\circ}$.

RHAMNOHEXITE C,H,,O, i.c.

ÇH(OH).CH(OH).CH(OH).CH, CH(OH).CH(OH).CH₂(OH) [c. 173°]. [a]_D = 11·6°. Formed by reducing rhamnohexose [c. 173°]. with sodium-amalgam (Fischer a. Piloty, B. 23, 3106). Small colourless prisms (from hot alcohol). Does not reduce Fehling's solution.

RHAMNOHEXOSE C.H_{1.}O₆ i.e. CH..OH(OH).CH(OH).CH(OH).CH(OH).CH(OH).CH(OH).CH(OH).CH(OH).CH(OH).CHO. [181°]. [α]_D = -61°. Formed by reducing the lactone of isodulcite carboxylic acid with sodiumamalgam in acid solution at a low temperature (Fischer a. Piloty, B 23, 3104). Small tables, sl. sol. alcohol. Its aqueous solution tastes Yields a di-phenyl-di-hydrazide [200°]. sweet.

RHAMNOSE v. ISODULCITE.

RHATANINE C₁₀H₁₂NO₃. S. ·8 at 100°. S (alcohol) ·01 at 15°; ·04 at 78°. Occurs in the extract of rhatany root (Wittstein, J. 1854, 656; Ruge, Viertelj. d. nat. Ges. in Zürich, 6, Helt 3; Gintl, Sits. W. [2] 60, 668; Kreitmair, Its aqueous solution is ppd. by H.SO., forming A. 176, 69). Spherical groups of white needles,

insol. ether. Not ppd. by lead acetate or subacetate. Resembles tyrosine. On heating with a little solution of Hg(NO₂)₂ it becomes rose-red, and gives a brown pp. on further addition of $Hg(NO_3)_2$. Boiling HNO_3 gives a red colour changing to blue, and finally to a fluorescent green. Sulphuric acid forms a sulphonic acid C₁₀H₁₂NO_s(SO₃H) aq crystallising from alcohol in

plates yielding BaA', 21aq and BaA', 5aq. Salts.—H,A"HCl: monoclinic prisms, decomposed by treatment with water or alcohol. (H₂A")₂H₂PtCl₂.—Na₂A": deliquescent amorphous mass.—K₂A".—BaA" 2aq.—SrA" 2aq.— CaA".—MgA".—Ag₂A": minute needles, sl. sol. cold water.—H₂A"HNO₂.—H₂A"H₂SO₄: trimetric crystals.—H₂A"H₃PO₄: small prisms.

RHATANITANNIC ACID C₂₀H₂₀O₉ (Raabe,

J. 1880, 1060). Occurs in rhatany extract from the root-bark of Krameria triandra (Wittstein, J. 1854, 656; Grabowski, A. 143, 274). Amorphous, sol. water. Reduces Fehling's solution. Gives a dark-green colour with FeCl_s. Yields phloroglucin and protocatechuic acid on fusion with potash. Boiling dilute H2SO, forms 'rhatania-red ' C20H18O8, which yields pyrocatechin on distillation.

EHINACANTHIN C₁₄H₁₈O₄. Occurs in the root of *Rhinanthus communis* (Liborius, J. 1881, 1022). Tasteless resin, sol. alcohol. Does not form glucose on boiling with HClAq.

RHINANTHIN C₂₉H₂₂O₂₀ (L.) or C₆₄H₃₆O₄₀. A glucoside occurring in the seeds of the yellow rattle (Rhinanthus crista-galli), and giving rise to the violet colour of bread prepared from rye contaminated with these seeds. Occurs also in the seeds of Alectrolophus hirsutus (Ludwig, Z. [2] 5, 803; Ar. Ph. [2] 142, 199), and in the leaves and stalks of the snap-dragon (Antirhinum majus) (Phipson, C. N. 58, 99). late groups of prisms, with bitter-sweet taste, v. sol. water and alcohol. Gives a bluish-green solution on warming with alcoholic HCl. Resolved by dilute HClAq into brown amorphous rhinanthogen and a sugar.

RHIZOPOGONIC ACID C28H36O4. [127°]. Obtained from the mushroom Rhizopogon rubescens by extracting with ether (Oudemans, R. T. C. 2, 155). Red needles, insol. water, v. e. sol. ether, chloroform, and boiling alcohol. alkaline solutions are intensely violet.—KA' aq:

minute dark-violet crystals.

RHODAMMINES RHODIUM-AMMONIUM

COMPOUNDS, p. 405.
RHODANIDES, another name for Sulphocyanides (q. v. vol. ii. p. 848).

RHODANIC ACID C.H. NS.20 i.e.

 $HS.CH_{2}.CO.S.CN$ or $CH_{2} < \stackrel{S-CS}{<} CO.NH$ (Andreasch, M.10,73). [169°]. Formed by warming an aqueous solution of chloro-acetic acid (1 mol.) with ammonium sulphocyanide (3 mols.) (Nencki, J. pr. [2] 16,1; B. 17, 2279; Ginsberg a. Bondzynski,
 B. 19, 113). Formed also by passing HCl into an alcoholic solution of thioglycollic acid and potassium sulphocyanide (Freydl, M. 10, 82). Yellow six-sided prisms and tables, v. sl. sol. cold water, v. sol. alcohol and ether. Acid in reaction. Boiling baryta-water splits it up into HCyS and thinglycollic acid. Water at 200° forms CO₂, H₂S, NH₂, and thinglycollic acid. Reacts with aldehydes in presence of HCl, form-

ing compounds of the type R.CH:C(SH).CO.S.CN. Ethylidene-rhodanic acid C₅H₅NS₂O $[148^{\circ}]$ forms yellow needles, sol. hot water. Benzylidene-rhodanic acid C₁₀H,NS₂O [200°] also forms yellow needles, converted by hot H₂SO₄ (4 pts.) into C₁₀H,NSO₃, crystallising in needles, yielding NaA', KA', and NH₄A', and converted by HNO₃ into C₁₀H₀N₂S₂O₇, crystallising in yellow needles yielding NaA' aq. o-Nitro-benzylidenerhodanic acid C₁₀H₆N₂S₂O₈ [189°] crystallises from dilute alcohol, and may be reduced to o-amido-benzylidene-rhodanic acid, which yields $C_{10}H_{2}AcN_{2}S_{2}O$ [280°-285°] and $C_{10}H_{6}Ac_{2}N_{2}S_{2}O$ [189°]. p-Nitro-benzylidene-rhodanic acid [252°] is also crystalline (Bondzynski, M. 8, 357). FeCl, added to a hot solution of rhodanic acid forms a brown pp., from which alcohol extracts C,H,N,S,O, as a brownish-red powder, forming a red solution in alkalis.

Salts.—CuA'₂ aq: yellowish-green amorphous pp.—(C₂H₂NS₂O)₂CuCl: golden needles.

RHODATES No selts have been isolated

RHODATES. No salts have been isolated the acidic radicle of which is composed of Rh and O; but there is some reason to think that such salts exist in the solution obtained by passing Cl into an alkaline solution of $Rh_2(OH)_a$; v. under Hydrated rhodium dioxide, p. 405.

RHODINOL C10H18O i.e. CH₂:C(C₈H₇).CH:CH.CHMe CH₂OH. (217°). S.G. 15 88. Occurs in German and Turkish oil of roses (Eckart, B. 24, 4205). Oxidised by chromic acid mixture to an aldehyde, rhodinal, and rhodinolic acid. P2O, yields a terpene. KMnO, forms valeric, butyric, acetic, oxalic, and carbonic acids and C₆H₁₄O₅, possibly an alcohol.

RHODIUM. At. w. 102.7. Mol. w. unknown. Melts at c. 2000° (Pictet, C. R. 88, 1317). S.G. 12.1 (Deville a. Debray, J. 12, 240). S.H. 05803 (10° to 97°; specimen contained trace of Ir) (Regnault, A. Ch. [3] 63, 1). C.E. 0000085 at

40° (Fizeau, C. R. 68, 1125).

Occurrence.-With the other Pt metals in platinum-ore; the p.c. of Rh varies from 0 to c. 5; as much as 12.3 p.c. was found in a South American ore by Deville a. Debray (A. Ch. [2]

29, 187; cf. Kern, C. N. 35, 88).

Del Rio (v. D. a. D., l.c.) found an alloy of Rh and Au in Mexico, containing from 34 to 43 p.c. Rh. Rhodium was separated from the other Pt metals in 1803 by Wollaston (T. 1804. 419), at the same time as he isolated Pd. The name rhodium was given by W. because of the rose-

colour of the salts (bosov = a rose).

Formation.—1. By reducing RhCl₂.3NaClAq by Zn, H, formic acid, &c.—2. By strongly heat-

ing RhCl. 3NH,Cl.

Preparation.—Rh is generally prepared by adding iron to the mother-liquors from which Pt has been extracted (v. Platinum, this vol. p. 286), and then treating the solid so ppd. processes for the treatment of this residue are many; that described here is the one employed by Claus (J. pr. 85, 129), and by Gibbs (J. pr. 84, 65; 94, 19), and is a modification of Claus's older method (cf. Deville a. Debray, C. R. 78, The platinum-residue—which contains Rh, Ru, Pd, Ir, Cr, Cu, and Pb-is fused at a bright red heat with 1 pt. Pb and 1 pt. PbO; the regulus, after separation of slag, is treated with nitric acid (equal vols. conc. acid and water); the insoluble residue is washed, dried, and mixed RHODIUM. 408

with its own weight of NaCl, and this mixture is heated to low redness, in a large porcelain tube, in a stream of Cl, for some time; the treatment with NaCl and Cl is repeated; after cooling, the contents of the tube are lixiviated with water, when RhCl, 3NaCl dissolves with small quantities of the corresponding salt of Ir and traces of some of the other metals. The solution is heated with HNO, Aq, to transform IrCl, into IrCl, and Ir is then removed, as IrCl, 2AmCl, by fractional ppn. with conc. AmClAq. The filtrate from the last pp. of IrCl..2AmCl, which is almost quite free from all metals except Rh, is evaporated to dryness with HNO, Aq, to decompose the AmCl; the residue is mixed with 3 or 4 times its weight of S, and heated to bright redness in a covered porcelain crucible, which is packed in powdered charcoal in a larger earthen crucible. The regulus thus obtained is boiled for some time with aqua regia, and then with conc. H₂SO₄; it is then fused, at a low red heat, with 3 or 4 pts. Zn, whereby an alloy of Rh and Zn is formed, with the production of so much heat that part of the Zn is volatilised. After cooling, the alloy is digested with conc. HClAq, to remove impurities, and is then dissolved in aqua regia. NH, Aq is added to this solution till the pp. which forms has dissolved; the solution is boiled, and evaporated until yellow Rh2(NH3)10Cls separates; the whole is then evaporated to dryness at 100°, and the residue is treated with warm dil. HClAq until the washings show no yellow colour; the residue is dissolved in boiling water, and the liquid is filtered, hot, into dil. HClAq; the small yellow crystals of Rh₂(NH₃)₁₀Cl₆ which separate are re-crystallised several times from hot, dil. NH, Aq, whereby the salt is obtained quite pure. The pure Rh₂(NH₃)₁₀Cl₆ is decomposed by heating strongly in a graphite crucible, and the Rh thus obtained is fused in a lime crucible in the O-H flame (v. vol. iii. p. 642). The fusion removes the last traces of Os and Si. For other methods of preparing Rh v. Bunsen, A. ,146, 265; Fremy, C. R. 38, 1008; Lea, Am. S. 38, 81, 248; Schneider, P. Supp. 5, 261; Philipp, D. P. J. 220, 95; Martius, A. 67, 357; Wilm, Bl. [2] 34, 679.

Properties .- A very hard, white metal, with a tinge of blue-grey; about as ductile and malleable as Ag. Melts only in the full O-H flame, without volatilising; spits on cooling; is superficially oxidised when very strongly heated in air. Insoluble in acids when compact, but the finely divided metal ppd. from solutions is sol. HNO, Aq, and also sl. sol. HClAq in presence of air (Wilm, Bl. [2] 34, 679). Alloys of Rh with Bi, Cu, Pb, Pt, or Zn are attacked by HNO, Aq. Rh is obtained as a black porous solid by reducing solutions of Rh compounds by Zn, Hg, EtOH, or HCO.H (v. Claus, J. pr. 85, 139; Wilm, l.c.). The finely divided Rh obtained by heating the purpureochloride takes up c. 13 p.c. O when heated in a stream of air (RhO requires 13.5 p.c. O), and the O is given up by heating in H.

In its chemical relationships Rh is closely allied to Ru and Pd, and less closely to Os, Ir, and Pt; v. Noble metals, vol. iii. p. 628.

The at. w. of Rh has been determined (1) by analyses of RhCl, 8KCl (Berzelius, P. 18, 442 [1828]); (2) by analyses of a large number of

different Rh compounds (Claus, J. pr. 85, 139 [1862]); (3) by analyses of Rh₂(NH₃)₁₀Cl₄ and Rh₂(NH₃)₁₀Br₆ (Jörgensen, J. pr. [2] 27, 433, 489 [1883]); (4) by reducing pure Rh₂(NH₃)₁₀Cl₅ in H, and determining the residual Rh (Seubert a. Kobbé, A. 260, 314 [1890]); (5) by determining S.H. of Rh (Regnault, A. Ch. [3] 63, 1). No compound of Rh has been gasified. The older values for at. w. of Rh varied from 104:3 to 103. Seubert a. Kobbé have shown that the true value is less than 103.

Reactions and Combinations.-1. Rhodium black heated in air combines with O to form RhO, according to Wilm (B. 15, 2225). Leidié (Bl. [2] 50, 664) says that the products contain from 14.5 to 17.8 p.c. O (RhO requires 13.5 p.c. O).—2. Oxidised to Rh₂O₃, and then to RhO₂, by fusion with potash and nitre (Claus, l.c.); also by heating strongly with barium peroxide (D. a. D., l.c.). -8. Fusion of Rh black with potassiumhydrogen sulphate produces what is probably a double Rh-K sulphate (Claus, l.c.). -4. Heated in chlorine to c. 400°, RhCl, is formed (v. Leidié, Bl. [2] 50, 664).—5. Mixed with sodium-chloride and heated in chlorine, RhCl, 3NaCl is produced; KCl and Cl, and BaCl, and Cl, act similarly (Berzelius, P. 13, 435; Bunsen, A. 146, 266).— 6. RhS is formed by heating Rh black in vapour of sulphur (B., l.c.).-7. Rh dissolves in molten phosphoric acid (Fischer, P. 18, 257), also in molten acid phosphates (Rose-Finkner, Hand. d. anal. Chem. 6th ed., 1, 354).—8. Finelydivided Rh dissolves in molten zinc (Jörgensen, J. pr. [2] 27, 433, 489).—9. Hydrogen is absorbed in large quantities by Rh. The finely divided metal obtained by heating RhCl₂.3AmCl becomes sensibly hot when H is passed over it (Wilm, l.c.) .- 10. Rh black decomposes chlorine water, forming HClAq and O, and solutions of hypochlorites, forming chlorides and O (Jörgensen, l.c.). It also decomposes hydrogen peroxide, giving off O .- 11. Rh black decomposes formic acid solution to H and CO2; alcohol in presence of alkali is oxidised to alkali acetate, and H is given off, at c. 30° (D. a. D., A. Ch. [2] 29, 137; 56, 385; cf. Hoppe-Seyler, B. 16, 117).—12. When Rh black obtained by heating RhCl. 3AmCl is heated in coal-gas, or in ethylene, the volume of the Rh increases and C is absorbed (Wilm, Bl. [2] 84, 679).

Detection.—Rh may be brought into solution in water, either by repeated fusion with KHSO, or by mixing with NaCl and heating to low redness in Cl. To detect Rh in solution, Demarcay C. R. 101, 951) recommends to add AmCl to a neutral, or an almost neutral, solution which has been boiled with HClAq, then to add fairly cone. freshly prepared NaClOAq, and then a 20 p c. acetic acid solution, drop by drop, with constant shaking, till any pp. formed by the NaClOAq dissoves to an orange-coloured solution; on standing, this solution decolourises, deposits a greyish pp., and finally acquires an intense sky-blue colour, which remains for some time. Free HNO, or H.SO, should be absent, not much acetic acid should be added, and the temperature should not be raised. By taking two equal portions of the liquid to be tested. diluting one with water to the same bulk as the other occupies when the reagents are added, and comparing the two, '00019 g. Rh can be

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detected in 3 c.c. liquid. The other Pt metals give no reaction with NaClOAq under the above conditions.

Rhodium, alloys of. Rh alloys with many metals, by fusion with them. Some of the alloys seem to have definite compositions, e.g. Rh_Pb and RhZn_2. Descriptions of the alloys will be found in the papers referred to:—As (Wollaston, T. 1804. 419); Bi (W., l.c.); Au (W., l.c.; van Riemsdyk, J. 1880. 1269); Fe (Faraday a. Stodart, G. A. 66, 167); Pb (Debray, C. R. 90, 1195; Deville a. D., C. R. 44, 1101; 94, 1557); Pt (D. a. D., l.c.); Ag (W., l.c.); Sn (D., l.c.; Leidié, C. R. 106, 1076); Zn (D. a. D., l.c.; Bunsen, A. 146, 271; Jörgensen, J. pr. [2] 27, 434).

434).

Rhodium, ammonio-salts of, v. RhodiumAmmonium compounds, p. 405.

Rhodium, chlorides of. The only chloride of Rh which has been isolated with certainty is RhCl₂. The lower chloride RhCl₂ probably exists.

RHODIUM DICHLORIDE RhCl₂. This chloride probably exists. Berzelius obtained a greyishred powder, to which he gave this composition, by heating Rh in Cl, warming the product with KOHAq, and treating the hydrated oxide so formed with HClAq, when RhCl₂ dissolved and RhCl₂ (?) remained (P. 13, 487). According to Fellenberg (P. 50, 63) RhCl₂ is obtained, as a rose-red powder, insol. water, HClAq, or HNO₂Aq, undecomposed by KOHAq, by heating RhS in Cl. Claus (J. 1855. 423) and Leidié (C. R. 106, 1076; 107, 234) regard the existence of RhCl₂ as unproved. According to L., no substance of constant composition is obtained by F.'s method.

(?) Double salt, Am, RhCl, xH₂O. The compound to which Wilm gave this composition, and which he said was obtained by adding AmCl to RhCl, Aq in presence of much aqua regia (B. 16, 3033), is almost certainly identical with Am, RhCl, NH, NO, 3½H₂O, obtained by Leidié (v.

AMMONIUM CHLOR-RHODITE, infra).

(Rhodo-TRICHLORIDE RhCl RHODIUM chloride. Rhodo-rhodic chloride, RhCl. RhCl. Rhodium sesquichloride, Rh₂Cl₆). RhCl, is the simplest formula possible, but it is not necessarily molecular. Leidie (C. R. 106, 1076; 107, 234) prepared RhCl, by heating the alloy RhSn, in a rapid stream of dry, airfree Cl at the B. P. of S, until SnCl, ceased to be formed, allowing to cool, and removing Cl by a current of CO₂. The chloride is thus obtained as a sealing-wax-red powder. The alloy RhSn₂ is formed by fusing 1 part finely divided Rh with 30 to 50 parts Sn, and treating the product for a day or two with fairly conc. HClAq at 0° (Debray, C. R. 104, 1470, 1577). RhCl₃ was also formed by Jörgensen (J. pr. [2] 27, 433) by heating Cl₂Rh₂(NH₃)₁₀Cl₄ in a stream of Cl. Leidié (Bl. [2] 50, 664) says that RhCl, is not obtained pure by Claus' method of heating Rh in Cl, as part of the RhCl, is decomposed at the temperature which must be employed to cause combination of Rh and Cl. Leidié (l.c.) could not obtain pure RhCl, by heating Rh, S, in Cl, nor by heating the double chlorides of Rh and alkali metals with H2SO4, as recommended by Claus; in the latter reactions L. obtained alkali sulphate and Rh₂(SO₄)₃. RhCl₄ is a red powder, insol. water and acids; decomposes to Rh and

Cl at c. 450°-500° (L., l.c.). When moist RhCl_x is heated in vacuo, or in a stream of N, it is partially decomposed at 100°, and wholly at 360°, to Rh₂O₂ and HCl (L., l.c.). Slowly reduced by H at low redness (v. Claus, J. 1855, 423).

HYDRATE OF RHODIM TRICHLORIDE RhCl₃·4H₂O. Said to be formed by Claus (l.c.)-by dissolving freshly ppd. Rh(OH)₃ in HClAq, and evaporating; by Berzelius (P. 13, 437) by decomposing K₃RhCl₆Aq by H₂SiF₆Aq, filtering, and evaporating. Described as a dark-red, glassy, delfquescent solid; e. sol. alcohol, insol. ether. Leidié (Bl. [2] 50, 664) could not obtain this, nor any definite, hydrate of RhCl₃.

Double salts. Chloro-rhodites

RhCl_a·3MCl, or M₃RhCl_a· M = Am, ½Ba, ½Pb, Hg, K, Ag, Na. Also M₂RhCl_a: M = ?Am, and K. Ammonium chlor-rhodite 2Am₃RhCl_a·3H₂O· (Ammonium-rhodium trichloride). Large, red, rhombic prisms (v. Keferstein, P. 99, 275); by adding conc. HClAq to Na,RhCl_aAq, filtering

from NaCl, and adding conc. AmClAq (Claus. l.c.; Wilm, B. 16, 3033; Leidié, l.c.). Sol. water, or dil. AmClAq; insol. alcohol. The salt Am_RhCl_xH_2O, said to be formed by heating a solution of the preceding salt (Vauquelin, A. Ch. 93, 204), could not be obtained by Leidié (l.c.).

Double salt with ammonium nitrate, 2(Am,RhCl₀,NH₄NO₄).7H₂O. Obtained, as violet-red crystals, by adding HNO₃Aq to Am,RhCl₆Aq, and then excess of AmClAq, concentrating, and drying the crystals in vacuo. Scarcely sol. HNO₃Aq; when boiled with water gives off same gases as are obtained by boiling. AmClAq with aqua regia.

Potassium chlor-rhodites (1) K₂RhCl₂.3H₂O. Dark-red triclinic prisms; by adding conc. KClAq to RhCl₂Aq, and allowing to evaporate (Claus, *l.c.*). Obtained by Seubert a. Kobbé (B. 23, 2556) by saturating the filtrate from the salt. K₂RhCl₃ (infra) with HCl, filtering from KCl, and evaporating. Leidie (*l.c.*) failed to obtain this salt.

(2) K₂RhCl₃. H₂O (no water, according to L.). Brown triclinic prisms, sl. sol. water. Gives Rh, KCl, and Cl when heated to redness. Formed by heating a mixture of 1 part spongy Rh with 2 parts KCl, in a stream of Cl, so long as Cl is absorbed, dissolving in water, filtering, and evaporating under reduced pressure (S. a. K., l.c.).

Sodium chlor-rhodite Na,RhCl, 9H, O. Prepared like the K salt (Berzelius, Claus, Leidié).

The lead, mercurous, and silver salts are obtained by adding Pb(NO₃)₂Aq, HgNO₃Aq, or AgNO₃Aq, to solution of an alkali salt (Claus, l.c.). The barium salt is obtained like the K and Na salts (Bunsen, A. 146, 276). Compounds of RhCl₃ with methylamine chlorides are described by Vincent (Bl. [2] 44, 513).

Rhodium, cyanides of, and derivatives, v. vol.

ii. p. 346. Rhodium, hydrosulphide of,

RhS₂H₃(=Rh₂S₃·3H₂S). A brownish - black, voluminous pp., obtained by passing H₂S into solution of a Rh salt at c. 100° (Leidié, Bl. [2] 50, 664). Insol. acids, or alkali sulphide solutions; decomposed by Br and aqua regia. Gives off H₂S, and forms Rh₂S, when boiled with much water in N, or when heated with water in a sealed tube to 100°.

Rhodium, hydroxides of, v. Rhodium oxides | AND HYDRATED OXIDES.

Rhodium, iodide of. The black substance obtained by Jörgensen in making Rh2(NH2)10Is (v. Iodopurpureorhodium iodide, p. 406) was probably RhI₃; analysis gave 21.8 p.c. Rh, the tormula requiring 21.3 p.c. According to Claus (l.c.) an iodide of Rh is formed when KIAq is added to solution of a salt of Rh.

Rhodium, oxides and hydrated oxides of. Rh and O combine, when Rh is strongly heated in O, probably with formation of RhO. The only oxides of Rh which have been isolated with certainty are Rh₂O₃ and RhO₂; RhO also probably exists, and there are indications of the formation of RhO₃. The hydrated oxides, or hydroxides, are RhO₃.3H₂O [=Rh(OH)₃], and $RhO_2.2H_2O = Rh(OH)_4$.

RHODIUM MONOXIDE RhO (Rhodous oxide). Probably formed when Rh black is heated in a stream of air (Wilm, B. 15, 2225). Leidié (Bl. [2] 50, 664) says that from 14.5 to 17.8 p.c. O is taken up; RhO requires 13.5 p.c. O. The grey powder obtained by heating Rh(OH), was looked on by Claus as probably RhO. Deville a. Debray (A. Ch. [3] 61, 83) obtained what they regarded as RhO containing a little PbO by melting Rh with PbO, Pb, and borax, cupelling, and treating the residue with HNO, Aq. RhO is said to be fairly easily reduced to Rh by heating in H.

RHODIUM SESQUIOXIDE Rh2O3 (Rhodo-rhodic oxide). This oxide may be formed when Rh black is heated to redness in air. Leidié found from 14.5 to 17.8 p.c. O taken up (Bl. [2] 50, 664); Rh₂O₃ requires 18.92 p.c. O. Rh₂O₃ was obtained by Claus (J. pr. 76, 24; 80, 282; 85, 129) by heating Rh(NO₃)₃ in a Pt crucible till oxides of N ceased to come off; also by fusing Rh with BaO₂ and Ba(NO₃)₂. Fremy (A. Ch. [3] 44, 385) says that Rh₂O₃ is produced by heating Na₂RhCl₂ to redness in a stream of O. Rh₂O₂ is also formed by heating RhO₃H₃. Rh₂O₃ forms a grey solid; insol. acids; reduced by H to Rh (Claus, l.c.). Fremy obtained it as lustrous, metal-like, crystals, somewhat volatile when heated in a stream of O, which decomposed ClAq giving HClAq and O.

HYDRATED RHODIUM SESQUIOXIDE. Rh₂O₃.3H₂O (Rhodo-rhodic hydroxide, RhO₃H₃). A black gelatinous pp. formed by adding excess of KOHAq, and a little alcohol, to Na,RhCl,Aq. Sl. sol. conc. HClAq (Claus, l.c.). (2) Rh₂O₈.5H₂O (Hydrated rhodo-rhodic hydroxide, RhO₃H₄.H₂O). Obtained by adding KOHAq to solution of RhCl, the latter being in excess, allowing to stand till the pp. is citron-yellow (the first rose-red pp. contains the hydrate mixed with K_sRhCl_s), and washing with hot water (Claus, *l.c.*). The whole of the KOH cannot be removed from the pp. by washing. Sol. HClAq, HNO,Aq, H,SO,Aq, H₂SO₂Aq, and when freshly ppd. also in acetic Also sol. conc. KOHAq, but reppd. on dilution (Descotils, Gm.-K. 3, 1261). Said by Leidié (C. R. 107, 234) to be sol. in solution of alkali hydrogen oxalates. Solutions of this hydrate in acids yield the salts Rh₂8X on evaporation; $X = 2NO_3$, C_2O_4 , SO_4 , SO_3 , &c.

RHODIUM DIOXIDE RhO, (Rhodi- or rhodic oxide). A brown solid; obtained by repeatedly fusing finely divided Rh with KOH and KNO. porated with excess of NHAq a compound

Insol. acids, or boiling KOHAq. Reduced to Rh by H at a high temperature (Claus, l.c.).

HYDRATED RHODIUM DIOXIDE RhOx.2H2O (Rhodic hydroxide, RhO, H4). Obtained by Claus (l.c.) by leading Clinto a solution of Rh, O, 5H, O in KOHAq till a dark-brown gelatinous pp. was formed, then adding pieces of KOH from time to time, and continuing the passage of Cl until pp. became more compact, and green, and the solution became blue-violet. A greenish solid, sol. HClAq, forming a greenish-blue solution, which evolves Cl and becomes the deep-red colour of RhCl₃ solution (cf. Jörgensen, J. pr. [2] 27,

Rhodic acid and rhodates (?). blue-violet solution obtained by passing Cl into Rh₂O₃.5H₂O in KOHAq (supra) is fairly permanent, but after some time it ppts. a blue powder, gas is given off, and the liquid becomes colourless. When the blue powder is dried it becomes green RhO₂.2H₂O. A blue pp. is also obtained by carefully neutralising the blueviolet solution by HNO, Aq; this powder, when moist, is sol. HClAq, with evolution of Cl; on heating, more Cl is evolved, and the liquid becomes the red colour of RhCl, solution. Claus (l.c.) regarded the blue-violet solution as containing potassium rhodate, and the blue pp. he looked on as rhodium trioxide or rhodic anhydride, RhO,.

Rhodium, salts of. But few salts are known obtained by replacing H of oxyacids by Rh; the chief are nitrate, nitrite, oxalates, phosphates, and sulphates; a few double salts are also known.

Rhodium, sulphides of. Rh and S combine when heated, probably forming RhS. Rh₂S₃ is formed by ppn. from Rh salts.

RHODIUM MONOSULPHIDE RhS (Rhodous sulphide). Said by Berzelius to be formed by heating finely divided Rh in S vapour; also, accord ing to Vauquelin (A. Ch. 88, 167), by heating Am₃RhCl₆ with S. Fellenberg (P. 50, 63) obtained RhS by heating dry Rh₂S, in CO₂ (?). A white-blue, lustrous solid; heated in air gives spongy Rh; heated in Cl forms RhCl, and S2Cl, (F., l.c.). RhS is insol. aqua regia, according to Debray (C. R. 97, 1333.) (v. also Gibbs, J. pr. 91, 171; Lecoq de Boisbaudran, C. R. 96, 152).

RHODIUM SESQUISULPHIDE Rh.S. (Rhodorhodic sulphide). Formed, by passing H2S into solution of a Rh salt, as a black pp. according to Leidié (C. R. 106, 1076, 1533) the pp. is Rh.S.3H.S (v. Rhodium, hydrosulphide of, p. 404), which becomes Rh.S. by heating for some time with water at 100°, in a closed vessel or in N. Leidié (l.c.) obtained Rh₂S₂, in black crystalline tablets, by heating RhCl, to 360° in a stream of H₂S; according to L., this is the only method whereby pure Rh₂S, can be prepared. Loses 8 p.c. S when heated in N to dull redness; all S expelled only in O-H flame (L., l.c.). Insol. alkali solutions; not acted on by HClAq, HNO,Aq, BrAq, or aqua regia (L., l.c.). Forms the double salt Rh₂S₃.8Na₂S, by adding excess of conc. Na₂SAq to Na₂RhCl₆Aq; decomposed by water, stable in presence of excess of alkali sulphide (Leidié, l.c.). M. M. P. M.

RHODIUM - AMMONIUM COMPOUNDS. (Rhodammines. Ammoniacal rhodium bases.)
When a solution of Rh in aqua regia is evaRh₂Cl₆.10NH₃ is obtained. By treating this body with moist Ag₂O and water in the cold Rh₂Cl₄O₂H₂.10NH₈ is formed; if heat is used, the product is Rh₂O₆H₆.10NH₃. These compounds react as definite salts, which exchange their more negative radicles without losing Rh or NH₃. The classification of the rhodium-ammonium compounds generally adopted is similar to that used for the Cr and Co compounds (v. vol. ii. pp. 158 and 222). In the following general formulæ M and X represent negative radicles, which may be the same or different:

M₂·Rh₂·10NH₃·X₄ or N₄H₆(NH₄)₆(Rh₂M₂).X₄

Purpureo-rhodium series.

Rh₂·10NH₃·X₆·2H₂O or N₄H₆(NH₄)₅Rh₂·X₆·2H₂O

Roseo-rhodium series.
Rh₂.12NH₃.X₆ or N₄H₄(NH₄)₈Rh₂.X₆
Luteo-rhodium series.

The Rh compounds were investigated by Berzelius (Lehrbuch, 5th ed., 3,956); Vauquelin (A. Ch. 88, 167); Claus (Beitrage zur Chemie der Platinmetalle, Dorpat, 1854); and more recently by Jörgensen (J pr. [2] 27, 633; 34, 394; 44, 48).

I. PURPUREO-RHODIUM COMPOUNDS

M₂.Rh₂.10NH₃.X₄. Changed to roseo salts, without loss of NH₃, by boiling with alkali solution. The two atoms of radicle represented by M₂ are not removed by the ordinary reagents for removing this radicle from salts.

Chloro-purpureo- series (Jörgensen, l.c.).

Chloropurpureorhodium chloride Cl. Rh. 10NH₃.Cl. This salt is prepared by dissolving the alloy RhZn₂ in aqua regia, evaporating to dryness, treating the residue with water, filtering, adding excess NH₂Aq. evaporating to dryness at 100°, heating with dil. HClAq, and filtering while hot; the insoluble salt is washed with dil. HClAq, dissolved in boiling water, and the hot solution is filtered into HClAq (1:1 by volume); the purpureo-chloride which separates is washed with dilute HClAq, and then with alcohol. Small lustrous yellow rhombic crystals; isomorphous with chloropurpureo-cobalt chloride (v. Topsöe, J. pr. [2] 27, 488; Keferstein, A. Ch. 99, 275). S.G. 2.079 at 18°. S. 56 cold water, considerably more sol. hot water. Sol. without decomposition in hot conc. H2SO4, also in boiling NaOHAq. Easily reduced to Rh, but not acted on by such oxidisers as aqua regia or HClAq and KClO₃. Heated in CO₂ gives Rh, NH₄Cl, NH₃, and N; heated in H, gives Rh, NH, Cl, and NH, heated in HCl gas, gives Rh and NH,Cl; heated in Cl gives RhCl, and NH,Cl. Reacts with various substances to exchange Cl, for equivalent quantities of other negative radicles. Combines with PtCl₄ to form the double salt Cl.Rh.10NH,.Cl.2PtCl,.

Chloropurpureorhodium hydroxids Cl₂.Rh₂.10NH₃.(OH)₄. Known only in solution, which is obtained by rubbing the chlorochloride with a little water and moist Ag₂O for a few minutes, and filtering. Reacts strongly alkaline; gives salts by neutralisation with various acids. After warming, the solution contains roseo-chloride, and hydroxide Rh₂.10NH₃.(OH)₄, along with Rh₂.10NH₃.(OH)₄, and gives a pp. with AgNO₂Aq.

The other salts of this series are the

chloro-carbonate $M(CO_3)_2.2aq$, chloro-nitrate $M(NO_3)_4$, silicoftworide $M(SiF_6)_2$, and chloro-sulphates $M(SO_4)_2.4aq$ and $M_2(SO_4)_2.3H_2SO_4$. $M=Cl_2.Rh_2.10NH_3$.

Bromopurpureo- series (Jörgensen, l.c.).

Bromopurpureor hodium bromide

Br₃·Rh₂·10NH₂·Br₄. Deep-yellow rhombic crystals; S.G. 2·65 at 17·5°; scarcely sol. water. Prepared similarly to the chlorochloride, using HBrAq in place of HClAq; also by dissolving the chlorochloride in 7 p.c. hot NaOHAq, digesting for about an hour at 100°, adding excess-conc. HBrAq, washing the pp. which forms with dil. HBrAq and then with alcohol; also by saturating solution of roseo-hydroxide, Rh₂·10NH₃·(OH)₆, with dil. HBrAq and heating for some time at 100°. Forms a double salt with platinic bromide, Br₂·Rh₂·10NH₃·Rr₂·2PtBr₄. The bromonitrate, Br₂·Rh₂·10NH₃·(NO_{d)4}, and bromosilicoftuoride, Br₂·Rh₂·10NH₃·2SiF₆, have been isolated.

Iodopurpureo- series (Jörgensen, l.c.). Iodopurpureorhodium iodide I₂.Rh₂.10NH₃.I₄. Prepared by heating the roseohydroxide with excess of HIAq for some time; then transforming the impure product into iodochloride by washing with water, then with alcohol, rubbing repeatedly with dil. HClAq, dissolving in hot water, and filtering into dil. HClAq. The iodochloride is then dissolved in hot water and poured into KIAq; the ppd. iodo-iodide is crystallised from hot water. Reddish-yellow, rhombic crystals; S.G. 3-11 at 14-8°; very sl. sol. cold water. Forms a double salt with 2PtI₄. The other salts of this series are the iodo-chloride MCl₄, iodo-nitrate M(NO₂), iodosilicofluoride M(SiF₆)₂, and iodo-sulphate M(SO₄)₂,6aq. M=I₂.Rh₂.10NH₃.

Nitratopurpureo- series (Jörgensen, J. pr. [2] 81, 49). Nitratopurpureorhodium nitrate (NO₃)₂Rh₂.10NH₃.(NO₃)₄. Formed by heating the roseonitrate Rh₂.10NH₃.(NO₃)₅ to 100°; also by adding an equal vol. conc. HNO₃Aq to an aqueous solution of the roseonitrate, and heating at 100°. Yellowish-white crystals; deflagrates when heated. The other salts of this series are nitrato-chloride MCl₄, and nitrato-dithionate M(S₂O₉)₂. 2aq. M = (NO₃)₂.Rh₂.10NH₃.

Nitritopurpureo- series or Xanthoseries (Jörgensen, J. pr. [2] 31, 49). Nitrito-

purpureorhodium nitrate

(NO₂)₂Rh₂·10NH₃·(NO₃)₄ (Xanthorhodium nitrate). 5g. chlorochloride are heated with 50 c.c. water and 30 c.c. NaOHAq (7 p.c.) till dissolved; after cooling, 10 g. crystallised NaNO₂ are added and then a slight excess of dil. HNO₂Aq (1 acid: 2 water). The powder which separates after a time is washed with dil. HNO₂Aq, and then with alcohol, and is crystallised from hot water. Octahedral crystals, isomorphous with corresponding Co salt. Insol. alcohol and dilute HNO₂Aq; sl. sol. cold water; easily sol. hot water and cono. HNO₂Aq. Noreaction with ordinary reagents for nitrites, e.g. KIAq, KMnO₄Aq, &c. Conc. H₂SO₄ does not evolve NO₂.

The other salts of this series are: xantho-bromide MBr, xantho-bromide MCl₄, xantho-dithionate M(S₂O₄)₂, 2aq, oxalate M(C₂O₄)₃, silicoftuoride M(SiF₄)₂, sulphates M(SO₄)₂ and M(SO₄)₂,3H₂SO₄. M = (NO₂)₂,Rh₂.10NH₃,

II. Roseorhodium compounds

Rh₂.10NH₃.X₆.2H₂O. All the atoms of the negative radicle in these salts react similarly; e.g. AgNO, Aq ppts. 6Cl from the chloride. Heated to 100° 2H,O is given off, and purpureo- salts remain. These compounds are generally formed by treating the chloropurpureo- compounds with hot alkali solutions. Many are prepared from the hydroxide by the action of acids (Claus, l.c.; Jörgensen, l.c.).

Roseorhodium hydroxide Rh2.10NH2.(OH)s. Prepared by heating chloropurpureochloride with excess of moist Ag₂O and water for some days, filtering from AgCl, and evaporating in vacuo. A yellow deliquescent solid, with strongly alkaline reaction; ppts. metallic hydroxides from salts; decomposes NH, salts, giving off NH, Neutralises acids, forming roseo- salts.

Roseorhodium bromide Rh. 10NH .. Br. 2H, O. Ppd. by adding cone. HBrAq to solution of the hydroxide; also by boiling chloropurpureochloride with NaOHAq, and then adding excess of conc. HBrAq. Pale-yellow crystals; changes to the purpureo- salt at 100°, or by crystallising from hot water. The other salts of this series are: carbonate M(CO₂), aq; iodo sulphate MI₂(SO₄)₂; nitrate M(NO₅)₆, and the double salt M(NO₃)2.PtCl₄; phosphate M(HPO₄)3.4aq, and the sodium pyrophosphate M(NaP₂O₂)_{2.23aq}; sulphate M(SO₄)_{3.3aq}, and the double salt M(SO₄)_{2.}PtCl₈.

HI Lumpopurous Annual Solution of the sulphate M(SO₄)_{2.2}PtCl₈.

III. LUTEORHODIUM COMPOUNDS Rh2.12NH3.X8. These salts are very stable. Their colours are the same as those of corresponding roseo- salts. The chloride can be prepared by heating the chloropurpureochloride with conc. NHAq; most of the other salts are obtained from the chloride (Jörgensen, J. pr. [2] 44, 48, 63). The negative radicles are all acted on similarly by

reagents.

Luteorhodium chloride Rh₂.12NH₂.Cl₂. Prepared by heating chloropurpureochloride with c. 30 times its weight NH3Aq (25 p.c. NH3) for several days at 100°-102°, dissolving in hot water, and ppg. by conc. HClAq. S. c. 12 at 8°. Not changed to chloropurpureochloride by digestion with HClAq at 100°. Forms the double salt Rh₂.12NH₂.Cl₂.3PtCl₄.6aq. The other salts of this series described by Jörgensen are: bromide MBr., nitrates M(NOs), and M(NOs), HNOs, phosphate M(PO₄)₂Saq and sodium luteo-pyrophosphate M(NaP₂O₇)₂.23aq, and sul-

phate M(SO4)3.5aq Rhodium-pyridine compounds. Compounds of Rh with pyridine, C,H,N, and negative radicles have been prepared by Jörgensen (J. pr. [2] 27, 483). The compounds examined contain 8C, H, N, and correspond with the chromtetrammonium series of ammoniacal compounds (vol. ii. p. 159), and with the octammine series of ammoniacal Co compounds (vol. ii. p. 225). The composition of these pyridine compounds is represented by the formula M,Rh2(C,H,N),X2; where M = Cl and X = Br, Cl, OH, NO, and SO, The salts resemble the NH, compounds generally. The chloride (X = Cl) is prepared similarly to chloropurpureorhodium chloride, using pyridine in place of ammonia; the other salts are prepared from the chloride by the action of acids. AgNO, Aq ppts. one-third of the halogen only from the chloride and bromide. М. М. Р. М.

RHODIZONIC ACID v. DI-OXY-DIQUINOYL.

RHEADINE $C_{21}H_{21}NO_{e}$. [232°]. S. (80 p.c. alcohol) ·09. S. (ether) ·08 at 18°. An alkaloid occurring in all parts of the red poppy Papaver Rhaas, and in the white seed capsules of the white poppy Papaver somniferum (Hesse, A. 140, 145; 149, 35; 153, 47). Small white prisms, nearly insol. water, alcohol, and ether. May be sublimed. Tasteless, and not poisonous. Its solution in very weak HCl is colourless, but in conc. HClAq and in H2SO4 it forms purple solutions, the base being partially decomposed. The colourless solution is ppd. by tannin and by HgCl2.

Salts.—B'2H2PtCl6 2aq: yellow amorphous pp.—B'HI 2aq: mass of minute prisms.

Rhoagenine $C_{21}H_{21}NO_{4}$. [223°]. S. (80 p.c. alcohol) 07. S. (ether) 06. Formed by boiling rhoeadine with dilute H2SO4. Small crystals (from alcohol), sl. sol. ether, alcohol, and water, v. sol. dilute acids. Its alcoholic solution turns litmus blue. Cannot be sublimed. Not coloured by acids. Its salts taste bitter. - B'2H2PtCle-B'HI: short prisms, m. sol. hot water.

R1BONIC ACID C.H., O. i.e. CH2(OH).CH(OH).CH(OH).CH(OH).CO2H. Tetra-oxy-valeric acid. Formed by heating CH, (OH).CH(OH).CH(OH).CO,H, arabonic acid, with pyridine in a closed vessel at 130° (E. Fischer a. Piloty, B. 24, 4216). When set free from its salts it changes into the lactone.—

 HgA'_{2} : needles.— CdA'_{2} . $[\alpha]_{D} = 0.6^{\circ}$. Needles. $Lactons\ C_{s}H_{s}O_{s}$. $[72^{\circ}-76^{\circ}]$. $[\alpha]_{D} = -18^{\circ}$. Colourless prisms, v. e. sol. water and alcohol, v. sl. sol. ether. Converted by heating with pyridine and water at 135° into arabonic acid. HNO, yields tri-oxy-glutaric acid. Phenyl

hydrazine forms $C_3H_4O_3(N_2H_2Ph)$ [164°]. RIBOSE $\times C_3H_{10}O_3$. Formed by reducing the lactone of ribonic acid with sodium-amalgam (Fischer a. Piloty, B. 24, 4220). Colourless syrup. Yields a phenyl-hydrazide [155°] and a p-bromo-phenyl-hydrazide C₁₁H₁₅O₄N₂Br [165°].

ACID C18H24O3. RICINELAIDIC Formed by the action of nitrous acid on ricinoleic acid (Boudet, A. 4, 16; Playfair, A. 60, 322; Bouis, A. Ch. [3] 44, 82; Ulrich, Bl. [2] 9, 225). White Got also by saponifying ricinelaidin. silky needles, v. sol. alcohol and ether. Reddens Yields n-heptoic acid on oxidation with litmus. HNO, (Krafft, B. 21, 2735). Alkaline KMnO, yields two tri-oxy-stearic acids [120°] and [116°] (Grüssner a. Hazura, M. 10, 196; Mangold, M. 13, 326). Forms oily C₁₈H₃₁Br₂O₃, converted by KOHAq in the cold into C₁₈H₃₂BrO₃, whence alcoholic potash forms an acid melting at 71°.

Salts.—BaA'2: unctuous powder.—AgA'. white powder, sol. NH₃Aq.

Ethyl ether EtA'. [16°]. Crystalline. A mide C₁₈H₂₅NO₂. [93°] (Rowney, Chem. Gaz. 1855, 361).

Reference. BROMO-RICINELAIDIC ACID.

BICINELAIDIN. [45°] (Bouis); [66°] (Boudet). Formed by the action of nitrous fumes on castor oil. Solid, v. sol. alcohol and ether. Split up by boiling KOHAq into glycerin and potassium ricinelaidate. Yields heptoic aldehyde (œnanthol) on dry distillation (Bertagnini, A. 85, 282). Yields octyl alcohol and potassium sebacate on distillation with potash (Bouis).

RICINIC ACID C₁₈H₃₄O₃ *i.e.* H₃₂(OH).CO₂H. [81°]. (252° at 15 mm.). C, H, (OH).CO, H. Formed by distilling barium ricinoleate in vacuo (Krafft, B. 21, 2786). Plates (from dilute alcohol). Yields, on oxidation by HNO, n-heptoic acid and an acid [c. 103°] not volatile with steam. Converted by HCl into diricinic acid C, H₃₂(OH).CO.O.C, H₃₂.CO₂H, triricinic acid C₁₇H₃₂(OH).CO.O.C, H₃₂.CO.O.C, H₃₂.CO.O.C, H₃₂.CO.D.A, and polyricinic acids. Sulphuric acid forms these as well as SO₃H.O.C₁,H₃₂.CO₂H, SO₃H.O.C₁₇H₃₂.CO.O.C₁₇H₂₂.CO₂H, &c. H2SO4 acting on castor oil forms the compound

acids, constituting Turkey red oil. The polyricinic acids are converted by boiling KOHAq but not by Na₂CO₃ into ricinic acid. Hot water decomposes the polyricino-sulphuric acids into H₂SO₄ and polyricinic acids (Juillard, Bl. [3] 6, 638).

RICININE. An alkaloid occurring, according to Tuson (C. J. 17, 195; C. N. 22, 229; cf. Werner, C. N. 22, 203), in the seeds of the castor oil plant (Ricinus communis). Rectangular prisms or laminæ, insol. water, nearly insol. ether, sol. alcohol.

RICINISOLIC ACID v. TRI-OXY-STEADIC ACID.

RICINOLETC ACID C₁₈H₃₄O₃ i.e. CH₃:[CH₂]₃,CH(OH).CH:CH[CH₂]₃.CO₂H (?) [17°]. Occurs as glyceride in castor oil (Bussy a. Lecanu, J. Ph. 13, 70; Saalmüller, A. 64, 108; Svanberg a. Kolmodin, J. pr. 45, 431; Bouis, A. Ch. [3] 44, 103; 48, 99; Petersen, A. 118, 69). Prepared by saponification of castor oil by KOH or HCl and purified by means of its Ca or Ba salt (Claus, B. 9, 1916; Krafft, B. 21, 2731). Crystalline mass, miscible with alcohol and ether. Its alcoholic solution is alkaline in reaction. Does not absorb oxygen from the air. Yields $C_{18}H_{28}AcO_{2}$. Yields polymerides when heated with water in closed vessels (Scheurer-Kestner, C. R. 113, 201).

Reactions.-1. Oxidised by HNO_s to azelaic acid C₀H₁₈O₄, oxalic acid, and heptoic acid.— 2. Alkaline KMnO, forms two isomeric tri-oxystearic acids (Dieff, J. pr. [2] 39, 345; B. 20, 1211; Hazura a. Grüssner, M. 8, 475; Mangold, M. 13, 326).—2. Nitrous acid forms ricinelaidic acid.-3. Yields methyl hexyl ketone (173°) and an acid $C_{1a}H_{2a}O_3$ [81°] on distillation of the Ba salt.—4. The Na salt on distillation with NaOH yields methyl hexyl ketone, sec-octyl alcohol and sodium sebacate Na₂C₁₀H₁₀O₄.-5. Bromine forms oily C₁₈H₃₄Br₂O₈, converted by alcoholic potash into $C_{1s}H_{32}BrO_3$, whence Br forms oily $C_{1s}H_{32}Br_3O_3$, converted by alcoholic potash at 120° into ricinostearolic acid $C_{1s}H_{32}O_3$, which combines with bromine forming unstable oily $C_{18}H_{32}Br_4O_3$.—6. PCl₈ forms $C_{18}H_{32}Cl_2O$.—7. HI and P give C18H38IO2.

Salts.—BaA'₂: soft scales (from alcohol), sl. sol. water.—SrA'₂: small grains (from alcohol).—CaA'₂ (dried at 100°). [80°].—MgA'₂: slender needles, v. sol. alcohol.—PbA'₂. [100°]. Crystalline mass, v. sol. ether.—ZnA'₂.—AgA': ourdy pp.

Ethyl ether EtA'. Oil. Amids C, Har. CONH. [66°]. Nodules (from alcohol).

References .- Bromo- and Oxy- RICINOLEÏO ACID

RICINOLIC ACID v. TRI-OXY-STEARIC ACID. RICINOSTEAROLIC ACID C18H32O3. [51°]. Formed by heating the dibromide of ricinoleic acid with alcoholic potash (Ulrich, Z. 1867, 547). Groups of needles (from alcohol), v. sol. ether. Volatile.—BaA'2: plates (from alcohol) [135°].—AgA': granular pp.

RICINOSTEAROXYLIC ACID C18H32O1. [78°]. Formed by heating ricinostearolic acid with moist Ag₂O (Ulrich, Z. 1867, 550). Dendritic groups of needles (from alcohol), v. sol. ether.—BaA'2.—AgA'; granular pp. decomposed by hot alcohol.

ROBININ C₂₅H₃₀O₁₈. [195°]. Occurs in the blossoms of the acacia (Robinia pseudacacia) (Zwenger a. Dronke, A. Suppl. 1, 257). Thin yellow needles (containing 5) aq). Neutral in reaction. Tasteless, sl. sol. cold water and alcohol, insol. ether. Sol. alkalis, forming yellow The ammoniacal solution turns solutions. brown in air. FeCl₃ gives a brown colour. Its alcoholic solution is ppd. by lead subacetate. Reduces hot Fehling's solution. HNO₃ forms oxalic and picric acids. Split up by boiling dilute acids into quercetin and a sugar.

ROCCELLIC ACID C₁,H₁₂O₄. [132°]. S. (boiling alcohol of S.G. 819) 55. Occurs in various species of Roccella (Heeren, S. J. 59, 346; Liebig, P. 21, 31; Schunck, A. 61, 64; Hesse, A. 117, 332). Four-sided prisms, insol. water, v. sol. alcohol and ether, sl. sol. warm benzene.

Sol. Na, CO, Aq and borax.

Salts.—BaA" (dried at 100°).—CaA" aq: amorphous pp. -Pb, A", (OH), 2aq: white powder. -Ag₂A'': amorphous pp. Ethyl ether Et_2 A''. Oil.

Anhydride C₁₁H₁₀O₂. Got by heating the acid above 220°. Oil, v. sol. hot alcohol and ether.

Anilide C₁₇H₃₀O₂(NHPh)₂. [55°]. Got by heating the acid with aniline at 190°. Colourless laminæ (from alcohol), insol. water, NH,Aq, and HClAq

ROCCELLININ C₁₈H₁₆O, Occurs in Roccella tinctoria (Stenhouse, A. 68, 69). Silky needles (from alcohol), nearly insol. cold alcohol and ether. Its solutions in alkalis are not coloured by air. Not decomposed by boiling KOHAq. Does not pp. metallic salts.

ROCHELLE SALTS. Potassium-sodium tartrate; v. TARTRATES.

ROSANILINE C20H21N3O i.e.

 $([4:1]C_6H_4(NH_2))_2C(OH).C_6H_3Me(NH_2)[1:3:4].$ Tri-amido-di-phenyl-tolyl-carbinol. Formed by oxidation of a mixture of aniline, o-toluidine, and p-toluidine (Hofmann, J. pr. 77, 190; 87, 226; Pr. 12, 2; Mühlhäuser, D. P. J. 266, 455, 503, 547). SnCl₄, As₂O₅, HNO₅, Hg(NO₅)₂, FeCl₅, and nitrobenzene may be used as oxidising agents. For the production of a red colouring matter it is essential that there should be present a base with methyl in the p-position to amidogen (p-toluidine, (1,3,4)-xylidine, or mesidine), and also a base with H in the p- position to amidogen (e.g. aniline, o-toluidine, or (1,3,2)xylidine) (Rosenstiehl, C. R. 82, 415; 94, 1319; 95, 238; 98, 433; A. Ch. [6] 2, 360). When nitro-benzene is used as oxidising agent it appears to be itself reduced to benzene and

nitrogen, and not to take part in the formation ! of rosaniline. Thus, when chloro-nitro-benzene, nitro-aniline, or di-nitro-benzene is used, a substituted rosaniline (e.g. chloro-rosaniline) is not formed (Lange, B. 18, 1918). Colourless needles or plates, obtained by ppg. a solution of one of its salts with NH₃Aq and recrystallising. Nearly insol. water, sl. sol. NH,Aq, m. sol. alcohol, insol. ether. Turns red in air. Dyes silk and wool in acid, neutral, and even in slightly ammoniacal solutions.

Reactions .- 1. Reduced by ammonium sulphide or powdered zinc to tri-amido-di-phenyltolyl-methane (leucaniline) (Hofmann, Pr. 12, 2).--2. Yields alkyl derivatives on heating with alkyl iodides and phenyl derivatives on heating a salt with aniline (Hofmann, Pr. 13, 9). Benzyl chloride, methyl iodide, and MeOH form $C_{20}H_{10}(C,H,)_sN_sMeI$, crystallising in lustrous green needles (Hofmann, B. 6, 263).—3. The hydrochloride, heated with water at 235°, forms NH, Cl, phenol, red crystals of C₂₀H₂₀N₂O₂ [176°], and colourless crystals, which turn red in air and then crystallise from alcohol in red needles C₂₀H₁₉NO₃ (Liebermann, B. 5, 144; 6, 951).— 4. The hydrochloride, heated with dilute HClAq at 240°, yields aniline and toluidine (L.).-5. Water at 270° forms di-oxy-benzophenone, di-amido-phenyl-tolyl-ketone, oxy-amido-phenyltolyl-ketone, phenol, and NH, (Liebermann, B. 6, 951; 11, 1435; 16, 1927).—6. Potassium cyanide added to rosaniline acetate in alcohol yields C21H20N4, a white crystalline powder, sol. HClAq, and reppd. by NH3Aq. It separates from alcohol in monoclinic crystals. It forms crystalline salts .-- 7. A solution of a rosaniline salt is decolourised by SO₂, and the solution is turned red, and afterwards violet, by aldehydes, and gradually deposits copper-coloured scales of alkylidene derivatives (Schiff, Bl. [2] 7, 518). C.H. CH: C.O.H., N., cenanthol forms Thus yields B'HAsO₂ B'H,PtCl, which and 8. Œnanthol reacts with rosaniline acetate in the cold, forming $(C_7H_{14})_s(C_{20}H_{16}N_s)_2$ (Schiff, Bl. [2] 5, 291).—9. By exhaustive chlorination it yields CCl, and per-chloro-benzene (Merz a. Weith, B. 16, 2876).—10. Boiling Ac₂O yields a tetra-acetyl derivative as a reddish amorphous powder. The same body is got by treating triacetyl-tri-amido-di-phenyl-tolyl-methane K₂Cr₂O₇ and HOAc (Renouf, B. 16, 1303). A mono-acetyl derivative C20H18AcN3, got by heating rosaniline hydrochloride with acetamide, is reddish-brown and yields dark-blue B'HCl with metallic lustre (Beckerhinn, J. 1870, 768).-11. May be sulphonated by H₂SO₄ containing SO₅ at 130° (Schoop, *Chem. Zeit.* 11, 572).—12. By diazotisation followed by reduction with tin and HCl in the cold it is converted into 'roshydrazide,' the hydrochloride of which forms green crystals, and dyes cotton brownish-red (Ziegler. B. 20, 1557).—13. Br forms a tetra-bromo-rosaniline (?) crystallising from benzene in prisms (Caro a. Graebe, A. 179, 203).—14. Aldehyde forms a blue product which is converted by Na₂S₂O₂ into aldehyde green C₂₂H₂₇N₂S₂O, an amorphous green mass, sol. alcohol and ppd. by ether (Cherpin; Usèbe, J. pr. 92, 337; Hofmann, B. 3, 761; Gattermann, B. 22, 227). Salts.—These are derived from the anhy-

drous base C,H,N:C(C,H,.NH2)2.—C,H19N2HČl.

Red crystals with green lustre. Insol. ether, sl. sol. water, m. sol. alcohol. Its solution in water or alcohol is crimson. Dyes wool and silk magenta. The absorption-spectrum has been studied by Hartley (C. J. 51, 169).—C₂₀H₁₉N₃3HCl. Ppd. by conc. HClAq. Yellowish-brown needles, more sol. alcohol and water than B"HCl .-Decomposed by much water, forming B"HCl.-C₂₀H₁₉N₃4HCl or (C₇H₆NH₃Cl).CCl(C₆H₄NH₃Cl)₂. Got by passing dry HCl into rosaniline powder (Rosenstiehl, Bl. [3] 9, 122). Reddish-brown and hygroscopic, forming a magenta solution. Fumes in air, giving off HCl. After keeping it is not in air, giving off HCl. After keeping it is not completely soluble. $-C_{20}H_{1p}N_34HBr$. Resembles the preceding salt. $-C_{20}H_{1p}N_3+HBr$. According to the preceding salt. $-C_{20}H_{1p}N_3+DAc$. Large crystals with green metallic lustre; v. sol. water and alcohol. $-(C_{20}H_{1p}N_3)_2H_2PtCl_y$. Amorphous. $-(C_{20}H_{1p}N_3)_23H_2PtCl_y$. $-(C_{20}H_{1p}N_3)_23H_2PtCl_y$. Green needles, v. sol. water. $-(C_{20}H_{1p}N_3)_2H_2NG_1$ (dried at 130°). Crystals with green lustre. Does not form an alum with green lustre. Does not form an alum view. $Al_2(SO_4)_3$ (Wood, C. N. 38, 1).— $C_{20}H_{10}N_3C_6H_3N_3O_7$. Red needles, very slightly sol. water. $(C_{20}H_{19}N_3)_2H_2C_2O_4$ aq. — Tannate: ppd. by adding tannin to an aqueous solution of a salt of rosaniline. Red mass, insol. water, sol. alcohol and HOAc (E. Kopp, J. 1862, 694) .-Aurinate: Got by heating equivalent quantities of rosaniline and aurin with alcohol (Dale a. Schorlemmer, C. J. 43, 186). Hexagonal crystals with green lustre (Dyson, C. J. 43, 471). More sol. alcohol than either constituent. Dyes silk magenta.—Phenate C20H10N3C6H6O. Got by heating rosaniline with phenol, dissolving in alcohol, and ppg. with water (Dyson, C. J. 43,

Para-rosaniline C19H19N3O i.e. ([1:4]C₀H₄(NH₂))₃COH. Tri-amido-tri-phenylcarbinol.

Formation. — 1. From tri-nitro-tri-phenylmethane by oxidation by CrO₃ and HOAc followed by reduction of the resulting tri-nitro-triphenyl-carbinol with zinc-dust and HOAc (E. a. O. Fischer, B. 11, 195, 473, 612, 1079, 1598; A. 194, 242).—2. By heating a mixture of aniline and p-toluidine with arsenic acid (Rosenstiehl, A. Ch. [5] 8, 192); formed therefore in the preparation of rosaniline (Graebe, B. 12, 2241).—3. By heating p-nitro-di-amido-tri-phenyl-methane with FeCl, at 170° (O. Fischer, B. 15, 678).—4. By heating p-nitro-benzylidene bromide wi aniline (Zimmermann a. Müller, B. 17, 2936). 5. Hy heating aurin with NH3Aq at 180° (Dale a. Schorlemmer, B. 10, 1016).-6. By heating di-amido-di-phenyl-methane (from aniline and formic aldehyde) with aniline, aniline hydrochloride, and nitro-benzene or arsenic acid

Properties. — Crystalline plates. colourless; sl. sol. water; absorbs CO, from air. Its solution in conc. HClAq is coloured grassgreen by a crystal of KClO.

Reactions.—1. Yields o-toluidine and diamido-benzophenone [237°] on boiling with HClAq for a long time (Wichelhaus, B. 19, 110). 2. HIAq at 190° forms aniline and p-toluidine 3. Nitrous acid forms the tri-diazo-derivative, which on boiling is converted into aurin. 4. Reduced by zinc-dust and HClAq to tri-amido-triphenyl-methane.-5. KCy added to its hydrochloride in presence of alcohol forms 'hydrocyanpararosaniline ' C, H, N, HCN, crystallising from hot alcohol in colourless prisms, forming B'H₃Cl₂2aq.—6. Pararosaniline (5 g.) is slowly converted by conc. HClAq (55 g.), water (55 g.), and aldehyde (22 g.) in the cold into aldehydeblue, a dark-blue hygroscopic powder, ppd. by adding NaCl to the product after dilution with water (Gattermann, B. 22, 227). Aldehyde blue yields, on distillation, a solid base C₁₀H₂N. Salt.—C₁₉H₁₈N₃Cl. S. 24 at 9°.

Tri-methyl-rosaniline v. TRI-METHYL-TRI-AMIDO-DI-PHENYL-TOLYL-CARBINOL.

Hexa-methyl-pararosanilinev. Hexa-methyl-TRI-AMIDO-TRI-PHENYL-CARBINOL

Phenyl-rosanilines v. HEXA-PHENYL-TRI-AMIDO-TRI-PHENYL-CARBINOL and TRI-PHENYL-TRI-AMIDO-DI-PHENYL-TOLYL-CARBINOL.

ROSEINE. A name for salts of Rosaniline. **ROSEMARY OIL.** S.G. 15 .905. $\alpha = +42'$ to +9° 48' in 100 mm. (Schimmel, Ph. [3] 21, 940; 22, 328). Obtained by steam-distillation from Rosmarinus officinalis (Kane, Trans. Irish Acad. 18, 135; Lallemand, A. 114, 197; A. Ch. [3] 57, 404; Gladstone, J. 1863, 549; Montgolfier, Bl. 1876, 18; Bruylants, J. Ph. [4] 29, 508). Contains a terpene (165°), dextrorotatory camphor, borneol, and cineol C10H18O (176°-179°) (Weber, A. 238, 89). The cineol forms a hydrochloride $(O_{\circ}H_{18}O)_{2}HCl$, a tetra-bromide [124°], and the iodide $O_{\circ}H_{18}I_{2}$ [79°].

ROSE OIL. Obtained by steam-distillation

from rose leaves. Contains a stearoptene C16H34 [34°] (365°) which is not attacked by chromic acid mixture and appears to be a paraffin (Markownikoff, B. 23, 3191; J. pr. [2] 48, 311; cf. Blanchet, A. 7, 154; Gladstone a. Dale, C. J. 17, 1; Salkülünski, N. R. P. 24, 129; Bauer, D. P. J. 204, 253; Flückiger, Z. [2] 6, 126; D. P. J. 204, 258; Fluckiger, Z. [2] 0, 120; Panajotoff, B. 24, 2700; Eckart, B. 24, 4205). Rose oil also contains rhodinol (q.v.). Bulgarian oil of roses contains rose of $C_{10}H_{20}O$ (225° cor.), which is oxidised by KMnO, to $O_{10}H_{10}(OH)$, (240°) S.G. $\frac{9}{6}$ 1·0445; $\frac{9}{6}$ 1·0343; and is reduced by HI to $O_{10}H_{22}$ (159°) S.G. $\frac{9}{6}$ ·7700; $\frac{39}{8}$ ·7554 (Ladenburg, J. pr. [2] 48, 293).

ROSEOCOBALT SALTS v. COBALTAMINES,

vol. ii. p. 226.

ROSEOCHROMIUM SALTS v. CHROM-AMMO-NIUM SALTS, vol. ii. p. 160.

ROSEORHODIUM SALTS v. RHODIUM-AMMO-

NIUM COMPOUNDS, this vol. p. 407.

ROSEWOOD OIL. The oil got by steam-distillation from the wood of Convolvulus scoparius contains a terpene C10H16 (249°) (Gladstone a. Dale). Rosewood also contains a black resin C₂₁H₂₁O₄ [95°] S.G. 15 1.266, which is insol. water, sol. alcohol, and gives a blood-red colour with H₂SO₄ (Terreil a. Wolff, Bl. [2] 33, 435).

ROSINDOLES. Red colouring matters,

ROSINDOLES. Red colouring matters, greatly resembling rosaniline, formed by heating indoles with BzCl, and by oxidation of benzylidene derivatives of indoles (E. Fischer a. Wagner, B. 20, 815). Thus methyl-ketole yields dimethyl rosindole, C. H. Which may possibly be ÇPh(C,H,NH)

NC,H,

ROSINDONE C.H. CO.CH: C.NPh C.H.

Rosindulone. [262°]. Formed by heating rosinduline with conc. HClAq at 170° under pressure (Fischer a. Hepp, A. 256, 288; 262, 244).

Formed also from oxy-naphthoquinone and phenyl-o-phenylene-diamine (Kehrmann, J. pr. [2] 43, 269; B. 24, 586). Red hexagonal tables, insol. hot water, m. sol. boiling alcohol, forming a scarlet solution with brick-red fluorescence. Reduced to naphthophenazine by distilling in a current of H over zinc-dust. Converted by p-toluidine at 130° into p-tolyl-rosinduline. Oxidised by CrO₃ to resindenic acid C₂₂H₁₄N₂O₃, which crystallises in colourless needles, sl. sol. alcohol, insol. water, v. sol. ether.

ROSINDULINE C22H15N, i.e.

-C:N C₆H₄<C(NH).CH:C.NPh>C₆H₄. [199°]. Formed by heating benzene- azo- (a)-naphthylamine with aniline and alcohol at 1650 under pressure (Fischer a. Hepp, A. 256, 236). Formed also by heating phenyl-o-phenylene-diamine with oxy-naphthoquinone imide, HOAc, and alcohol at 100° (Kehrmann, J. pr. [2] 43, 269; B. 24, 587). Reddish-brown plates, v. sol. alcohol and ether, insol. water. Conc. H2SO, forms a green solution, becoming red on dilution. The alcoholic solutions exhibit reddish-yellow fluor-

escence.—B'HCl 3 aq: red needles.

Rosinduline. This name has also been given to the phenyl derivative of the preceding body

(v. vol. iii. p. 9).

ROSOLIC ACID v. TRI-OXY-DI-PHENYL-TOLYL-CARBINOL and TRI-OXY-TRI-PHENYL-CARBINOL.

ROTTLERIN C₁₁H₁₀O₃? [200°]. A yellow substance occurring in the fruit of Rottlera tinctoria, which grows in India (Anderson, N. Ed. P. J. [2] 1, 300; Jawein, B. 20, 182). It is identical with Mallotoxin (q. v.).

RUBAZONIC ACID C₂₀H₁₇N₀O₂. Formed by

oxidation of oxy-amido-phenyl-methyl-pyrazole (Knorr, A. 238, 137). Got also by boiling pyrazole-blue with NH, Aq. Prepared from the phenyl-hydrazide of oxy-phenyl-methyl-pyrazole by reduction with zinc-dust and HOAc, followed by oxidation with FeCl, (Buchka a. Sprague, B. 22, 2551). Insol. water and dilute acids, v. sol. ether and benzene. Its solutions in alkalisare deep violet, but on boiling become yellow.

RUBERYTHRIC ACID C₂₄H₂,O₁₄ i.e. C₁₄H₄O₂(OH).O.C₁₂H₁₄O₃(OH),. Rubianic acid. [260°]. Obtained from madder-root by extracting with boiling absolute alcohol (Rochleder, A. 80, 321; 82, 205; Schunck, A. 66, 176; Liebermann a. Bergami, B. 20, 2241). Lemon-yellow needles. v. sol. hot water, v. sl. sol. alcohol, insol. ether. Its alcoholic solution is yellow. Alkalis form blood-red solutions. It does not ferment with yeast. Boiling dilute acids split it up into alizarin and sugar. Ac,O and NaOAc yield

 $C_{28}H_{26}\Delta c_0O_{14}$, crystallising in yellow needles $[230^\circ]_{-} - C_{28}H_{28}KO_{14}$.—BaA'₂ aq.

RUBIAN $C_{28}H_{34}O_{18}$. A glucoside in madder root, split up by acids, alkalis, or the madderferment (erythrozym) into glucose, alizarin, and other bodies (Schunck, P. M. [4] 12, 200, 270; J. pr. 59, 465). Amorphous resin, v. sol. water, m. sol. alcohol, insol. ether. Its solutions are very bitter. Yields ruberythric acid on treatment with baryta. It is probably a mixture of

ruberythric acid and other bodies.

RUBIANIC ACID v. RUBERYTHRIC ACID.
RUBIDINE. A general name for homologues of pyridine of the formula C, H, N, e.g. DI-METHYL-ISOBUTYL-PYRIDINE.

Rubidine C₁₁H₁₇N. (280°). S.G. ²² 1·017. Occurs in coal tar (Thenius, Rep. Chim. app. 4, 181). Liquid, sl. sol. water, miscible with alcohol and ether. Its salts redden in the air. Bleaching-powder gives a red colour. Colours acidified fir-wood red. — B'₂H₂PtCl₄: reddish

crystalline powder.

RUBIDIUM. Rb. At. w. 85·2. Mol. w. unknown. Melts at c. 38·5° (Bunsen, A. 125, 367). S.G. c. 1·52 (B., l.c.). Refraction-equivalent 12·1 (Gladstone, Pr. 18, 49); 11·6 (Kannonikoff, J. R. 1884 [1] 119). Chief lines in emission-spectrum are two in the red, Rb, and Rb, 7800 and 6297; and two in the violet, Rb, and Rb, 4216 and 4206 (Kirchoff a. Bunsen, P. 118, 337; Lecoq de Boisbaudran, Spectres lumineux [1874] 46; v. also B. A. 1884. 438). Beketoff (J. R. 1888. 363) gives [Rb²,O] = 94,900.

Occurrence.-Never free. Salts are widely distributed, but in very small quantities, along with salts of Cs, and frequently also salts of Li, K, and Na. Many lepidolites contain Rb salts amounting to c. 24 p.c. Rb (K. a. B., P. 113, 337; 119, 1; Grandeau, A. Ch. [3] 67, 155; Allen, Am. S. [2] 34, 367). Traces of Rb salts have been found in specimens of orthoclase (Erdmann, J. pr. 86, 377, 448; Laspeyres, A. 134, 349; petalite (Grandeau, l.c.); micas (Schrötter, J. pr. 85, 458); basalt (Engelbach, A. 135, 126); carnallite (Erdmann, l.c.); in alum obtained on the island of Volcano (Cossa, Acc. dei Lincei, 11, 9); saltpetre (Dieulafait, C. R. 98, 1545). Small quantities of Rb salts, varying from c. ·0002 to c. ·03 g. RbCl per litre, occur in mineral springs (v. Bunsen, A. 122, 347; Grandeau, A. Ch. [3] 67, 155; Schrötter, J. pr. 85, 458; Redtenbacher, J. pr. 85, 458; Böttger, J. pr. 89, 378). Traces of Rb salts have been found in sea water (Sonstadt, C. N. 22, 25, 44); in fumaroles giving out boric acid (Hofmann, J. C. T. 1863. 354); in coffee, tea, cocoa, and tobacco (Grandeau, l.c.); in the ashes of beetroot (G., l.c.; Lefèvre, C. R. 55, 430); in the ashes of various plants (von Than, A. Suppl. 2, 84); in normal human urine (Schiaparelli a. Peroni, G. 10, 390). According to Lockyer a well-marked Rb line occurs in the solar spectrum (Pr. 27, 279).

Historical.-In 1860 Bunsen applied the method of spectroscopic analysis, then being worked out by him along with Kirchoff, to the examination of the residue obtained by evaporating very large quantities of the water from a mineral spring at Dürkheim. Two red lines appeared which were not known to belong to any element. By further investigation Bunsen separated the chloride of the element, and on electrolysing the molten chloride he obtained metallic globules, which rose to the surface and took fire. By strongly heating an intimate mixture of the carbonate of the new element with finely divided charcoal and distilling into a receiver under rock-oil, Bunsen obtained the metal in some quantity. The name rubidium was given because of the characteristic lines in the red part of the spectrum (ruber = dark red) (v. Bunsen, A. 113, 337; 122, 347).

Formation.—1. By distilling the mixture of Rb₂CO₂ and C obtained by carbonising Rb.H tartrate (Bunsen, l.c.; Setterberg, A.211, 100).—2. By distilling a mixture of RbOH and Al (Beke-

toff, J. R. 1888. 363).—3. By passing a strong electric current through molten RbCl, the positive electrode being graphite, and the negative an iron wire, globules of Rb rise to the surface, and there take fire. If the negative electrode is surrounded with H (v. Lithium, vol. iii. p. 148), a smalt-blue substance is formed, which diss ves in water with formation of RbOH and H; this blue solid is probably a lower chloride than RbCl, perhaps Rb₂Cl (Bunsen, l.c.).

Preparation.—The saline residues from the manufacture of lithium compounds from Saxon lepidolite (at Struve's works in Leipzig) were found by Bunsen (l.c.), Heintz (J. pr. 87, 310), and Erdmann (J. pr. 86, 294) to be rich in Rb salts. The specimen examined by Bunsen contained 19.75 p.c. RbCl. Cossa (B. 11, 811) recommended the natural alum from the island of Volcano as a good raw material for preparing

Rb salts.

1. Bunsen's method (A. 122, 351) for separating Rb is based on the different solubilities in water of the platinochlorides of K, Rb, and Cs; these solubilities at 17° are in the ratio of 15:2:1. One kilo. of the saline residue from the preparation of Li compounds from Saxon lepidolite is dissolved in 2.5 kilos. water, and ppd., when cold, by 30 g. Pt in aqua regia. After settling the liquid is drawn off, and the pp. is boiled 25 times in succession with small quantities of water (1.5 kilos. in all), each portion being poured into the liquid which was drawn off from the pp. by the PtCl4. The 4 kilos. of liquid and washings are evaporated to 2.5 kilos. While this evaporation is proceeding, the washed platinochloride pp. is dried at 100°, heated nearly to redness in a stream of H until a mixture of Pt with RbCl, and perhaps a little CsCl, remains (if the solid melts reduction is incomplete); and the RbCl is dissolved out in hot. water. The Pt which remains is dissolved in aqua regia, and added to the liquid which has been evaporated. The pp. is washed with small quantities of water as before, dried, and reduced in H, the supernatant liquid and washings being evaporated. The reduced Pt is dissolved and added to the liquid, and so on. These processes are repeated seven or eight times. From 1 kilo. material Bunsen obtained .125 kilo. RbCl, containing 8-4 p.c. KCl and a little CsCl. To separate KCl, the approximately pure RbCl is dissolved in water (36 g. in 1 litre), heated to boiling, and a boiling solution of PtCl₄ (30 g. Pt in aqua regia, diluted to 1 litre) is added; the pp. is washed repeatedly with water at 40°-50°, then dried, and reduced in H; the RbCl is dissolved in water, and the Pt in aqua regia, and these processes are repeated until the RbCl shows no trace of the red lines of K in the spectroscope. Finally, Bunsen separated CsCl by transforming the chlorides into carbonates, and treating the powdered salts with boiling absolute alcohol until the insoluble RbCl ceased to show the blue lines of Cs in the spectroscope (RbCl is insoluble, while CsCl dissolves, in absolute alcohol). This process has been modified by Heintz. (J. pr. 87, 310), Grandeau (A. Ch. [3] 67, 155), Piccard (J. pr. 86, 449), Böttger (J. pr. 89, 878), Schrötter (J. pr. 85, 458), de Boisbaudran (Bl. [2] 17, 551), and others.

2. Redtenbacher's method (J. pr. 95, 148) is

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based on the different solubilities of the alums of K, Rb, and Cs; the solubilities in water, at 17°, are as 22:4:1. The method has been modified by Stolba (J. pr. 99, 49), Godeffroy (A. 181, 176), Petersen (D. P. J. 224, 176), Cossa (B. 11, 811), Setterberg (A. 211, 100), Robinson a. Hutchins (C. N. 49, 253). The mother-liquor from which Li₂CO₃ has been separated by Schrötter's method (vol. iii. p. 149) is evaporated with H₂SO₄Aq, with addition of Al₂(SO₄)₃, whereby Rb and Cs alums are obtained mixed with K alum. Or lepidolite may be decomposed by heating with H2SO4 and CaF2, the residue boiled with five or six volumes water, KOH added equal to 5 p.c. of the quantity of lepidolite used, the whole boiled with water, and filtered hot, when the alums crystallise out on cooling (Stolba, l.c.). The alums are dissolved in half their weight of boiling water, half as much cold water is added, and after stirring the whole is let cool to 45°, when almost all the Rb and Cs alums separate, leaving much K alum in solution. The solution in water, and cooling, are repeated, temperature being allowed to fall to 40°; the alum which separates is dissolved in water equal to half the weight of the original quantity of mixed alums, boiled for some time, and allowed to cool to 60° while being stirred. The crystals which separate are almost pure Cs alum; the mother-liquor, when evaporated, yields almost pure Rb alum.

To prepare the metal, RbCl or Rb alum can be transformed into RbOH, and Rb prepared therefrom; or the RbOH may be dissolved in the proper quantity of tartaric acid to form RbH tartrate, which may then be charred, and Rb obtained from the mixture of Rb₂CO₂ and C

so formed.

To prepare RbOH, the RbCl is converted into Rb₂SO₄ by H₂SO₄Aq, the Rb₂SO₄ is dissolved in water, boiled, BaOAq is added, the liquid being kept boiling. BaSO, is removed by decantation, and the liquid is evaporated to dryness in a silver dish as quickly as possible. RbOH is obtained from Rb alum by dissolving in hot water, ppg. Al₂O₃ by slight excess of NH, Aq, filtering hot, evaporating to dryness in a Pt dish, strongly heating till (NH₄)₂SO₄ is removed, and then proceeding to decompose the Rb₂SO₄ by BaOAq as already described. RbOH is mixed with Al clippings in the ratio 2RbOH:Al (=1 part Al to 7.5 parts RbOH), and the mixture is heated to full redness in a furnace, in an iron cylinder, connected by an iron tube with a glass receiver. The Rb distils over, and appears like Hg in the receiver. As H is given off, the Rb is surrounded by a non-oxidising atmosphere (4RbOH + 2Al = Rb₂O.Al₂O₈ + 2Rb + 2H₂). From 28 to 33 p.c. of the Rb in RbOH is obtained (Beketoff, J. R. 1888. 363; abstracts in C. J. 58, 108, and B. 21, Ref. 424). For a description of the method of preparing Rb from Rb₂CO, and C v. Bunsen (A. 125, 367) and Setterberg (A. 211, 100). An explosive compound similar to the compound of K and CO (this vol. p. 300) is liable to be formed.

For a method of separating Rb from Cs and K, based on the formation of double compounds of RbCl with SbCl, and with SnCl, v. Muthmann (B. 26, 1019 [1893]).

Properties .- A very lustrous, white metal,

with an extremely faint tinge of yellow. Soft as wax, even at -10° ; melts very easily (c. 38.5°). Lighter than water. Bunsen (A. 125, 367) made one determination of S.G., which gave 1.52, but the result is not final. Heated in absence of O, the metal distils below redness, forming a blue vapour. Rb dissolves in liquid NH₃ (Seeley, C. N. 23, 169). When exposed to air Rb is at once covered with a greyish-blue film, which is supposed to be a suboxide; much heat is produced, and the metal soon takes fire. Decomposes cold water very rapidly, with evolution of H, which takes fire. Rb is the most positive element next to Cs. It closely resembles K in its chemical relations (v. Alkalis, METALS of the, vol. i. p. 114). The at. w. of Rb has been determined (1) by determining Cl in RbCl by Bunsen (P. 113, 339; 115, 584 [1861]), Piccard (J. pr. 86, 454 [1862]), and Godeffroy (A. 181, 189 [1875]); (2) by determining V.D. of RbCl and RbI at c. 1200° (Scott, Pr. E. 14, 410). The S.H. of Rb has not been determined; but from observations of the molecular heats of RbCl and Rb₂CO₃, and comparisons of these with those of the corresponding salts of metals that have normal atomic heats, it is probable that the atomic heat of Rb is c. 6.4, and, therefore, that the S.H. is c. .075. The atom of Rb is monovalent in the gaseous molecules RbCl and RbI.

Reactions and Combinations.—1. Rapidly oxidises in air; much heat is produced, and the metal ignites. Beketoff (J. R. 1888. 363) gives [Rb',O] = 94,900.—2. Decomposes cold water, with rapid evolution of H and production of RbOHAq; Beketoff (l.c.) gives [Rb,H*OAq] = 96,400, with formation of RbOHAq and H.—3. Burns in vapour of chlorine, bromine, iodine, sulphur, and arsenic, forming compounds with

those elements (Bunsen, l.c.).

Detection and Estimation. — Phosphomolybdic acid gives a yellow pp. insol. acids (Debray, Bl. [2] 5, 404). Silicotungstic acid (SiW₁₂O₁₂H₈.xH₂O) ppts. Rb salts, but not salts of K (Godeffroy, B. 9, 1365). As Cs salts are ppd. by SbCl₃ in conc. HClAq, added to solutions containing conc. HCl (as 6CsCl.SbCl₃), while Rb salts give no pp., Rb can be separated from Cs salts by addition of HClAq and SbCl₃ in HClAq (G., B. 8, 9). Rb is best detected by the spectroscope. The most characteristic lines are in the blue-violet, Rb₂(λ =4206), and Rb₂(λ =4216): 0002 mgm. RbCl can be detected (Bunsen, l.c.). Rb is determined in the form of RbCl. For the separation of Rb from K, and estimation of the Rb, v. Bunsen (l.c.). Bunsen determined RbCl in presence of CsCl by finding the sum of the two chlorides, then estimating the total Cl, and calculating the quantity of each salt.

Rubidium, amalgam of. When conc. RbClAq is electrolysed, using Hg as negative electrode, a solid, white, lustrous, crystalline amalgam is formed. Decomposes cold water rapidly; becomes oxidised superficially in air, and then deliquesces with formation of RbOH. Is electropositive to K amalgam (Bunsen, A. 122, 347).

347).

Rubidium, bromide of. RbBr. Formula probably molecular, from analogy of RbCl and RbI. Formed by burning Rb in Br. Also by saturating HBrAq with Rb₂CO₂, and evaporating.

White, lustrous, regular octahedra. S. 98 at 5°, 104.8 at 16° (Reissig, A. 127, 33). Melts at c. 683° (Carnelley, C. J. 33, 279). Combines with IBr, to form RbBr.Br; and with Br to form RbBr.Br₂ (Wells a. Wheeler, Am. S. [3] 43, 475).

Rubidium, chloride of. RbCl. Mol. w. 120·57. V.D. at 1200° -1500° 69·7 (Scott, Pr. E. 14, 410). Melts at c. 710° (Carnelley, C. J. 33, 279). By burning Rb in Cl. Also by saturating HClAq with Rb₂CO₃, evaporating, and crystallising from water. Also by reducing Rb₂PtCl₆ in H, and dissolving out RbCl in water (cf. Preparation, p. 411). White, lustrous, cubical crystals; unchanged in air; decrepitates White, lustrous, cubiwhen heated, and then melts, and volatilises at a very high temperature. S. 76.38 at 1°, 82.89 at 7°; sol. alcohol (Bunsen, l.c.). Molten RbCl is a good conductor of electricity (Hampe, Chem. Zeitung, 1887. No. 54). Double compounds: RbCl combines with many metallic chlorides; some at any rate of the products, e.g. the platinoand platini-chlorides Rb₂PtCl₄ and Rb₂PtCl₆, are best regarded as Rb salts of acids containing metals and Cl. The following compounds have been prepared: M=RbCl. — 6M.SbCl, (1); owogkCl (6): 6M.BiCl, (1); 2M.CdCl₂, M.CdCl₂.xrH₂O(1); 2M.CrCl₃.aq(2); 2M.CuCl₂(1); M.AuCl₃ (3); 3M.FeCl₃ (1), 2M.FeCl₃.aq(2); 2M.MnCl₂. 3aq (1); 2M.HgCl₂, 2M.HgCl₂. 2aq, M.2HgCl₂, M.4HgCl₂(1); 2M.PtCl₂(4), 2M.PtCl₄(5); 2M.NiCl₂(1); 3M.TlCl, (2); 2M.SnCl₄ (1); 2M.ZnCl₂ (1).

References to above.—(1) Godeffroy, B. 8, 9; (2) Neumann, A. 244, 329; (3) Rosenbladt, B. 19, 2535; (4) Nilson, Bl. [2] 27, 210; (5) Bunsen, A. 122, 347; (6) Muthmann, B. 26,

1019, 1425.

RbCl also combines with IBr to form RbCl.IBr; with ICl, to form RbCl.ICl; with BrCl, to form RbCl.BrCl; and with Br to form RbCl.Br₂ (Wells a. Wheeler, Am. S. [3] 43, 475).

RbCl.Br₂ (Wells a. Wheeler, Am. S. [3] 43, 475).

Rubidium, cyanide of. Setterberg (A. 211, 100) probably obtained a cyanide of Rb by passing dry HCN into an alcoholic solution of RbOH.

Reissig (A. 127, 33) failed to obtain the salt

free from products of decomposition.

Rubidium, ferrocyanide of; v. vol. ii. p. 335. Rubidium, fluoride of, RbF. Formula probably molecular, from analogy of RbCl and RbI. S.G. 3·202 at 16·5° (Clarke, Am. S. [3] 13, 293). Melts at c. 753° (Carnelley, C. J. 33, 279). Forms a double compound with uranyl fluoride, 4RbF.UO₂F₂.6aq (Ditte, C. R. 91, 166).

Rubidium, hydroxide of, RbOH. A white, brittle solid, with slight greyish tinge; very alkaline and caustic; easily sol. water, with production of much heat; sol. alcohol. Formed by decomposing H₂O by Rb; also by boiling Rb₂SO₄Aq till air is driven out, adding BaOAq gradually, keeping the liquid boiling till BaSO₄ oceases to be formed, decanting from BaSO₄, which separates out rapidly, and evaporating in a silver dish. Melts below redness; is not decomposed by heating. When molten attacks Pt. Beketoff (J. R. 1888. 863) gives [Rb,O,H] = 164,800; [Rb²O,H²O] = 69,900.

Rubidium, iodide of, RbI. Mol. w. 211:53. V.D. 110:8 at 1200°-1500° (Scott, Pr. E. 14, 410). Melts at c. 642° (Carnelley, C. J. 33, 279). Lustrous, white, regular octahedra; unchanged in air. S. 187.5 at 6.9°, 152 at 17.4°. Formed by saturating HIAq with Rb_2CO_3 , evaporating, and crystallising from water. Also by heating Rb in I vapour (Reissig, A. 127, 33). RbI combines with I to form $RbI.I_2$ (Wells a. Wheeler, Am. S. [3] 43, 475).

Rubidium, oxides of. No oxide of Rb hasbeen isolated. Rb burns in air, possibly with formation of an oxide and peroxide similar to K₂O and K₂O₄. The greyish-blue film which is formed on the surface of the metal when exposed to the air is perhaps a suboxide. According to Beketoff (J. R. 1888. 363), the thermal value of [Rb',O] is 94,900.

Rubidium, salts of. The principal salts formed by replacing H of oxyacids by Rb are borate, carbonates, chlorate and perchlorate, chromate and dichromate, molybdate, nitrate, oxalate, silicotungstate, sulphate, and thiosulphate (v. Carbonates, Nitrates, &c.).

Rubidium. silicofluoride of. Rb₂SiF₈. White, regular crystals. S.G. 3·338 at 20°. S. c. ·16 at 20°, 1·35 at 100°. By ppg. hot Rb alum solution by CuSiF₈ (Stolba, J. pr. 102, 1).

M. M. P. M. BUBIJERVINE v. JERVINE.

RUE 01L. The essential oil (c. 229°) obtained by distilling Ruta graveolens contains methyl ennyl ketone C₁₁H₂₂O (224°), a ketone C₁₂H₂₁O (232°), and a small quantity of terpene (Gerhardt, C. R. 26, 225, 361; Cahours, C. R. 26. 262; Greville Williams, T. 1858 [1] 99; A. 107, 374; Hallwachs, A. 113, 108; Harbordt, A. 123, 293). By boiling oil of rue with nitric acid (S.G. 1·2) an acid C₆H₁₈N₂O₄ may be obtained (Chiozza, A. 85, 225; Alexejeff, Z. 1865, 736). This acid is a heavy oil, forming KA' and NaA', crystallising from alcohol in greenishyellow tables (Limpach, A. 190, 298).

RUFICOCCIN v. CARMINIC ACID.
RUFIGALLIC ACID v. HEXA-OXY-ANTHRAQUINONE.

RUFIN C₂₁H₂₀O₈. A product of the action of heat on phlorizin (Mulder, Rev. Scientif. 3, 50; Stas, A. 30, 198). Dark-red resin, sol. alcohol, nearly insol. ether and water. Its alkaline solutions are red. Ac₂O forms C₂₁H₁₀AcO₈ (Schiff, A. 156, 5).

RUFIOPINE is Tetra-oxy-anthraquinone. RUFOL v. DI-OXY-ANTHRACENE.

RUTHENAMMINES v. RUTHENIUM-AMMONIUM COMPOUNDS, p. 418.

RUTHENATES v. RUTHENIUM, SALTS OF OXYACIDS OF, p. 417.

RUTHENITES v. RUTHENIUM, SALTS OF

oxyacids of, p. 417.

RUTHENIUM. At. w. 101.4. Mol. w. unknown. Melts at highest temperature attainable by O-H flame, which is above 2000° (Deville a. Debray, A. Ch. [3] 56, 385). S.G. 12.26 at 0° (crystallised) (D. a. D., C. R. 83, 928); older determinations varied from 8.6 to 11.4. Joly (C. R. 116, 430) gives S.G. of Ru melted in an electric furnace as 12.63 at 0° referred to water at 4°. S.H. .0611 from 0° to 100° (Bunsen, P. 141, 1). C.E. at 40° (linear) .00000963 (Fizeau, C. R. 68, 1125).

Occurrence.—In many platinum ores; Ru is a constant ingredient of osmiridium (vol. iii. p. 47), the quantity varying from 3 to 6 p.c. (v. Claus, A. 56, 257; 59, 234; Gibbs Am. S. [2] 29,

427; 81, 63; 34, 341). Wöhler found Ru₂S₃ in laurite, a rare mineral occurring in Borneo and

in Oregon (J. pr. 98, 226).

In 1828 Osann (P. 13, 283; 14, 329; cf. 64, 197) thought he had discovered three new metals in the portion of a Pt ore insol. in aqua regia; to these metals he gave the names polin, pluranium, and ruthenium (from Ruthenia = Russia). In 1845 Claus (A. 56, 257; 59, 234; cf. 63, 359) examined the substance called ruthenium oxide by Osann, and found it to consist chiefly of SiO₂, TiO₂, Fe₂O₃, and ZrO₂; from this material Claus isolated a new metal, to which he assigned Osann's name, ruthenium.

Formation. — 1. By reducing Ru₂O₃ by heating in H or C.—2. By strongly heating (NH₄)₂RuCl₈, (NH₄)₂RuCl₂, Ru(NH₃.NH₃Cl)₂, or Ru(NH₃.NH₃Cl)₂.HgCl₂.—3. By heating RuSn₂ in

a graphite boat in a stream of HCl.

Preparation.—Deville a. Debray (C. R. 82, 926) prepared pure Ru by heating the ordinary metal in a stream of O till every trace of Os was removed as OsO, (vol. iii. p. 646), fusing with KOH and KNO, dissolving in water, saturating with Cl, and distilling off RuO, into KOHAq, by heating in a stream of Cl on a water-bath, ppg. Ru,O, by adding alcohol to the solution of KRuO, heating the pp. in coal-gas till reduced to Ru, melting in a graphite crucible with 5 to 6 parts pure tin, treating with boiling HClAq, washing and drying the crystalline RuSn, which remained, and heating this strongly in a graphite boat in a stream of HCl.-2. Osm-iridium is mixed with 2 parts KNO, and 1 part KOH, and the mixture is heated to redness in a large silver crucible, embedded in MgO in an earthenware crucible; the molten portion is poured off, and the residue is heated with more KNO, and KOH. The fused mass is lixiviated with water; the orange solution, which contains KOH and K salts of HNO₂, HNO₃, H₂OsO₄, and H₂RuO₄, is neutralised by HNO₃Aq, when a black pp. forms containing OsO₂.xH₂O and Ru₂O₃xH₂O. This pp. is filtered off, the filtrate being set aside, and is washed and boiled with HClAq and HNO, Aq in a retort connected with a receiver which is kept very cold; the boiling is continued till a drop of the distillate, when brought alternately into the oxidising and reducing parts of the Bunsen flame, ceases to give a bright flash in the former, i.e. until OsO4 ceases to distil off. The residue, which contains RuCl₂ and RuCl₄, is dissolved in a little water, excess of NH₄Cl is added to the hot conc. solution, the liquid is poured off from (NH₄)₂RuCl₃, evaporated and mixed with more NH₄Cl, when (NH₄)₂RuCl₆ ppts. Excess of NH, Cl is removed from the pps. by washing with alcohol. By strongly heating the pps. Ru is obtained. The solution filtered from the pp. of OsO2.xH2O and Ru2O3.xH2O contains OsO4 and RuO,; it is made strongly acid by HClAq, OsO, is distilled off, the solution is evaporated till most of the KNO₃ crystallises out, the mother-liquor is evaporated to dryness, the residue is dissolved in water, and Ru₂S₈ is ppd. by adding (NH₄)₂SAq and then acidifying; after washing and drying, the Ru₂S₂ is roasted, and the Ru₂O₃ so formed is reduced by heating in H, or by strongly heating in a graphite crucible; or the Ru₂O₂ may be dissolved in HClAq, and NH₄-Ru chlorides obtained from this solution and decomposed by

heat (Claus, J. pr. 85, 129; Carey Lea, Am. S. [2] 38, 83).

For other methods of separating Ru from osm-iridium v. Deville a. Debray (A. Ch. [3] 56, 385; C. R. 83, 927; Fremy, A. Ch. [3] 44, 385; Gıbbs, Am. S. [2] 34, 343; 37, 61). Debray obtained ruthenium in crystals by dissolving the spongy metal in molten lead, heating strongly, and, after cooling, dissolving out the lead (with which Ru does not alloy) in HNO₂Aq (C. R. 90, 1195)

Properties.—A white, lustrous, hard, heavy, brittle metal. As obtained by heating the double ammonium chlorides, Ru forms a white spongy mass; the metal formed by reducing the oxide by H appears in white porous fragments, which can be powdered; after fusion in an electric furnace Ru appears greyish (Joly, C. R. 116, 430). Next to Os, Ru is the most infusible metal; small pieces can be melted at the highest temperature of the O-H flame (D. a. D., A. Ch. [3] 5b, 385); a little RuO₄ is formed and then decomposed to RuO₂ (D. a. D., C. R. 80, 457). Insoluble in all acids if in compact pieces; very finely divided Ru dissolves slightly and slowly in boiling aqua regia. Not attacked by molten KHSO4; oxidised to K2RuO, by fusion with KOH, KNO, or KClO, Ru dissolves in molten Zn or Pb; treatment with HNO, Aq leaves the Ru undissolved. Small quantities of Ru when alloyed with Pt metals soluble in acids go into solution, with the Pt metals, in these acids. Finely divided Ru combines with O, when heated in O or in air.

Ru is closely related to Rh and Pd, and less closely, but very distinctly, to Os, Ir, and Pt;

v. Noble metals, vol. iii. p. 628.

The atomic weight of Ru has been determined (1) by determining the ratio of Ru to KCl in K₂RuCl₃ (Claus, P. 65, 218 [1845]; (2) by analyses of the salts M₂Ru(NO)Cl₃, where M=K, Na, and NH₄ (Joly, C. R. 107, 994; 108, 946 [1889]; (3) by determining V.D. of RuO₄ (Debray a. Joly, C. R. 106, 328); (4) by measuring S.H. of Ru (Bunsen, P. 141, 1). Claus's determinations gave values from 102-2 to 104-8 for the at. wt.

The only compound of Ru whose mol. weight is known in the gaseous state is RuO₄; the valency of the atom of Ru cannot be determined from this datum.

Reactions and Combinations.—1. Finely divided Ru, when heated to redness in air or in oxygen, rapidly absorbs 18.5 p.c. O (Claus, J. pr. 42, 364); after 10 hours, c. 23 p.c. O has combined (D. a. D., C. R. 87, 441); the product is a mixture of RuO₂ and Ru (Claus failed to obtain a lower oxide than RuO₂ in this way). Crystals of RuO₂ are obtained by heating the metal in a tube in a rapid stream of O; according to Debray and Joly (C. R. 106, 1494), RuO₄ is formed and then decomposed (v. Ruthenium Tetroxide, p. 417).—2. Heated in chlorine, a small portion is changed to a chloride; mixed with sodium chloride and heated in chlorine, Na_RuO₄ is formed (Claus, l.c.).—3. Spongy Ru is said to form HClAq when placed in chlorine water; and to produce chlorides in solutions of hypochlorites, O being given off (Schönbein, J. pr. 98, 76).—4. Very finely divided Ru dissolves slightly when kept in boiling aqua regia for some time, RuO₄ being formed.—5. M_RuO₄

soluble in water. is formed by fusing Ru with caustic alkalis, alkali chlorates, or alkali nitrates. Alkaline earth nitrates or chlorates also form ruthenates (v. Ruthenates, p. 417).

Detection.—A few mgms. of the substance to be tested are heated, in a spoon of Pt, with excess of KNO, until the whole melts and ceases to give off gas; when cold, the mass is dissolved in water, when a strongly coloured orange-yellow solution is obtained; a little HNO, Aq is added to this solution, followed by HClAq, and warming till the voluminous black pp., produced by HNO₈Aq, dissolves to an orange-yellow solution; H₂S is passed into this liquid, until it appears black from the presence of suspended Ru sulphide (or oxysulphide); on filtering, an azureblue liquid is obtained, probably containing RuCl₂ (Bunsen, A. 146, 265). For estimation v. Deville a. Debray (C. R. 83, 927).

Ruthenium, alloys of. An alloy with iridium was obtained by Deville a. Debray (A. Ch. [3] 56, 385). A crystalline alloy with tin, RuSn_s, is formed by melting Ru with 10 to 15 parts Sn, and when cold treating with HClAq. The tin is removed, and crystalline Ru remains, by heating, in a graphite boat, in HCl gas (D. a. D., l.c.; v. also C. R. 83, 927). Ru alloys with zinc; much heat is produced (D. a. D., l.c.). Ru dissolves in molten lead, but separates out again on cooling (Debray, C. R. 90, 1195).
Ruthenium, ammonio-salts of; v. Ruthe-

NIUM-AMMONIUM COMPOUNDS, p. 418.

Ruthenium, chlorides of. Two chlorides, RuCl, and RuCl, are obtained by heating finely divided Ru in a stream of Cl. No other chloride has been isolated; the existence of double compounds of RuCl, with alkali chlorides is doubtful. The molecular weights of the chlorides are not known.

RUTHENIUM DIGHLORIDE RuCl₂. (Ruthenous oride. Ruthenoso-chloride.) Prepared by chloride. heating very finely divided Ru in a stream of Cl. Action begins at c. 360° (Joly, C. R. 114, 291) with formation of RuCl₂, which is removed by subliming, and the temperature is kept at dull redness for an hour or two; the residue is powdered, and again heated in Cl; and this treatment is repeated several times (Claus, A. 59, 234). A black crystalline mass; insoluble water, acids, or alkali solutions. The azureblue liquid formed by the action of reducers, such as Zn or H₂S, on solutions of RuCl₂, was thought by Claus (*l.c.*) to contain RuCl₂.

RUTHENIUM TRICHLORIDE RuCl. (Rutheno-, or ruthenoso-ruthenic, chloride. Sesquichloride of ruthenium.) Obtained, as a brown, crystalline, very hygroscopic solid, by dissolving Ru(OH), (Claus, A. 59, 234), or RuO, (Joly, C. R. 107, 994), in HClAq, and evaporating to dryness. Joly (C. R. 114, 291) obtained RuCl, by heating finely divided Ru at 360° to 440° in a mixture of CO and Cl, the Cl being in excess. Claus says that RuCl, is sol. water, with partial decomposition to an insol. oxychloride; and that, on heating, the solution decomposes with separation of a black powder (partly oxychloride, partly RuO,H,), the colouring power of which is so great that 1 mgm. suffices to make 500 c.c. water appear quite black. Joly (C. R. 114, 291) says that RuCl_s, prepared by heating Ru in Cl and CO, is insol. cold water, acid solutions, CCl_s, CS_p, CHCl₃, Et₂O, or PCl₃; it is decomposed slowly by hot water; slowly dissolved by digesting with 50 times its weight of absolute alcohol, in a sealed tube. Solution in alcohol is purple-violet; slowly decomposes in moist air, more rapidly at 60°, giving RuCl₂.OH (v. RUTHENIUM HYDROXY-CHLORIDE, p. 416). RuCl, absorbs NH, to form 2RuCl₂.7NH₃; another compound containing NH₃, viz. Ru₂Cl₄(OH)₂.7NH₃.3H₂O, is formed by throwing RuCl, in small successive quantities into cold saturated NH, Aq and then heating to 40° (Joly, C. R. 115, 1299).

Double salts. Chlor-ruthenites, RuCl₃.2MCl, or M₂RuCl₅; M=NH₄, K, Na (Claus, A. 59, 234; 63, 359). Formed by evaporating RuO,H, in HClAq nearly to dryness, dissolving in water, and adding conc. solution of the alkali chloride; also by heating a mixture of Ru with KCl or NaCl in Cl (Joly, C. R. 107,994). The potassium salt, which seems to be the most definite, is a brownish-violet, crystalline powder, scarcely sol. cold water, somewhat sol. boiling water, insol. alcohol of 80 p.c.

Joly (C. R. 108, 854) obtained ruthenium nitrosochloride RuCl, NO.H.O, by heating RuCl, with a large excess of HNO, Aq, and evaporating at 120°. Heated in vacuo, or in CO₂, at 440°, gives off NO, and leaves a mixture of RuCl₂ and RuO₂. Solution of the nitroso-chloride is not ppd. by alkalis in the cold; but on boiling with enough alkali to combine with 3Cl, a brown gelatinous pp. of nitroso-oxide, Ru₂O₃.2NO.2H₂O, is obtained. For compounds of RuCl, with NO and alkali chlorides, v. Nitrosochlor-ruthenates, infra.

RUTHENIUM TETRACHLORIDE RuCl. (Ruthenichloride. Ruthenic chloride.) Claus (A. 59, 234) supposed that a solution of RuO,H, in HClAq contained this chloride; he also described various alkali chlor-ruthenates, M.RuCl., obtained by adding alkali chlorides to the supposed solution of RuCl. Joly, however (C. R. 107, 994), has shown that Claus' salts contain NO; and he asserts that the compound used by C. and supposed to be RuO, H, itself contained NO. Joly failed to obtain either RuCl, or the salts M.RuCl. The compounds described by C. as M2RuCle, but shown by J. to be M2Ru(NO)Cls, may be called nitrosochlor-ruthenates.

Nitrosochlor-ruthenates, $M_2Ru(NO)Cl_4(RuCl_2.NO.2MCl)$. M = Am, K, Na. These salts were obtained by Joly (C. R. 107, 994) by heating RuCl, in HNO, Aq for some time and adding alkali chlorides, also by adding alkali nitrites to warm solutions of RuCl, in HClAq. According to Joly, the salts described by Claus as M.RuCl, are in reality nitrosochlor-ruthenates. These salts yield NO when heated with CaCO. Solutions of these salts are not ppd. by alkalis in the cold; on boiling with alkali sufficient to combine with 3Cl, a pp. of Ru₂O₃(NO)₂.2aq is slowly formed (J., C. R. 108, 854). The group NO evidently forms part of the acidic radicle of the salts (cf. RUTHENIUM-AMMONIUM COMPOUNDS, p. 418).

Potassium nitrosochlor-ruthenate, K₂Ru(NO)Cl₃. Prepared by fusing Ru with KOH and KNO, dissolving in HClAq, and evaporating somewhat till excess of KCl and KNO, crystallise out; by evaporating the mother-liquor a mixture of K₂RuCl₃ and K₂Ru(NO)Cl₃ is obtained, from which water dissolves the latter salt. Purified by crystallisation from water, wherein the salt is fairly soluble. Black, orthorhombic prisms; transparent when in thin crystals. Solution in water is violet-red.

Ruthenium, cyanide of, and derivatives; v.

vol. ii. p. 346.

Ruthenium, hydroxides of; v. RUTHENIUM,

OXIDES AND HYDRATED OXIDES OF, infra.

Ruthenium, hydroxychloride of, Ru(OH)Cl₂. Described by Joly (C. R. 114, 291) as a solid, obtained by heating, in moist air to 60°, a solution of RuCl, in absolute alcohol. Very solwater; the deep indigo-blue solution slowly deposits RuO₃H₃.

Ruthenium, iodide of, RuI,. This compound is formed, according to Claus (A. 59, 234), as a black pp. by adding KI to K₂RuCl₃Aq.

Ruthenium, nitrosochloride of, Ru.NO.Cl., v. under Ruthenium trichloride.

Ruthenium, nitrosopentoxide of,

Ru₂(NO)₂O₃.2H₂O, v. under Ruthenium pentoxide.

Ruthenium, oxides and hydrated oxides of. Claus described four oxides of Ru: RuO, RuO,, RuO2, and RuO4; and the hydrated oxides Ru2O2.3H2O and RuO2.2H2O. According to Debray and Joly, RuO and Ru2O3 have not been isolated, but the oxides Ru2O3 and Ru4O9 exist, besides RuO2 and RuO4. According to Joly, Claus' RuO2.2H2O contains NO. Ru combines with O when melted in the O-H flame, with formation of RuO2; it is probable that RuO4 is formed, and then decomposed to RuO2+O2.

RUTHENIUM MONOXIDE RuO. (Ruthenous oxide.) According to Claus (A. 59, 234), this oxide is formed, as a dark-grey, metal-like solid, by calcining RuCl₂ mixed with rather more than an equivalent of Na₂CO₃ in a stream of CO₃, and washing with water as long as anything dissolves. Debray and Joly (C. R. 106, 328, 1424)

could not obtain this oxide.

RUTHENIUM SESQUIOXIDE Ru₂O₃. (Ruthenoxide.) When Ru black is strongly heated in a Pt crucible, c. 18·5 p.c. O is rapidly taken up; O is then slowly absorbed till from 23 to 24 p.c. has combined, and a blue-black mass is formed. This blue-black solid is Ru₂O₃, according to Claus (A. 56, 257; 59, 234); according to D. a. J. (l.c.)

it is a mixture of Ru and RuO2.

Hydrated ruthenium sesquioxide Ru₂O₂·3H₂O or RuO₂H₃. (Black ruthenium hydroxide.) A black powder; prepared by ppg. solutions of RuCl₂ by alkali, or by alkali carbonate, phosphate, or borate (Claus, A. 59, 234). Also ppd.from an aqueous solution of Ru(OH)Cl₂ (v. Hydroxychloride, supra). Even after washing for several days the pp. retains c. 1 p.c. alkali Reduced, but incompletely, by H at ordinary temperature. Insol. KOHAq or NaOHAq, sl. sol. NH₄Aq (Claus, Lc.); sol. in acids. The only salts of oxyacids derived from this oxide which have been prepared are Ru(NO₂)₂·3KNO₂ and the corresponding Na salt (Claus, J. 1863, 697; Gibbs, Am. S. [2] 29, 427; 31, 63; 34, 341).

RUTHENIUM DIOXIDE RUO... (Ruthenic oxide.) Formed by strongly heating Ru sulphide, or Ru(SO.),, in air (Claus, A. 59, 234). Ru(SO.), is obtained by ppg. RuCl, solution by H₂S, and oxidising the pp. by HNO₂. RuO₂ is also formed, as a crystalline sublimate, by roasting osm-

iridium containing Ru in a porcelain tube (Fremy, A. Ch. [3] 44, 385). Also obtained by strongly heating finely divided Ru for some time in a stream of O (D. a. D., A. Ch. [3] 56 385; C. R. 83, 927; 87, 441). The whole of the Ru is not converted into RuO, but some metal remains unoxidised (D. a. J., C. R. 106, 1494). At a very high temperature RuO, is formed, and at a lower temperature this is resolved into RuO, and O (D. a. D., C. R. 80, 457; D. a. J., C. R. 106, 1424). D. a. J. (C. R. 106, 328) obtained RuO, partly amorphous and partly crystalline, by heating RuO, to 440°.

Dark-violet, quadratic crystals; isomorphous with rutile (Fremy, l.c.; D. a. J., l.c.). S.G. 7-2 (D. a. D., A. Ch. [3] 56, 385). Insol. acids, sol. molten KOHAq. Easily reduced by H. The only salt of an oxyacid corresponding with RuO₂ which has been isolated in Ru(SQ). (Claus Lc.)

which has been isolated is Ru(SO₄)₂ (Claus, l.c.).

Hydrated ruthenium dioxide RuO₂.5H₂O or RuO₄H₄.3H₂O. The existence of this compound is doubtful. Claus (A. 59, 234) gave this composition to the gelatinous, yellow-brown pp. formed by decomposing K₂RuCl₆Aq by Na₂CO₃; but Joly has shown that Claus' K₂RuCl₆ contains NO (cf. Ruthenium tetrachloride, p. 415). Claus (also obtained hydrated RuO₂ by evaporating Ru(SO₄)₂Aq with KOHAq; but as the sulphate employed was obtained by oxidising Ru sulphide by HNO₂, it may have contained NO. Joly (C. R. 107, 994) gives the formula Ru₂O₃(NO)₂·2H₂O to the brown gelatinous pp. formed by boiling Ru(NO)Cl₃ or K₂Ru(NO)Cl₃ with KOHAq or K₂CO₂Aq. It seems probable that Claus' RuO₂·5H₂O is the same as the nitroso-compound obtained by Joly (cf. infra Hydrated nitroso-pentoxide).

RUTHENIUM PENTOXIDE $\mathrm{Ru}_2\mathrm{O}_5(=\mathrm{RuO}_2\mathrm{\cdot}\mathrm{RuO}_3)$. When $\mathrm{RuO}_4\mathrm{Aq}$ stands for some time in a closed vessel, a black pp. settles down and a black crystalline crust forms on the sides of the vessel, while O is set free and escapes when the vessel is opened. The composition of the pp. and crystalline crust after drying at 100° is given by Debray a. Joly as $\mathrm{Ru}_2\mathrm{O}_5$ (C. R. 106, 328). This oxide is also obtained by allowing a dilute solution of a ruthenate to stand for a long time, or by adding a dilute acid to such a solution. $\mathrm{Ru}_2\mathrm{O}_5$ dissolves in HClAq, giving off Cl; heated

to 360° Ru₄O_p is formed.

Hydrated nitroso-pentoxide Ru₂O₃(NO)₂.2H₂O. A black solid, formed by boiling Ru(NO)Cl₃ or K₂Ru(NO)Cl₃ (v. Nitrosochloride, and Nitrosochlor-ruthenates, p. 415) with sufficient alkali or alkaline carbonate to combine with 3Cl, and drying the pp. at 150° (Joly, C. R. 108, 854). Slowly decomposes when heated to 360° in CO₂, giving Ru₂O₃; above 440° decomposes violently, giving off N oxides; reduced by H below 100°, giving off NH₂. Sol. in acids, forming nitrososalts.

RUTHENIUM NONOXIDE Ru₄O₅ (= 3RuO₂.RuO₃). This composition was given by D. a. J. (C. R. 106, 328) to the black, lustrous, crystalline crust obtained by heating RuO₄Aq to 100°. The same oxide is formed by heating Ru₂O₅ to 860°. Heated to 440° this oxide gives amorphous RuO₂ and O.

RUTHENIUM TRI-OXIDE RuO. (Ruthenic anhydride.) This oxide has not been isolated.

but salts derived from it are known. RuO, is the hypothetical anhydride of ruthenic acid (H2RuO4), which has not been isolated. For the salts corresponding with this oxide v. Ruthenates, under Ruthenium, salts of oxy-ACIDS OF, infra.

RUTHENIUM HEPTOXIDE Ru2O7. (Per-ruthenic anhydride.) This oxide, the hypothetical anhydride of the non-isolated per-ruthenic acid (HRuO₄), has not been prepared. For the salts corresponding with this oxide v. Perruthenates, under RUTHENIUM, SALTS OF OXYACIDS or, infra.

RUTHENIUM TETROXIDE RuO. (Sometimes called per-ruthenic acid, and frequently ruthenic acid.) Mol. w. 165.24. V.D. 83.3 at 100° and 106 mm. pressure (Debray a. Joly, C. R. 106,

Formation.—1. By passing Cl into a solution of the product of fusing Ru with KOH and KNO₃.—2. By the action of KClO₂ and HCl, not aqua regia, on K, RuCl, Aq (Claus, A. 59, 234).-3. By heating finely divided Ru in a stream of O to c. 1000°, and cooling rapidly by means of If the temperature is allowed to fall slowly the RuO, formed decomposes explosively at c.

108° (Debray a. Joly, C. R. 106, 100).

Preparation.—A mixture of 1 part finely divided Ru, with 8 parts KOH and 1 part KNO, is heated to dull redness in a silver crucible till fused; the product, when cold, is dissolved in 16 parts water; the solution is placed in a tubulated retort connected, by a long tube, with a very well cooled receiver (the connecting tube being also cooled), and a rapid stream of Cl is passed into the solution; a considerable amount of heat is produced, and RuO, passes over and solidifies in the receiver (Deville a. Debray, A. Ch. [5] 4, 537). To free the preparation from water D. a. J. (C. R. 106, 328) place it in a tube with fused CaCl₂, contract the tube in the middle, pump out the air, and seal off the tube. RuO, slowly sublimes into the upper part, which is then separated from the rest. The RuO, is finally distilled in vacuo into tubes which have been very carefully cleansed from all traces of organic matter, and dried by heating to redness in a stream of H. RuO, may be obtained in well-formed crystals by subliming in vacuo at the ordinary temperature.

Properties.—(D. a. J., C. R. 106, 328.) Golden yellow crystals, melting at 25.5° to an orange-red liquid, which solidifies slowly to a vitreous solid. Smell resembles that of ozone; the vapour causes coughing, but does not affect the eyes (Claus, J. pr. 80, 282). Sublimes at 3 to 4 mm. pressure on the slightest rise of temperature; the vapour is golden yellow. Decomposes at 106°-107° without boiling. The vapour-pressure of the crystals is almost nil at 0° , 20 mm. at 42°, and 182 mm. at 100.8°. May be distilled in water-vapour containing Cl or HClO. Gradually reduced to RuO, by sunlight (Joly, C. R. 113, 693). Dissolves in water, forming a goldenyellow solution, which gradually decomposes with ppn. of Ru₂O₅.xH₂O. RuO, attacks Hg, and slowly acts on glass. RuO, does not form corresponding salts, either by reacting with acids or alkalis. Moist BuO₄ is very easily reduced, e.g. by contact with paper.

Reactions.—1. Decomposed by heat; no so-

tion below 106°; at 107° sudden decomposition occurs, with a smoky flame and formation of amorphous RuO2 on the walls of the vessel, and crystalline RuO2 from the part that has melted (D. a. J., l.c.). According to D. a. J. (C. R. 106, 100), RuO, is formed when O is passed over spongy Ru at c. 1000°, and the product is rapidly cooled. If cooling is allowed to take place slowly the RuO, decomposes at c. 108°.-2. Dissolves in water; solution keeps unchanged for some time, but slowly deposits Ru,O,.xH,O. Decomposition is more rapid at higher temperatures; at 100° Ru,O,xH,O is formed (D. a. J., l.c.).-3. Solution in water, or moist RuO, is quickly reduced by alcohol and several other organic compounds (Claus, J. pr. 80, 282).—4. With solutions of alkalis forms ruthenates and per-ruthenates, with evolution of O. Alcoholic solution of potash ppts. Ru₂O₃.3H₂O.— 5. Heated with hydrochloric acid, Cl is given off and RuCl_s formed in solution.—6. Hydrogen sulphide gradually throws down a black pp. of an oxysulphide, from RuO, Aq (Claus, l.c.)

Ruthenium, oxyacids of. No oxyacid of Ru has been isolated, but some salts of the hypothetical acids H2RuO4 and HRuO4 are known (v. RUTHENIUM, SALTS OF OXYACIDS OF, infra).

Ruthenium, oxysulphide of. A current of H₂S passed into RuO Aq slowly ppts. all the Ru as a black substance, which is said by Claus (J. pr. 80, 282) to be an oxysulphide, whose composition varies according to the quantity of H.S. passed in. The pp. suddenly glows when dried at 100°, and burns, giving off SO2.

Ruthenium, salts of. Very few salts are known obtained by replacing the H of oxyacids by Ru. The double nitrites Ru(NO.) 3MNO2, where M = K or Na, are the only salts which have been prepared corresponding with Ru₂O₃; and Ru(SO₄)₂ is the only salt of RuO, which has

been isolated; the double sulphite RuSO₃.K₂SO₃ corresponds with RuO.

Ruthenium, salts of oxyacids of. A few salts of the hypothetical acid H.RuO4, and two salts of the hypothetical acid HRuO, have been isolated.

RUTHENATES, $M_2^1RuO_4$. Salts of hypothetical H_2RuO_4 (formerly called ruthenites). These salts are formed by heating Ru, or an oxide of Ru, with an oxide, hydroxide, nitrate, or chlorate of an alkali metal (Claus, A. 56, 257; 59, 234), or of an alkaline earth metal (Deville a. Debray, A. Ch. [3] 56, 385). Ruthenates in solution are easily reduced by organic bodies. with ppn. of black Ru, O, 3H,O; acids added to conc. solutions form per-ruthenates, MIRuO.

Barium ruthenate BaRuO, aq. Obtained by adding RuO, to BaOAq. The solution is at first green, probably from formation of perruthenate; a black pp. is thrown down, which soon changes to the cinnabar-coloured crystal-line Ba salt. This salt is also formed by adding BaCl, Aq to a solution of the K salt (Debray a. Joly, C. R. 106, 1494).

Potassium ruthenate K, RuO, aq (D. a. J., l.c.). 50 g. RuO, melted under water, are slowly added to 70 g. KOH in 500 c.c. water at 60°; temperature is kept at 60° until the evolution of O has ceased, when the solution is evaporated in vacuo; long, black, orthorhombic prisms, reflecting green light, are formed (for crystalline measurements v. D. a. J. (l.c. p. 1497). After drying in vacuo, the crystals absorb moisture and CO₂ from the air. Unchanged at 106° in vacuo; loses H2O at 200°, and may then be heated to 400° without decomposition. Very sol. water; solution is orange yellow; it soon decomposes in air, depositing Ru2O, xH2O, and probably forming KRuO,Aq. Organic bodies dipped into the solution become black from deposition of Ru₂O₃.3H₂O. Acids hasten formation of KRuO, Aq and Ru,O,; from conc. K,RuO, Aq acids ppt. KRuO,, which decomposes to Ru₂O₅ and RuO₄ (D. a. D., C. R. 83, 927).

The magnesium, silver, and strontium salts are obtained by adding MgCl₂Aq, AgNO₃Aq, and SrCl₂Aq respectively to K₂RuO₄Aq. The sodium salt has not been isolated; it cannot be separated

from excess of soda.

PER-RUTHENATES, MIRUO. Salts of hypothetical HRuO. (formerly called *ruthenates*). Obtained by the action of Cl on ruthenates, or of acids on conc. solutions of ruthenates. Alkalis produce ruthenates from solutions of

per-ruthenates.

Potassium per - ruthenate KRuO₄ (Deville a. Debray, C. R. 83, 927). 50 g. RuO₄, melted under water, are slowly added to 60 g. KOH in 250 c.c. water at 60°; O is evolved, and the solution becomes green; after O ceases to come off, the liquid is allowed to cool in a closed vessel; the cold (red) liquid, which contains K₂RuO₄, is poured off, and the crystals of KRuO₄ that adhere to the sides of the vessel are dried in vacuo on an unglazed porcelain plate. Black, opaque, quadratic pyramids (for crystallographic measurements v. D. a. J., C. R. 106, 1496). Unchanged in air after drying in vacuo; decomposes suddenly at 440°, giving off O (? with formation of K2RuO4 and RuO2). reacts in the cold to form RuO4 and KCl. Slightly sol. water; solution is blackish-green and soon decomposes, KOH being formed. Alkalis added to KRuO, Aq produce K2RuO, Aq, with evolution of O.

The sodium salt, NaRuO aq, has been isolated; it forms black crystals. Addition of chlorides of alkaline earths to solution of KRuO, or NaRuO, ppts. MIRuO, not MI(RuO), and

the solutions contain RuO₄.

Ruthenium, sulphides of. No sulphide of Ru has been isolated with certainty. could not obtain a sulphide by heating Ru with S in CO₂ (A. 56, 257; 59, 234). The pps. formed by H2S in solutions of Ru compounds vary in composition, and contain free S. By ppg. RuCl. solution by H2S, and heating the solid in CO2, S and H₂O are given off, and a greyish-black metal-like substance remains, which is perhaps Ru₂S₃ (C., *l.c.*). The pp. which H₂S produces in RuO, Aq is probably an oxysulphide (q.v.). The mineral laurite, from Borneo and Oregon, contains Ru.S. with Os sulphide (Wöhler, A. 139, 116; 191, 374) M. M. P. M.

RUTHENIUM-AMMONIUM COMPOUNDS. (Ruthenammines. Ammoniacal ruthenium bases.) These compounds have not been obtained by treating Ru chlorides with NH, Aq. The starting-point for the compounds described by Claus was RuCl, 2NH, Cl. By treatment with NH, Aq, Claus obtained RuCl, 4NH, 3aq, and from this he prepared various salts RuX.4NH,

where X = monovalent acidic radicle (N. Petersb. Acad. Bull. 1, 97; 2, 158). By evaporating a solution of the base Ru(OH)2.4NH3 in vacuo over H2SO4, Claus obtained a new base, to which he gave the formula Ru(OH), 2NH, 4aq; he did not prepare any salts of this base. Joly (C. R. 107, 994) found that Claus' RuCl, 2NH, Cl is a nitroso-compound, and has the composition RuCl, NO.2NH, Cl. Joly also showed (C. R. 108, 1300) that Claus' RuCl 4NH has the composition RuCl₂NO.OH.4NH₃, and that the salts obtained by Claus belong to the form RuX₂.NO.OH.4NH₃. Whether the base described by Claus as Ru(OH)₂.2NH₃ contains NO or not has not been determined; it is described here provisionally.

The nomenclature adopted in this article

must be looked on as only provisional.

NITROSO-RUTHENI-DIAMMINES,

X.NO.Ru(NH₃.NH₃R)₂; R may or may not be the same as X. (Joly, C. R. 108, 1300; 111, 969.) Described by Claus as ammonium ruthenoso-diammonium compounds, N₂H₄(NH₄)₂Ru.X₂, or ruthenoso-diammines, Ru(NH₃.NH₃X)₂.

Series I. Hydroxy-compounds, OH.NO.Ru(NH₃.NH₃R)₂. The chloride, R = Cl, was obtained (Joly, C. R. 108, 1300) by boiling Ru(NO)Cl₃ (described under RUTHENIUM TRI-CHLORIDE, p. 415) with excess of NH3Aq, until the liquid became golden yellow, and deposited orange-yellow crystals of the salt. The bromide R = Br, and *iodide* R = I, were prepared similarly. The carbonate $R = \frac{1}{2}CO_s$, nitrate R NO_s, and sulphate $R = \frac{1}{2}SO_4$, were prepared by the reaction of the chloride with Ag salts. By reacting on the chloride with moist Ag₂O, Joly obtained a strongly alkaline liquid which probably contained the hydroxide, R=OH. The chloride

forms a double compound with PtCl.

Series II. Chloro - compounds,
Cl.NO.Ru(NH₃.NH₃R)₂ (Joly, C. R. 111, 969). The chloride, R = Cl, was obtained by dissolving the hydroxy-chloride in much HClAq, evaporating, and crystallising from boiling water; it combines with PtCl₄. The bromide and iodide, R = Br and I, were obtained similarly to the

chloride.

Series III. Nitrato - compounds, NO₃.NO.Ru(NH₃.NH₃R)₂. The nitrate, R = NO₃, was formed by the action of AgNO, Aq, or boilling HNO, Aq, on the chloro-chloride (J., l.c.).
Series IV. Sulphato - compounds

Series IV. Sulphato-compounds, SO, 2[NO.Ru(NH₃.NH₃R)₂]. Two sulphates, the normal salt where $R = \frac{1}{3}SO_4$ and the acid salt $2(SO_4.2[NO.Ru(NH₃.NH₃)₂SO_4]).H_2SO_4$, were obtained by the action of H_2SO_4Aq on the chloro-

These compounds give off NH, when heated in KOHAq, and separate Ru nitroso-hydroxide Ru(OH),NO, from which nitroso-salts are ob-

tained by the action of acids.

??RUTHENOS-AMMINES, Ru(NH₂X)₂₂ or ruthenos-diammonium compounds, NH₂(NH₄)Ru.X₂. The only compound of this class which has been isolated is the hydroxide, Ru(NH₃.OH)₂.4aq, obtained by Claus (N. Petersb. Acad. Bull. 1, 97; 2, 158) by evaporating, over H2SO, in vacuo, a solution of the base to which he gave the formula Ru(NH_s.NH_s.OH)_s. As the compound from which Claus obtained the supposed ruthenos-ammine hydroxide has been found to contain NO, it is probable that the product of evaporation was also a nitroso- compound.

M. M. P. M.

RUTIN C₄, H₅₀O₂₈. [above 190°]. S. ·58 at

100°. S. (bolling alcohol) 20. S. (ether) ·29.

Extracted by alcohol or HOAc from the leaves
of the garden rue (Ruta graveolens) (Weiss,
Pharm. Centr. 1842, 903; Bornträger, A. 53,
385; Förster, B. 15, 217). Occurs also in
capers, the flower-buds of Capparis spinosa
(Rochleder a. Hlasiwetz, A. 82, 197; 96, 123;
Zwenger a. Dronke, A. 123, 145), in safflower
(Stein, J. pr. 58, 399; 88, 280), in rose leaves
(Filhol, J. 1863, 594), and in the leaves of the buck-

wheat (Polygonum Fagopyrum (Schunck, C. J. 53, 262). Crystallises from water in pale-yellow needles (containing 3aq). Sol. alkalis, forming a yellow solution, from which it is reppd. by acids. FeCl, gives a dark-green colour. Lead acetate added to its alcoholic solution forms a yellow pp. Pb₃C₄₂H₄₄O₂₃. Does not reduce Fehling's solution. Decomposed by boiling dilute H₂SO₄ into quercetin (1 mol.) and isodulcite (3 mols.).

RUTYL. A name sometimes used for decoyl

C10H19O or decyl C10H21.

RUTYLENE v. Decinene. RUTYLIDENE v. HENDECINENE.

S

SACCHARIC ACID C_eH₁₀O_k i.e. CO₂H.CH(OH).CH(OH).CH(OH).CH(OH).CO₂H. Mol. w. 210. Formed by the action of nitric acid (6 pts. of S.G. 1·15) on dextrose (glucose), and therefore on cane sugar and on milk sugar (Scheele, Opuscula, ii. 203; Trommsdorff, A. 8, 36; Guérin-Varry, A. Ch. [2] 49, 280; 52, 318; 65, 332; Erdmann, A. 21, 1; Hess, A. 26, 1; Thauloff, A. 27, 113; Liebig, A. 30, 313; 113, 1; Heintz, P. 61, 315; 105, 211; 106, 93; 111, 265, 291; A. 51, 185; Tollens, B. 21, 2149). Formed also by oxidation of dextrin, maltose, and dextrose by Br, and subsequent addition of ZnCO₃ (Herzfeld, A. 220, 352), and by the action of Br on glycuronic acid (Thier-

felder, B. 19, 3148).

Preparation.—1. Dextrose (5 g.) is evaporated with HNO₃ (30 c.c. of S.G. 1·15) to a thick syrup. The syrup is dissolved in water (20 c.c.) and neutralised with K₂CO₃; HOAc is then added and the solution evaporated until the acid K salt crystallises out (Tollens, A. 249, 218).—2. Starch (100 g.) is ground up with water (100 c.c.), poured into HNO₃ (500 c.c. of S.G. 1·15), and heated on the water-bath till red fumes begin to come off. The temperature is then lowered and kept at 65° until syrupy. The product is converted as above into the acid K salt (20 g.), which is dissolved in water, neutralised with ammonia, and ppd. by AgNO₃. The Ag salt is then decomposed by HCl (Sohst a. Tollens, A. 245, 4).

Properties.—Brittle deliquescent mass, v. sol. water and alcohol, v. sl. sol. ether. Dextrorotatory, [a]p varying from 8° to 38°. Reduces auric chloride and ammoniacal AgNO₃ (forming a mirror) but not Fehling's solution (Kiliani, B. 14, 2529).

Reactions.—1. Oxidised by HNO₂ to dextrotartaric, racemic, and oxalic acids.—2. Potash at 250° yields acetic and oxalic acids.—3. PCl₃ forms chloromuconic acid C₆H₄Cl₂O₄ [260°] (Bell, B. 12, 1272).—4. Boiling dilute (30 p.c.) HClAq forms dehydromucic acid (S. a. T.; Schrötter, M. 9, 442).—5. HIAq and P at 150° forms some adipic acid (De la Motte, B. 12, 1572).—6. The ammonium salt decomposes at 160° into CO₂, NH₂, and pyrrole (Bell a. Lapper, B. 10, 1961). The ethylamine salt yields, in like manner, ethyl-pyrrole.—7. Phenyl-hydrasine acetate at 100° forms C₆H₂O₄(CO.N₂H₂Ph)₂, a

yellowish substance [210°], insol. water, alcohol, and ether; not decomposed by alcoholic potash (Maquenne, Bl. [2] 48, 721).

Salts.—KHA": trimetric crystals; a:b:c = 1.763:1:2.234. S. 1·1 at 7°.—K₂A": crystal-line crusts, v. sol. water.—(NH₄)HA". S. 1·22 at 15°; 24·4 at 100°. Four-sided prisms.—BaA": minute crystals, v. sl. sol. water.—BaA"3aq: amorphous.—CaA"aq.—SrA"1½aq.—MgA"3aq: crystalline, m. sol. hot water.—ZnA"aq.—CdA".—PbA".—Pb₂C₁₂H₁₄O₁₈.—Pb₂C₁₂H₁₄O₁₈.—Pb₂C₁₂H₁₄O₁₈.—Pb₂C₁₄H₁₄O₁₈.—Pb₂C₁₄H₁₄O₁₈.—Wite pp. Ethyl ether Et₂A". Crystalline mass, v. sol. water and alcohol.—(Et₂A")₂CaCl₂. Prisms,

v. sol. water, sl. sol. alcohol, insol. ether.

Tetra-acetyl derivative of the ethyl
ether C₆H,Ac₄Et₂O₅. [61°]. Monoclinic tables
(from alcohol), insol. Aq, v. e. sol. hot alcohol.

Anhydride C.H.O., i.e.

CH(OH) CH(OH).CO₂H. Sac-

charolactone. [132°]. Formed by allowing syrupy saccharic acid to stand over H₂SO₄ for some days (Sohst a. Tollens, A. 245, 5). Thin plates (from water). Yields pyromucic acid on heating. Reduced in acid solution by treatment with sodium-amalgam to glycuronic acid CO₂H.CH(OH).CH(OH).CH(OH).CH(OH).CH(OH).CH(OH).CHO(Fischer a. Piloty, B. 23, 937; 24, 521).

Di-acetyl derivative of the double

anhydride C10H10Os i.e.

CH(OAc).CH.O.CO CO.O.—CH.—CH(OAc). [188°]. Formed from Ac.O, acid potassium saccharate, and H.SO. (Maquenne, Bl. [2] 48, 720; cf. Baltzer, Bl. [2] 10, 263; A. 149, 238). White plates, v. sl. sol. alcohol and ether.

Amide C₈H₈O₈(NH₂)₂. Amorphous powder. Isosaccharic acid C₈H₁₀O₈. [185°]. [a]_D = 46° at 20°. Formed by careful oxidation of glucosamine hydrochloride by dilute HNO₈ (S.G. 1·2) (Tiemann, B. 17, 246; 19, 1258, 1273). Trimetric crystals (from water), v. sol. water and alcohol, sl. sol. ether. Dextrorotatory. Converted by heat into CO₂ and pyromucic acid. Reduced by HI to adipic acid. Heated in a current of dry HCl it yields furfurane (aa')-dicarboxylic acid. PCl₈ forms C₄H₃ClO(CO₂H)₂, which yields Et₄A'' [44°].—

KHA" \(\frac{1}{2}\text{aq} : \neq \text{sol. water.} -(\text{NH}_4)_2\text{A".} -\text{BaA".} -\text{CaA".} -\text{SrA".} -\text{PbA"} 2\text{aq.} -\text{CuA".} -\text{Ag.A".} \)

Ethyl ether $\text{Et}_2\text{A}''$. [73°]. (250°). [a]_D -+35.5°. Needles, v. sol. water. Yields a tetra-acetyl derivative C₄H₄(OAc)₄(CO₂Et)₂ [47°] v. sol. water and alcohol.

Amide of the anhydride $C_1H_1O(OH)_2(ONH_2)_2$ [226°]. [a]_D = 7.2°.

Anilide of the anhydride. [231°]

Acetyl derivative C,H,(OAc),(CO,H),aq.

[101°]. Needles (from water).

Parasaccharic acid C₆H₁₀O₈. Formed, together with glycyrrhetin, by boiling glycyrrhizin with dilute H2SO4 (Habermann, C. C. 1880, 253). Hygroscopic mass, sol. water and alcohol. Its salts are amorphous.

Metasaccharic acid C,H10Os. The salt CaA" aq, got by boiling an aqueous solution of the dianhydride with CaCO₃, forms crystals, sl. sol. water. KA' and Ag₂C₈H₈O₈ have also been prepared (Fischer, B. 23, 2621). Solutions of metasaccharates turn deep red when exposed to air.

Dilactone $C_6H_6O_6$ i.e.

ÇH(OH).ÇH.O.ÇO 1- Mannosaccharic CO . O.CH—CH(OH) . acid. [68°]. S. 5.5 at 15°. $[a]_D = -202^\circ$ (Fischer, B. 24, 541). Formed by oxidation of arabinose-carboxylic (mannonic) acid $C_{\rm e}H_{12}O_{\rm r}$ by digesting its lactone with HNO, (S.G. 1·2) for 24 hours at 50° (Kiliani, B. 20, 341, 2710; 21, 1422). Long colourless needles (containing 2aq), sl. sol. alcohol, insol. ether. Readily reduces Fehling's solution. Reduced by HI to n-adipic acid. Sodium-amalgam reduces it to mannite. Phenyl hydrazine hydrochloride and sodium acetate solution in the cold form CH(OH).CH(OH) [192°] crys-

CO . O.CH.CH(OH).CO.N₂H₂Ph tallising in minute plates (containing 2 aq), v. e. sol. hot water. Phenyl hydrazine hydrochloride and boiling sodium acetate solution forms C₄H₄(OH)₄(CO.N₂H₂Ph)₂, crystallising in minute

plates [213°], v. sl. sol. water.

Di-acetyl derivative of the dilactone CH(OAc).CH.O.CO CO.O -- CH-CH(OAc) [155°]. Formed by adding a few drops of H2SO4 to a mixture of the dilactone and Ac.O (Kiliani, B. 22, 524). Prisms, v. sol. hot HOAc.

A mide $C_4H_4(OH)_4(CO.NH_2)$ [190°]. Formed from the lactone and NH,Aq. Mono-

clinic tables.

Dextro-metasaccharic acid. d-Mannosaccharic acid $C_0H_{10}O_8$. Salts.—CaA": crystalline powder.—BaA": minute tables, m. sol. water.— SrA".-CdA": minute tables, v. sl. sol. water.

Lactone $C_6H_9O_6$. [180°-190°]. [α]_D = 202° at 23°. Formed by oxidation of mannose, of mannite, or of d-mannonic acid by HNO, (S.G. 1.2) at 50° (Fischer a. Wirthle, B. 24, 539; Easterfield, C. J. 59, 306). Long needles (from alcohol or water), v. sol. hot water. Readily reduces Fehling's solution. Turns yellow on boiling with potash. Phenyl-hydrazine acetate in the cold

forms CH(OH).CH(OH).CON,H.Ph [191°], while at 100° it forms C.H.(OH) (CO.N.H.Ph)

[212°], almost insol. hot water,

Amide C_eH₁₂N₂O₂. [189°]. Formed from the lactone and NH₂Aq in the cold. Crystals.

Inactive metasaccharic acid C.H.O.s. i-Mannosaccharic acid. The salts of this said greatly resemble those of the preceding acid.

Lactone C.H.O. [c. 190°]. Formed by mixing equal parts of the dextro- and lavolactones in aqueous solution. Formed also by oxidising i-mannonic lactone (E. Fischer a. Stanley Smith, B. 23, 2622; 24, 544). The aqueous solution is inactive to light, and gradually becomes acid on standing. Phenyl-hydrazine acetate reacts, forming in the cold CH(OH).CH(OH) CO.O — CH.CH(OH).CON,H,Ph [190°-195°]

and at $100^{\circ} C_{\downarrow}H_{\downarrow}(OH)_{\downarrow}(CON_{2}H_{2}Ph)_{2}$ [220°-225°].

Amide. [185°].

SACCHARIN. The lactone of saccharinic acid (v. infra). The name has also been applied to the imide of SULPHOBENZOIC ACID.

SACCHARINIC ACID C.H.2O. i.e. $CH_2(OH).CH(OH).CH(OH).CMe(OH).CO_2H.$

Formed by boiling an aqueous solution of dextrose, lævulose, or invert sugar with lime (Péligot, Bl. [2] 36, 226; C. R. 90, 1141; Scheibler, B. 13, 2212). On setting the acid free it changes, especially on heating, into the anhydride. The salts are v. e. sol. water. - KA': monoclinic tables.—NaA'. $[a]_D = -17.2^{\circ}$.—CaA'₂ (dried at 100°). Formed by boiling the lactone with water and CaCO₂. Amorphous. $[a]_D = -5.7^{\circ}$. — CuA'₂ 4aq: blue nodules (Kiliani, B. 15, 2955).

Anhydride $C_0H_{10}O_3$. [161°]. [a]_p = 94°. S. 13 at 15°. Electrical conductivity: Walden, B. 24, 2028. H.C. 656,900. H.F. 252,100 (Stoh-

mann, J. pr. [2] 45, 313).

Preparation. - A cold solution of 1 kg. of invert sugar in 9 litres of water is treated with 100 g. of powdered lime and allowed to stand, agitating at intervals. After 14 days 400 g. more CaO2H2 is added, and the mixture again allowed to stand for one or two months, until it no longer reduces Fehling's solution. It is then saturated with CO2, the remaining Ca precipitated with oxalic acid, and the filtrate evaporated nearly to a syrup. The saccharin which separates is recrystallised from hot water; the yield is 100 g. (Kiliani, B. 15, 2954).

Properties.-Prisms, with bitter taste, v. sol. hot water. Dextrorotatory. Converted by alkalis into lævorotatory salts of saccharinic acid. May be volatilised almost without decomposition. Can be extracted by ether, even from solutions containing Na₂CO₃. Does not ferment with yeast. Does not reduce Fehling's solution. even after long boiling with dilute H,SO4. Not attacked by dilute HNO3. Conc. HNO3 forms oxalic acid and saccharonic acid $C_0H_{10}O_7$. 1 g. reduces 4.6 g. of KMnO₄. Oxidised by moist Ag₂O to formic, acetic, and glycollic acids. Boiling HIAq reduces it to the lactone of γ-oxyisohexoic acid CH3.CH(OH).CH2.CHMe.CO2H, which is further reduced by HI at 200° to CHMePr.CO.H (Liebermann a. Scheibler, B. 16, 1821; Kiliani, A. 218, 371). I and KOH give iodoform. Potash-fusion forms formic and lactic acids (Hermann a. Tollens, B. 18, 1333). Boiling HClAq has no action. Phenyl-hydrazine forms C₂H₁₁O₃(N₂H₂Ph) [165°], crystallising from alcohol in needles, v. sol. water (Fischer a. Passmore, B. 22, 2733). Phenyl cyanate at 165° forms Ca4HaeN4Os, crystallising in silky needles.

[230°-240°], v. sol. hot aniline, sl. sol. alcohol | (Tesmer, B. 18, 2607)

Isosaccharinic acid CaH12Oa i.e.

 $CH_2(OH).CH(OH).CH_2.C(OH)(CO_2H).CH_2OH$? Formed by the action of lime on malt-extract at 20° or on milk-sugar (Cuisinier, Bl. [2] 38, 512). The free acid at once splits up into water and lactone. The Na salt is leevorotatory.—CaA'2. S. 1 in hot water. Crystalline powder.

Lactone C.H.O. Isosaccharin. $[a]_D = 63^\circ$ in a 10 p.c. aqueous solution; = 74° in HOAc. Electrical conductivity: Walden, B. 24, 2028. Prepared by allowing a solution of milk-sugar (1 kilo) in water (9 litres) mixed with slaked lime (450 g.) to stand for 6 weeks; filtering; saturating the filtrate with CO2; boiling, again filtering, and evaporating to 500 c.c. The calcium isosaccharinate (170 g.) which then crystallises is decomposed by oxalic acid (Kiliani, B. 18, 631). Large crystals, v. e. sol. water, alcohol, and ether. Dextrorotatory. Does not ferment with yeast or reduce Fehling's solution.

Reactions.-1. Reduced by HI and P to the lactone of oxy-hexoic acid, and finally to CHPrMe.CO₂H. -2. Oxidised by HNO₃ to dioxy-propane tricarboxylic acid C, H, (OH), (CO, H), which is split up by heat into CO, and ay-di-oxyglutaric acid CH₂(CH(OH).CO₂H)₂ (Kiliani, B. 18, 2514).-3. Phenyl cyanate at 165° forms $C_{s_1}H_{s_0}N_4\acute{O}_s$ [181°] a white amorphous powder (Tesmer, B. 18, 2609).

Anilide C₁₂H₁₇NO₅. [165°]. Formed by heating the lactone with aniline at 115° (Sorokin, Bl. [2] 47, 168; J. pr. [2] 37, 318). Needles, v. sol. water. Decomposed by acids and alkalis into aniline and isosaccharin.

Metasaccharinic acid C₆H₁₂O₆ i.e. CH₂(OH).CH(OH).CH(OH).CH(OH).CH₂.CO₂H. Formed, together with isosaccharinic acid, by the action of lime on a cold solution of milk-sugar, its Ca salt being contained in the mother-liquor from which calcium isosaccharinate has crystallised (Kiliani, B. 16, 2625; 18, 642). The free acid splits up at once into water and lactone. CaA', 2aq: crusts of minute prisms, v. sl. sol. cold water.—CuA'2 2aq: groups of minute green plates.

Lactone C.H.O. Metasaccharin. [142°]. $[a]_{\rm p} = -48^{\circ}$. Large trimetric plates; a:b:c = 624:1:899. V. sol. cold water and alcohol, v.

sl. sol. ether. Lævorotatory.

Reactions.—1. Reduced by HI to the lactone of oxy-n-hexoic acid.—2. Oxidised by HNO, to tri-oxy-adipic acid.—8. Phenyl cyanate at 165° forms C₈₄H₈₀N₄O₉, a white amorphous powder [205°-210] (Tesmer, B. 18, 2608).

SACCHAROLACTONE v. Anhydride of Sac-CHARIC ACID.

SACCHARONIC ACID C.H 10O, i.e.

CO,H.CH(OH).CH(OH).CMe(OH).CO,H. Formed by heating saccharin (1 pt.) with HNO. (3 pts. of S.G. 1.375) at 35°, diluting, adding CaCO, filtering, evaporating, and extracting with ether (Kiliani, A. 218, 361). The product so obtained is the lactonic acid; the free acid at once splits up into water and lactone.- Na,A". Formed by boiling the anhydride with Na₂CO₃Aq. Crystalline. — (NH₄)₂A".—CaA": gummy. Ag.A": flocculent pp.

Lactonic acid C.H.O. Saccharone. [a] = -6°. Formed as above. Large trimetric

crystals (containing aq), v. sl. sol. ether. Conc. HIAq converts it into a dibasic acid C.H.O. [139°] crystallising from water in small prisms, and finally to a-methyl-glutaric acid. Does not reduce Fehling's solution.—NaA' aq. Formed from saccharone (2 mols.) and Na₂CO₃Aq (1 mol.). Trimetric prisms with neutral reaction.—NH,A': crystals. Its aqueous solution is ppd. by lead subacetate, but not by lead acetate or AgNO,.

SACCHARUMIC ACID C11H18O11. A product of the action of baryta on dextrose (Reichardt, Vierteljahrschrift pr. Pharm. 19, 384, 503). Yellowish-brown powder with bitter taste, v. sol. water and alcohol, sl. sol. ether. Its solution becomes dark on exposure to air or on addition of alkalis. — BaH₂A'. — Ba₂A'. 5aq: pp. — Cu₂A'. 3aq: greyish-brown pp. — Pb₂A'. — Pb₂C₁₄H₁₂O₁₁. — SACCULMIC ACID C₁₁H₁₆O₄? Formed,

together with sacculmin, by boiling cane-sugar (300 g.) with H_2SO_4 (15g.) and water (420 g.) (Stein, A. 30, 84; Sestini, G. 10, 121). The yield of the two bodies is about 10 p.c. of the sugar used. Glittering black mass, sl. sol. water and alcohol, insol. ether. The alcoholic solution is red, with reddish-brown reflex. Sol. KOHAq and reppd. by acids. AgNO, gives brown amorphous C₄₄H_{se}AgO₁₈. BaCl₂ ppts. brown C₂₂H₁₈BaO₈ aq. Chlorine passed into water in which sacculmic acid or sacculmin is suspended gives yellow flocculent di-chloro-oxysacculmide C11H8Cl2O8. Di-chloro-oxy-sacculmide boiled with KOHAq forms oxysacculmic acid $C_{11}H_0O_0$, which is sol. water, but insol. water containing H_2SO_4 . CuSO, gives in its aqueous solution a brown pp. of $C_{44}H_{30}CuO_{21}$. Bromine and water form amorphous orange $C_{44}H_{26}Br_0O_{22}$ (Sestini, G. 12, 292).

Sacculmin C44H88O15. Formed as above. Black amorphous mass, insol. KOHAq. Reacts with chlorine water in the same way as sacculmic acid. According to Conrad and Guthzeit (B. 18, 443; 19, 2844), sacculmin and sacculmic acid are mixtures of variable composition.

SAFFLOWER v. CARTHAMIN.

SAFFRANINE (PHENO-) C18H18N4O. Phenosaffranine. The hydrochloride may be represented as $C_{e}H_{s}(NH_{z}) < \frac{N}{NPhCl} > C_{e}H_{s}NH_{z}$ (Bernthsen, B. 20, 179, 2690; cf. Andresen, B. 19, 2215; Nietzki, B. 19, 3017, 3163; Witt, B. 19, 3121), or by $C_sH_4 < \frac{N}{NCl(C_sH_*NH_*)} > C_sH_s(NH_*) \begin{bmatrix} 2\\1 \end{bmatrix}$ (Witt).

Formation.—1. Formed by gently warming phenylene-p-diamine (1 mol.) with aniline (2 mols.) and K₂Cr₂O₇ (Nietzki, B. 16, 466). A blue compound C_eH₄<\(\frac{NCl(NHPh)}{NCl(NHPh)}\)>C_eH₂NH₂(?) is first formed, and changes at 100° into the red saffranine (Barbier a. Vignon, Bl. [2] 48, 338 772; C. R. 105, 939). The blue compound is reduced by zinc and hydrochloric acid to C₆H₄<NH>C₆H₃.NH₂, which is oxidised by air to amido-phenazine. According to Nietzki, however, the blue intermediate compound is the indamine C.H. (NH2).N:C.H4:NH2Cl and yields di-amido-diphenylamine on reduction.—2. From benzene-azo-aniline (amido-azo-benzene) by treatment with nitro-benzene, Fe, and HClAq

(B. a. V.). -8. By oxidising a mixture of NH(C₂H₄.NH₂[1:4])₂ (1 mol.) and aniline (1 mol.)

with K2Cr2O, (B.).

Properties.—Golden plates. When freshly precipitated by baryta from the sulphate it is C18H18N4OH, but after frequent recrystallisation from hot water it approximates to C18H14N4 (Nietzki a. Otto, B. 21, 1590). V. sol. hot water, sol. alcohol, almost insol. ether. Conc. H2SO, gives a green colour, changing to blue and red on dilution. Conc. HClAq gives a blue colour. NaNO₂ added to a solution of saffranine hydrochloride forms the diazo- compound C18H15N5Cl2, which gives a blue solution and yields $C_{1s}^{\dagger}H_{1s}^{\dagger}N_{s}Cl_{2}^{2}$ AuCl₂. This diazo- compound on boiling with alcohol forms C18H13N3, the salts of which form red aqueous solutions and dye wool and silk a bluer shade than saffranine. The base $C_{18}H_{18}N_3$ is also got by adding H_2SO_4 till the colour is violet, then adding excess of NaNO2 and boiling: it yields B'HNO₂, B'H₂SO₄, and a violet acetyl derivative. Alcoholic potash forms NH, and saffranol. Zinc-dust and HCl give leuco-saffranine and, on long boiling, C18H18N8O [130°], crystallising from water in long needles yielding C₁₈H₁₈AcN₃O [173°] and converted by nitrous ether into C₁₈H₁₈N₂O [117°], crystallising from alcohol in needles.

Salts. — $(C_{18}H_{15}N_4Cl)_2PtCl_4$. — $C_{18}H_{15}N_4Cl$. Flat needles with green lustre, sl. sol. cold, v. sol. hot water, insol. HClAq.—B'HNO_s. Almost

insol. HNO,.

Di-acetyl derivative C18H14Ac2N4. B'HCl. Got by heating saffranine hydrochloride with Ac₂O and NaOAc (Nietzki, B. 16, 468). Lustrous brown plates, forming a violet solution

in alcoholic soda.—B'HI.

Di-methyl-saffranine. The chloride of this substance C20H10N4Cl is formed, together with N₂(C₆H₄NMe₂)₂, by the action of p-nitroso-dimethyl-aniline (1 mol.) on aniline (1 mol.) in alcoholic solution at 100° (Barbier a. Vignon, C. R. 105, 672). An isomeric (?) di-methylsaffranine got by oxidising a mixture of C₆H₁(NH₂)NMe₂ and aniline yields B'₂H₂PtCl₆ and B'HNO₃ (Nietzki, B. 16, 869; 19, 3163). An isomeride is formed by heating benzene-azo-xylidine with aniline at 150° (Menton, A. 263, 837). Crystalline, forming a rose-red aqueous solution. A tetramethyl-saffranine C2H2N4 is got by oxidising di-methyl-phenylene-green mixed with aniline acetate (Bindschedler, B. 16, 867). It yields B'HCl, v. e. sol. water and B'HNO, aq.

Ethyl-saffranine C18H13EtN4. Formed by boiling p-phenylene-ethyl-diamine with aniline, K.Cr.O., and dilute HOAc (Schweizer, B. 19, 150). Formed also by oxidising phenylene-pdiamine, aniline, and ethyl-aniline with K₂Cr₂O, and dilute HOAc (S.). The hydrochloride forms bluish-green hygroscopic crystals, sol. water and alcohol, insol. ether. Its solutions exhibit olivegreen fluorescence.-B'2H2PtCle: dark lustrous

needles.

Di-ethyl-saffranine C₂₂H₂₄N₄. Formed by oxidising a mixture of C₈H₄(NH₂)(NEt₂) (1 mol.) and aniline (2 mols.) with K₂Cr₂O₇ (Nietzki, B. 16, 470). Formed also by oxidising a mixture of phenylene-p-diamine, aniline, and di-ethyl-aniline with K₂Cr₂O₇ (N.). The alcoholic solution is fluorescent. The hydrochloride is sol. HClAq.

NaNO2 added to its solution in HClAq gives a greenish-blue colour due to the diazo-chloride which forms C22H23N3Cl2PtCl4. - B'2H2PtCl6: green needles.

Acetyl derivative C18H11Et2AcN4. hydrochloride is ppd. as glittering brown needles on adding NaCl to its aqueous solution.

 $-B'_{2}H_{2}PtCl_{6}$.

Tetra-ethyl-saffranine C28H32N4 or

C26H28N4.OH. Formed by oxidising a mixture of

phenylene-di-ethyl-p-diamine, aniline, and di-ethyl-aniline with $K_2Cr_2O_7(N.).-B'_2H_2PtCl_b$.

Para-saffranine $C_{20}H_{18}N_4$. Prepared by dissolving mauveine in dilute acetic acid and boiling with PbO. NaOH is added to slight excess, and the red filtrate boiled with some powdered zine and CaCO, and then NaCl added. Commercial saffranine contains some parasaffranine (Perkin, C. J. 35, 728).

Properties .- Red-brown pp., dissolving in water or alcohol. It is isomeric or identical with commercial saffranine, prepared from equal molecules of tolylene p-diamine, o-toluidine, and aniline. Salts.—B'HCl.—B'HNOs. These salts dye silk a shade more scarlet than saffranine.

Saffranine C21H20N4 or C21H21N4OH. Formed from o-toluidine by treatment with nitrous acid and oxidation of the product with K2Cr2O, (Mène, C. N. 25, 215; Hofmann a. Geyger, B. 5, 526; Dale a. Schorlemmer, C. J. 35, 683). Obtained by heating toluene-azo-o-toluidine with o-toluidine hydrochloride at 150°-160° (Witt, B. 10, 873), and by adding K2CrO4 to a hot solution of tolylene-p-diamine hydrochloride (1 mol.) and o- or p-toluidine hydrochloride (2 mols.) (Witt, B. 12, 939; Bindschedler, B. 13, 207). The commercial saffranine (saffranine hydrochloride) can be purified by recrystallisation from water containing HCl, or by treatment with alcohol (Böttger, N. R. P. 23, 181). Reddish-brown crystals, v. sol. water and alcohol, insol. ether. Conc. HCl changes the colour of its solution through violet to blue. H2SO4 turns it blue and finally green. Decolourised by zinc and HClAq. Saffranine is also decolourised by glucose and NaOHAq on heating, and hence may be used as a test for sugar in urine (Crismer, C. C. 1888, 1510); the colour is restored by atmospheric oxidation. Commercial saffranine (.5 g.) is fatal when administered to dogs by subcutaneous injection (Weyl, B. 21, 2191).—B'HCl. Thin reddish crystals, sol. water and alcohol, insol. ether and NaClAq.—B'₂H₂PtCl₆. Yellowish-red crystalline powder, almost insol. water, alcohol, and ether. -×B'HBr: minute needles.—B'HNO_s. Slender reddish-brown needles, v. sl. sol. cold water .-Sulphate: needles, m. sol. water. $B'C_{\epsilon}H_{\epsilon}N_{s}O_{\tau}$: brownish-red insoluble needles.

Substances of the saffranine class are formed by heating $NH(C_sH_4NH_2)_2$ with (4,3,1)- and (4,2,1)- xylidine and with (5,4,2,1)-cumidine, but not with mesidine, (5,4,3,1)-cumidine, or

(5,2,1)-, (6,2,1)-, (5,3,1)-, and (3,2,1)-xylidine (Nietzki, B. 19, 3163).

Saffranol $O_{18}H_{10}N_2(OH)_2$. Formed by boiling saffranine hydrochloride with alcoholic potash for 3 days (Nietzki a. Otto, B. 21, 1593). Ppd. as yellowish needles by adding HCl to its solution in alkalis. Nearly insol. water, alcohol, and HOAc. Conc. H₂SO₄ forms a brown solution. Yields a crystalline diacetyl derivative.

SALICIN. 423

SAFFRON. The dried stigmas of *Crocus* sativus, used as a colouring matter. It contains crocin (q.v.), crocose (q.v.), picrocrocin (q.v.) polychroite, and an oil $C_{10}H_{14}O$ (209°) miscible with alcohol and ether but decomposed by water (Quadrat, J. pr. 56, 68; Weiss, J. pr. 101, 65).

Polychroite C₁₈H₂₀O₁₈. Dried saffron is washed with ether and extracted with water. The extract is mixed with alcohol and filtered, and the filtrate ppd. by ether. Orange deliquescent mass, v. sol. water, insol. alcohol. Yields glucose on boiling with dilute acids.

SAFROLE $C_{10}H_{10}O_2$ i.e.

CH₂ < O.C.CH:CH CH.CH:CH₂ . Shikimole. [8°]. (232°). S.G. 18 1 0956 (S.); 11.3 1.0963 (Gladstone, C. J. 59, 290). V.D. 81 (obs. and calc.). $\mu_{\rm D} = 1.539$ at 11° (G.). $R_{\infty} = 45.57$ (Brühl, B. 21, 477). H.C.v. 1,243,800. H.C.p. 1,244,700. H.F. 40,300 (Stohmann a. Langbein, J. pr. [2] 46, 533). The chief constituent of the essential oil of sassafras (Sassafras officinalis) (Grimaux a. Ruotte, A. 152, 88; J. Schiff, B. 17, 1935; Poleck, B. 17, 1940; 19, 1094; 22, 2861). Occurs also in the oil obtained from the leaves and fruit of Illicium religiosum (Eykman, R. T. C. 4, 36, 45; B. 22, 2757; 23, 864), and in the oil of the camphor tree, Cinnamomum camphora (Flückiger, Ph. [3] 17, 989). Monoclinic crystals, sol. alcohol and ether. Smells like oil of sassafras. Inactive to light. Oxidised by dilute KMnO₄ to CO₂, formio, oxalic, and piperonylic acids, and piperonal. Alcoholic potash converts it into isosafrole. Does not react with hydroxylamine or BzCl. Does not combine with NaHSO_s. Bromine forms C₁₀H₅Br₅O₂ [170°] S. (chloroform) 7.

Isosafrole C₁₀H₁₀O₂ i.e.

CH₂ \bigcirc O₀C₆H₃.CH:CH.CH₃. (247°). V.D. 80·3 (obs.); 81 (calc.). H.C.v. 1,233,600. H.C.p. 1,234,500. H.F. 50,500. Formed by boiling safrole (100 g.) with KOH (250 g.) in alcohol (500 c.c. of 94 p.c.) for 24 hours (Schiff; Ciamician a. Silber, B. 23, 1160). Liquid, even at -18° , miscible with alcohol, ether, HOAc, and benzene, insol. water and alkalis. Conc. H₂SO₄ forms an intensely red solution. Oxidised by K₂Cr₂O₇ and H₂SO₄ to piperonal and acetic aldehyde. Alkaline KMnO₄ gives piperonylic acid. Bromine in CS₂ forms C₁₀Br₂H₂O₂ [110°], crystallising from ligroïn in colourless needles. Sodium reduces it in alcohol solution to C₁₀H₁₂O₂ or C₈H₃Pr \bigcirc OCH₂ (228°), a liquid miscible with

alcohol and ether. Isosafrole is converted by treatment with KOH and MeOH into a compound, apparently different from methyleugenol, but yielding isovanillic acid on oxidation, and [4:2:1] $C_aH_3(C_3H_3)(OAO)_2$ on treatment with Ac_2O (Ciamician a. Silber, B. 25, 1470).

Nitrosite C₁₀H₂N₂O₄ i.e. CH₂O₂:O₆H₂.C CMe ? (Angeli, B. 24, 3994;

25, 1956). Mol. w. 224 (by Racult's method). [124°]. Ac₂O forms a white modification [124°]. Alcoholic potash forms an isomeride [185°] which yields a mono-acetyl derivative [129°], and a benzoyl derivative [146°]. KMnO₄ oxidises

yields a di-acetyl derivative [138°].

SAGAPENUM. A gum resin imported from Egypt and Persia, consisting of yellow agglomerated granules, smelling like garlic and having an acrid bitter taste. Softens in the hand. Yields umbelliferone on distillation. HNO₂ forms styphnic acid. Potash-fusion forms resorcin (Brandes, N. Tr. 2, 2, 97; Pelletier, Bull. Pharm. 3, 481; Johnston, Tr. 1840, 361). After moistening with alcohol it dissolves in H₂SO₄, forming a brown liquid. Sagapenum is only partially soluble in alcohol and ether, but the ethereal extract is not clouded by addition of alcohol. FeCl, colours its solutions black (Hirschsohn, J. 1875, 859; Ar. Ph. 1810, 481; 11, 54, 152, 247, 312, 434; C. C. 1877, 182).

SAGE OIL. Obtained in the south of Europe by distilling sage (Salvia officinalis) with water, the plant being cut down in the autumn (M. M. P. Muir, C. J. 33, 292; 37, 678; cf. Rochleder, A. 44, 4; Herberger, R. P. 34, 138; Hlasiwetz, J. pr. 51, 355). Contains a terpene (156°), salviol (q. v.), $C_{10}H_{18}O$ (c. 200°), a camphor C10H18O, probably a little cymene, and, especially when prepared from English plants, cedrene, C₁₅H₂₄ (c. 260°). The amount of salviol and of camphor is very small at first and increases with the age of the oil, being formed by oxidation of the terpenes. Pure salviol, pure sage terpene, and pure cedrene do not resinify when singly exposed to heat and light; but a mixture of these rapidly darkens.

SAGO. Balls of starch got from the pith of certain species of *Sagus* and *Cycas* by stirring with water, allowing to settle, and rubbing the deposit through sieves.

SALAMANDRINE C₃₁H₄₀N₂O₅. An alkaloid obtained from the poisonous secretion of the cutaneous glands of the salamander (Salamandra maculata). Crystalline, v. sol. water and alcohol (Zalesky, Bl. [2] 6, 344). Alkaline in reaction. Poisonous.—B'H₂Cl₂: needles.

SALIGIN C, H₁₀O, i.s.

C₄H₁₁O₅.O.C₄H₄.CH₂OH. Mol. w. 286. [201°]
(Schiff, B. 14, 304). S. 3·3 at 11° (Piria, A. 96, 378). [a]_B = -62·6° (Tiemann, B. 18, 1600; cf. Hesse, A. 176, 116; Sorokin, J. pr. [2] 37, 331. Occurs in the bark of several species of willow and poplar, e.g. Salix helix, S. amygdalina, Populus tremula, P. græca (Leroux, A. Ch. [2] 43, 440; Braconnot, A. Ch. [2] 44, 296; Pelouze a. Gay-Lussac, A. Ch. [2] 44, 220; 48, 111; Piria, A. Ch. [2] 65, 281; [3] 14, 257; Bouchardat, C. R. 18, 299; 19, 602, 1179; 20, 610, 1635; Gerhardt, A. Ch. [3] 7, 215; Tischhauser, A. 7, 280). Occurs also in castoreum (Wöhler, A. 67, 360).

Formation.—1. By digesting an aqueous solution of helicin with sodium-amalgam (Lisenko, Z. 1864, 577) or with sine and H₂SO₄ (Michael,

Am. 5, 172).—2. By boiling populin with lime-

Preparation.—Willow bark (6 lbs.) is boiled with water; the filtrate evaporated (to 18 lbs.); mixed while hot with PbO (2 lbs.); digested for 24 hours and filtered; the filtrate evaporated to a syrup and left to crystallise (Duflos, A. 8, 200; cf. Peschier, A. Ch. [2] 44, 418; Erdmann, B. J. **8**3, 1, 136).

Properties .- Trimetric tables;

a:b:c = 927:1:2.494.Sol. water and alcohol, Tastes bitter. insol. ether. Lævorotatory. Neutral to litmus. Its solutions are not ppd. by lead acetate or subacetate, by gelatin, or by tannin. Conc. H₂SO₄ gives a red colour. IClAq forms a crystalline body (Stenhouse, C. J. 17, 327). After injection of salicin, salicylic aldehyde and salicylic acid are found in the urine (Laveran a. Millon, A. Ch. [3] 13, 145; Ranke, J. pr. 56, 1).

Reactions.—1. Decomposed at 240° into saliretin and glucosan.—2. Split up by emulsin and saliva at 40° into glucose and saligenin (Piria; Städeler, J. pr. 72, 350).—3. Boiling dilute H,SO, forms glucose and saliretin.—4. Oxidised by chromic acid mixture to CO2, formic acid, and salicylic aldehyde.-5. Cold dilute HNO, forms helicin. Hot conc. HNO, forms picric acid .- 6. Potash-fusion forms salicylic acid .-7. Boiling NaOHAq forms saliretin.—8. HCl and KClO, give tetra-chloroquinone.

Metallic derivatives. — C13H1, NaO, Formed from salicin and NaOEt in alcohol (Perkin, C. N. 18, 110). Crystalline. — Pb₂C₁₃H₁₄O₂. Got by adding NH₃Aq to a hot conc. solution of salicin, and then adding lead subacetate. Bulky white pp.

Acetyl derivative C13H14Ac4O,. Needles (from alcohol), nearly insol. water (Schiff, Z. [2]

5, 51).

Benzoyl derivative C20H22O8 i.e. C₁₁H₁,B₂O₂. Populin. [180°]. S. ·05 at 9° (Piria); ·04 at 15°; ·24 at 100° (Schiff); S. (alcohol) 1 at 16°. Occurs in the bark and leaves of the aspen (*Populus tremula*) (Braconnot, A. Ch. [2] 44, 296; Piria, A. Ch. [3] 34, 278; 44, 366; A. 81, 245). Formed by heating salioin with Bz₂O (Schiff, A. 154, 5). Prepared by boiling the leaves of the aspen with water, ppg. with lead subacetate, removing lead from the filtrate by H2S, and evaporating to crystallisation. Light silky needles (containing 2aq). Tastes sweet. Lævorotatory (Biot a. Pasteur, C. R. 34, 607). Sl. sol. water and alcohol, nearly insol. ether. K₂Cr₂O, and H₂SO₄ form salicylic aldehyde. HNO₄ (S.G. 1·3) forms benzoylhelicin. Conc. H₂SO₄ forms a deep-red solution, whence water produce a powder ("rutilin"). whence water ppts. a red powder ('rutilin'). Boiling dilute acids split it up into glucose, benzoic acid, and saliretin. Alcoholic NH, at 100° forms benzamide, benzoic ether, and salicin. Populin is not hydrolysed by emulsin.

C,3H,6Bz2O,. Di-bensoyl derivative Formed, together with the mono- and tetra-benzoyl derivatives, by heating salicin with Bz O (Schiff). Flocculent mass, nearly insol.

water, sl. sol. ether.

Tetra-bensoyl derivative C13H14Bz4O1. Amorphous resin, not coloured by cold H2SO4. Chloro-salicin C₁₂H₁₇ClO₇. Formed by passing Cl into water in which salicin is suspended

(Piria, A. Ch. [8] 14, 275). Long silky needles (containing 2aq), sol. water and alcohol, insol. Tastes bitter. Resolved by emulsin into glucose and chloro-saligenin. Yields a tetraacetyl derivative, crystallising from alcohol in prisms.

Di-chloro-salicin C₁₃H₁₆C₁₂O₂. Formed by the action of Cl on chloro-salicin. Silky needles (containing aq), nearly insol. cold water, m. sol. alcohol. Gives no colour with FeCl, or H,SO. Split up by emulsin into glucose and di-chloro-

saligenin.
Tri-chloro-salicin C₁₃H₁₅Cl₃O₇. Formed by chlorinating the preceding body in presence of water and CaCO₃. Yellowish needles (containing aq) (from dilute alcohol), almost insol. cold

water.

Bromo-salicin C₁₂H₁,BrO₇. [160°]. Formed by slowly adding Br to salicin (1 pt.) in water (20 pts.) (O. Schmidt, Z. [2] 1, 320). Four-sided prisms, v. sol. water and alcohol. Split up by emulsin into glucose and bromosaligenin.

SALICYL. The radicle o-oxy-benzoyl [1:2]C_sH₄(OH).CO. The same name is sometimes applied to the radicle o-carboxyl-phenyl [1:2]CO₂H.C₆H, and, rarely, to o-oxy-benzyl C,H,(OH).CH,,

SALICYLAMIC ACID. An old name for

o-Amido-Benzoic acid.

SALICYLAMIDE. The amide of o-Oxy-BENZOIC ACID.

SALICYLAMINE. A name for o-Oxy-BENZYL.

SALICYL-GLYCIDIC ACID v. OXY-PHENYL-GLYCIDIO ACID.

SALICYL-GLYCOLLIC ACID is o-CARBOXY-PHENOXY-ACETIC ACID

SALICYLIC ACID v. o-Oxy-BENZOIC ACID. Homosalicylic acid v. Oxy-TOLUIC ACID.

SALICYLIC ALDEHYDE v. o-OXY-BENZOIC ALDEHYDE.

SALICYL-LACTIC ACID v. DI-OXY-PHENYL-PROPIONIC ACID.

SALICYL-PHENOL v. DI-OXY-BENZOPHENONE. SALICYL-RESORCIN v. TRI-OXY-BENZOPHEN-

SALICYL-SULPHURIC ACID v. CARBOXY-PHENYL SULPHURIC ACID.

SALIGENIN v. o-OXY-BENZYL ALCOHOL.

SALIGENOL. Another name for o-Oxy-BENZYL ALCOHOL.

SALIRETIN v. o-Oxy-BENZYL ALCOHOL.

SALOL v. Phenyl ether of o-Oxy-Benzoic

SALT-FORMING OXIDES. Oxides which form salts, either by reacting with acids (or acidic oxides)—in which case the oxides are basic-or by reacting with basic oxides or hydroxides—in which case the oxides are acidic. For a table showing the compositions and general characters of the characteristic highest saltforming oxides of each group of elements, v. vol. M. M. P. M. iii. p. 661.

SALTPETRE. A common name for potassium

nitrate; v. vol. iii. p. 514.

SALTS. The earliest use of the word salt seems to have been to designate the solid obtained by evaporating sea-water. The term was afterwards applied to solids which more or less resembled sea-salt, especially to those solid bodies which were easily soluble in water. The

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study of the compositions of salt-like substances led to the view that they were made up of two parts, one of which was electro-positive with regard to the other. When the compositions of acids had been generalised in the statement that acids are compounds of hydrogen with nonmetallic, or negative, elements, the relation of salts to acids was indicated by calling salts me-tallic derivatives of acids. This conception of the composition of salts included the older view, inasmuch as the metallic derivative of an acid is necessarily a compound of a positive metal with a less positive non-metal, or with a (less positive) group of elements, which non-metal, or group of elements, was combined with hydrogen in the parent acid.

The most general conception of the composition of salts is that which formulates them as M_xR_y , where M is the positive, or basic, or basylous part, and R is the negative, or acidic, or chlorous part of the salt. Both M and R may be either simple or complex. Common salt, NaCl, is the type of all salts as regards composition; the positive radicle, Na, may be replaced by other metals, or by groups of elements which are positive with regard to the other part of the compound; so the negative radicle Cl may be replaced by other non-metals, or by groups of elements which are negative with regard to the other part of the compound.

When an aqueous solution of a salt is electrolysed, the positive radicle is separated at the negative electrode, and the negative radicle at the positive electrode (secondary reactions may occur). An element may form part of the positive radicle of a salt, and the same element may form part of the negative radicle of another salt. Thus in the electrolysis of a ferric salt the iron travels to the cathode, but in the electrolysis of K, Fe(CN), the iron travels, with the cyanogen, to the anode. In the electrolysis of PtCl₄ the Pt is separated at the negative electrode, but in the electrolysis of Na, PtCl, the Pt is separated, with the Cl, at the positive electrode. We must then regard the radicles of, say, FeCl, as Fe and 3Cl, the radicles of PtCl, as Pt and 4Cl, and the radicles of NaCl as Na and Cl; but we must look on the radicles of K4Fe(CN)6 as 4K and Fe(CN)6, and the radicles of Na₂PtČl_s, not as 2Na, Pt, and 6Cl, but as 2Na and PtCl_s. Again, neither the salt K, Fe(CN), nor the salt Na2PtCl, in aqueous solution, gives the ordinary reactions of iron or platinum; but if these salts were composed of the radicles 4K, Fe, and 6CN, and 2Na, Pt, and 6Cl respectively, we should expect them to show the same qualitative reactions as solutions of FeCl, and PtCl. In contrast with these salts is the compound formed by evaporating a mixture of K2SO, Aq and CuSO, Aq. The salt thus produced, K2SO4.CuSO4.6H2O, when dissolved in water, gives the ordinary reactions of sulphates, of K compounds, and of Cu compounds; moreover, on electrolysis the K and Cu appear at the cathode, and the group SO, is set free at the anode (where it reacts with H.O to give H.SO. and O).

But all these salts-NaCl, PtCl, FeCl, K,Fe(CN), Na₂PtCl, and CuSO, K₂SO,—are represented, as regards composition, by the general formula M.R. This formula satisfactorily ex-

presses the composition of all salts; but it says nothing about the properties of these compounds. When solutions of two salts are mixed. a more or less complete interchange of positive radicles generally takes place. This capability of reacting as if they were composed of two parts-or, one may say, of reacting in a binary way-is characteristic of salts. Salts in solution are most probably dissociated into two parts, each of which carries an electric charge. Even if the whole of the salt is not dissociated, the portion which readily takes part in chemical changes is most probably dissociated binarily (cf. ELECTRICAL METHODS, p. 184).

Salts, then, may be regarded as compounds of the form MzRv, which readily exchange M and R with other compounds of similar composition, and which in aqueous solutions are wholly or partially separated into their positive and negative radicles, each radicle carrying with it an electric charge.

Salts are often classified as normal, acid, basic, and double salts.

An acid salt is one whose negative radicle contains hydrogen that can be eliminated by causing the salt to react with an alkali or basic oxide, while a normal salt contains no replaceable hydrogen. As an acid is described as a compound containing hydrogen, all or part of which is replaceable by metal when the acid interacts with a metal or a basic oxide, it is evident that an acid salt comes under the description of an acid, and, strictly speaking, is an acid. According to the meaning already given to the term 'salt,' the class of salts includes acids. It is, however, convenient to give a special name to the compounds of hydrogen with negative radicles. A basic salt contains a positive radicle or radicles which can be replaced by a negative radicle with formation of a normal salt. This positive radicle is sometimes the group OH, sometimes it is a group M_xO_y , where M is the metal of the salt. Thus Bi(OH), NO₃ and Pb(PbO)NO₃ are basic salts. When treated with nitric acid the former gives the normal salt Bi(NO₃)₃, and the latter the normal salt Pb(NO₃)₂. The basic salt is an intermediate stage between the base and the normal salt, just as the acid salt is an intermediate stage between the acid and the normal salt.

Basic salts are formed by fairly weak bases; the very strong bases, Na₂O, K₂O, CaO, &c., do not form basic salts.

Normal salts frequently combine with other normal salts to form double salts. Generally speaking, one of the components of a double salt is a salt of a strong base which forms acid salts but not basic salts, and the other component is a salt of a weaker base which readily forms basic salts but does not form acid salts. That component which is itself the salt of a strong base -e.g. K2SO4, NaCl, &c.-may be regarded as the more acidic or negative radicle of the double salt, while the component which is itself the salt of a weaker base—e.g. Al₂(SO₄), Zn₂(PO₄)₂, &c.—may be regarded as the more basic or positive radicle of the double salt. The double salt belongs to the type M.R.; both M and R are themselves salts. And just as many normal salts are formed by the direct union of their radicles-e.g. NaCl by combining Na and 426 SALTS.

Cl, BaSO₄ by combining BaO and SO₅—so double salts are formed by the union of a more positive with a less positive radicle, which is, in such

cases, itself a salt.

The reactions of some double salts indicate that these salts are to be regarded as composed of the two simple salts which were brought together when the double salts were formed; the salt MgSO, K, SO, 6aq, for instance, gives, in solution, the reaction of sulphates, of Mg, and of K, and on electrolysis the Mg and K go to the negative electrode. On the other hand, the reactions of some double salts indicate that these salts are not to be regarded as composed of the simple salts by the union of which the double salts were formed; the salt $Cr_2(C_2O_4)_3.K_2C_2O_4$, for instance, does not give the reactions for instance, does not give the constant of oxalates, nor the reactions of Cr: this salt must be looked on as the potassium salt of the acid $H_2Cr_2(C_2O_4)_4$. If the salt (NH₄)₂Cr₂(C₂O₄)₄.2H₂O is treated with NH₃Aq, a new salt is formed having the composition $Cr_2(C_2O_4)_3.(NH_4)_2C_2O_4.4NH_3.xH_2O$, and the reactions of this salt show that it is an oxalate, and therefore that it has a different constitution from that of the salt (NH₄)₂Cr₂(C₂O₄)₄ from which it is obtained.

The salts of the haloid acids combine to form many double salts, e.g. PbI₂.KI, BiCl₃.3KCl, AuCl₂.KCl, &c. Most of these double haloids are generally regarded as double salts in the usual acceptation of the term; but several of them do not give the ordinary reactions of the less positive metal they contain, nor the reactions of the halogen which enters into their composition. For instance, PtCl₂.NaClAq reacts with two equivalents of AgNO₃ to give PtCl₄.2AgCl; if PtCl₄.2NaCl were a double salt, reacting in solution as if it were composed of the two radicles PtCl₄ and NaCl, it would interact with six equivalents of AgNO₃, and the products would be Pt(NO₃)₄, NaNO₃, and AgCl. It is evident, then, that the Na-Pt chloride is the Na salt of the acid H₂PtCl₆, and that the constitution of this salt is similar to that of the ordinary simple salts.

Remsen (Am. 11, No. 5 [1889]) proposes to regard all the so-called double haloids as simple salts similar to ordinary oxysalts. It seems to me that the constitutions of these salts cannot be determined by an examination of their composition only, but that the reactions of each salt must be examined. Some of the double haloids are undoubtedly simple salts, as some of the double oxysalts are certainly simple salts of metal-containing acids; but as some oxysalts containing two metals react so as to show that they are composed of two radicles, each of which is a simple salt, so some of the double haloids also give the ordinary reactions of both the metals, and also of the acidic radicle, which they contain, and must therefore be regarded as true M. M. P. M. double salts.

SALVIOL $C_{10}H_{18}O$. (200°). S.G. 15 .938, $\mu_D = 1.462$ at 20°. R_{∞} 74.2. (M. M. P. Muir, C. J. 33, 292; 87, 683). A constituent of oil of Sage (q, v). It is a dextrorotatory liquid. On distillation some of it splits up into H_2O and $C_{10}H_{10}$.

Reactions.—1. Chromic acid mixture or dilute nitric acid produces acetic and oxalic

acids and camphor [174°].—2. P_sO_s produces polymerides of $C_{10}H_{1s}$, a terpene (171°), an aromatic hydrocarbon (c. 130°), and a paraffin-like hydrocarbon (170°–180°). Cymene is not formed. Hence salviol is not $C_{10}H_{1s}O$. However, according to Semmler (B. 25, 3350), salviol is identical with tanacetone.

Constitution.—Refractive index appears to indicate absence of C:C groups.

SALYLIC ACID. A name given by Kolbe to a supposed isomeride of benzoic acid got from salicylic acid by successive treatment with PCl₂ and sodium-amalgam. It is identical with benzoic acid (Reichenbach a. Beilstein, A. 132, 309). The name is also applied by Städeler (A. Suppl. 7, 159) to two acids: (a) C₁₄H₁₄O₅ [101°] and (β) C₂₁H₂₂O₈ [95°], obtained by allowing salicylic aldehyde to stand for 12 years under water. The (a)- acid forms small four-sided prisms, gives no colour with FeCl₃, and yields Ag₂A". The (β)- acid forms aggregates of needles, gives an intense violet colour with FeCl₃, and yields Ag₃A". Both acids are v. sol. alcohol and ether.

SAMARIUM. This name was given by Lecoq. de Boisbaudran (C. R. 88, 322; 89, 212) to the metal of an earth which he separated from what was then known as didymia, occurring in samarskite. Delafontaine (C. R. 89, 632) obtained from didymia an earth which he called decipia, and when this had been removed by ppn. as sulphate, the new earth, samaria, remained in solution, and was obtained by fractionally ppg. by NH, Aq. Cleve (C. J. 43, 362) separated what he regarded as pure samaria from orthite; he prepared and described several salts of the new element. Cleve found the at. wt. of Sm to be 150, supposing the oxide to be Sm₂O₃; this number was confirmed by Bettendorff (A. 263, 164). It is still very doubtful whether samaria is a homogeneous substance; according to Crookes, it contains at least four different bodies (v. METALS, RARE, vol. iii. p. 244). Because of the great uncertainty attaching to the homogeneous nature of the so-called samaria it does not seem desirable to give an account of the bodies which have been described as compounds of Sm. The student is referred to the memoirs cited above, and also to the following:-L. de Boisbaudran, C. R. 114, 575; 116, 611 (spectrum and compounds of samaria); Marignac, Ar. Sc. [3] 3, 413 (spectrum, and compounds); Krüss a. Nilson, B. 20, 2134; 21, 2310 (spectrum); v. also, for spectrum, B. A. 1884. 438. The compounds of the supposed element are described by Cleve (l.c., and also Bl. [2] 43, 162) (cf. also Earths. vol. ii. p. 422). M. M. P. M.

SANDAL W00D. A dye wood furnished by Pterocarpus santalinus, which grows in tropical Asia. It dyes mordanted cotton red. It contains santalin (Meier, A. 72, 320). Yellow sandal wood (from Santalum album), yields by steam-distillation an oil containing santalol C_{1s}H_{2s}O, and santalal C_{1s}H_{2s}O (Chapoteaut, Bl. [2] 37, 303). After treatment with Na the oil yields a terpene C_{1o}H_{1e}, S.G. 919, μ_A = 1·487 (Gladstone, C. J. [2] 10, 6). The crude oil heated in sealed tubes at 310° forms C_{2s}H_{2s}O (c. 340°), C_{4s}H_{2s}O₂ (350°) and C_{2s}H_{2s}O (240°) whence P₂O₃ forms C_{1o}H_{1s} (c. 177°).

Santalol (310°) is converted by P₂O₅ into water and C₁₅H₂₄ (260°). HOAc at 150° forms the acetyl derivative C₁₅H₂₅OAc (298°), and a compound C₃₀H₄₀O (283°).

Santalal (300°) is converted by P₂O₅ into

C₁₈H₂₂ (248°).

Sandal wood oil from Santalum Preissianum contains a crystalline solid [105°] (Ph. [3] 22, 328).

SANDARACH. A gum resin which exudes from Thuja articulata growing in Barbary, and from various species of Callitris growing in Australia and New Zealand (Johnston, J. pr. 17, 157; Maiden, Ph. [3] 20, 562). Transparent yellowish mass with vitreous fracture, resembling pine-resin in taste and smell. Sol. alcohol. Completely sol. ether. Its alcoholic solution is ppd. by Pb(OAc)₂. It appears to contain several resins $(e.g.~\dot{C}_{40}H_{62}\dot{O}_{8},~C_{40}\dot{H}_{62}\dot{O}_{6})$ differing in solubility in alcohol.

SANGUINARINE C17H15NO4 (E. Schmidt, Ar. Ph. [3] 26, 622). Prepared from the root of Sanguinaria canadensis (Dana, Annals of the Lyceum of New York, 2, 245; Probst, A. 29, 120; 31, 241; Schiel, A. 43, 233; Am. S. [2] 10, 220; Naschold, J. pr. 106, 385). According to Probst and Schiel it is identical with chelerythrine, but Naschold and Schmidt assign the formula $C_{17}H_{15}NO_4$ to sanguinarine and $C_{18}H_{17}NO_4$ to chelerythrine. Prepared by exhausting the root with ether and ppg. the hydrochloride by HCl. Pearly grains, insol. water, sol. alcohol and ether. Poisonous.— Salts: B'HCl aq.—B'2H2PtCl6 (dried at 100°).- $B'_2H_2PtCy_6$. According to König (C. C. 1891, i. 321) sanguinarine has the formula $C_{20}H_{15}NO_4$, crystallises with jaq, melts at 211°, and forms the red salts B'HCl 5aq, B'HNO, aq, B'HAuCl, and B'2H2PtCl6.

SANTAL Č,H,O,. Obtained from sandal wood by extracting with very dilute KOH, ppg. with HCl, and extracting the pp. with ether (Weidel, Z. [2] 6, 83). It is accompanied by a scarlet crystalline powder C14H12O4? Santal crystallises from alcohol in colourless iridescent four-sided laminæ (containing $1\frac{1}{2}aq$), insol. water, sl. sol. cold alcohol, sol. KOHAq, sl. sol. NH, Aq. FeCl, colours its alcoholic solution dark red. Potash-fusion yields protocatechuic

SANTALIN $C_{15}H_{14}O_{5}$ (Meier, A. 72, 320); $C_{15}H_{16}O_{5}$ (Wegermann a. Häffely, A. 74, 226); $C_{17}H_{16}O_{5}$ (Franchimont, B. 12, 14). [105°]. The colouring matter of sandal wood and of coliatur wood. Extracted by ether or alcohol, and purified by ppg. its alcoholic solution with Pb(OAc), and decomposing the pp. by dilute H₂SO₄. Minute red prisms (M.) or amorphous (F.), without taste or smell, insol. water. Its alcoholic solution is blood-red. Sol. alkalis, forming violet solutions in which BaCl₂ ppts. Ba(C₁₅H₁₃O₂)₂ as a dark-violet amorphous pp. Pb(OAc), added to its alcoholic solution ppts. PbC₁₅H₁₄O₆ (dried at 100°). On heating with HCl, santalin gives off MeCl (1 mol.). Gives resorcin and HOAc on fusion with potash. KMnO, gives off an odour of vanilla.

SANTONIC ACID C15H20O, i.e.

C_eH_eCHMe.CH(OH).CH₂.CO₂H (?) (Cannizzaro, B. 18, 2748). [171°] (Hvoslef, B. 6, 1471).

S. \cdot 56 at 17°. [a]_D = -74° at 20° in chloroform (Nasini, B. 13, 2210; G. 13, 164). Formed by boiling santonin with baryta-water (Cannizzaro a. Sestini, G. 3, 241; Hesse, B. 6, 1280). Colourless trimetric plates (Strüver, G. 6, 349), m. sol. hot water, v. sol. ether, v. e. sol. alcohol. Acid to litmus. Leevorotatory. HIAq forms $C_{15}H_{26}$ (235°-245°) and $C_{15}H_{21}$ (144° at 3 mm.) (Cannizzaro a. Amato, B. 7, 1104). Boiling HIAq and P form (a)- and (B)-metasantonin. By boiling with HOAc and heating the residue it is converted into santonide and parasantonide. Sodium-amalgam yields hydrosantonic acid. Does not react with hydroxylamine. When its chloroform solution is heated with POCl, as long as HCl comes off, and a current of damp air passed through the product, there is formed PO(C₁₅H₁₈O₃Cl)₃ [98°] (Cannizzaro, G. 10, 409) Salts.—NaA'. Stellate groups of needles.

 $-BaA'_2$.—AgA': white pp.

Methyl ether McA'. (86°). Colourless needles (Cannizzaro, G. 6, 335).

Ethyl ether Eth'. [95°]. $[a]_D = -45^\circ$ at 26°. Trimetric crystals.

n-Propyl ether PrA'. (220° at 3 mm.). S.G. $^{20}_{4}$ 1·125. $[a]_{D} = -40^{\circ}$ at 20°.

Isobutyl ether CH,PrA'. [67°]. $[a]_D = -41^\circ$. Needles (Carnelutti a. Nasini, B. 13, 2208; G. 10, 530).

Allyl ether C_3H_3A' . [55°]. $[a]_p = -40^\circ$. Acetyl derivative $C_{15}H_{16}AcO_4$. [140°]. Got by boiling the acid for six hours with AcCl (Sestini, G. 5, 121). Prisms (from CHCl_a). It is accompanied by $C_{1a}H_{2a}O$, [128°].

Chloride $C_{15}H_{19}O_3CI$. [155°]. Bromide $C_{15}H_{19}O_3Br$. [145°]. Iodide $C_{15}H_{19}O_3I$. [136°]. Hydrosantonic acid $C_{15}H_{22}C$

C15H22O1 f170°1 Formed by reducing santonic acid with sodiumamalgam (Cannizzaro, G. 6, 341). Colourless trimetric crystals (from ether), less sol. alcohol than santonic acid. Dextrorotatory. Oxidised by Ag₂O to metasantonic acid. HOAc at 150 forms hydrosantonide C₁₅H₂₀O₃ [156°], which is reconverted into hydrosantonic acid by alcoholic potash (Cannizzaro a. Valente, G. 8, 309). AcCl converts hydrosantonic acid into acetyl-hydrosantonide C₁₅H₁₈AcO₃ [204°], which is converted by alcoholic NH₃ at 180° into hydrosantonamide C₁₈H₂₁O₃.NH₂ [190°]. BzCl fo hydrosantonide C₁₈H₁₉BzO₃ [157°]. BzCl forms benzoyl-

Salts.-NaA' Saq: trimetric crystals. -KA' 2aq: monoclinic crystals.

Santoninic acid C15H20O4 i.e.

CH:CH.CH.CHMe.CO CH:CH.CH.CHMe.CH.CH(OH).CH2.CO2H (Cannizzaro, B. 18, 2746). Santonous acid. $a_1 = -26^{\circ}$ in alcohol at 22°. Formed by heating santonin with KOHAq (Hesse, B. 6, 1280). Trimetric tables, sl. sol. cold water, m. sol. ether, v. e. sol. alcohol. Not turned yellow by light. Acid to litmus. Not coloured red by alcoholic potash. At 120° it splits up into water and santonin; dilute H2SO4 acts in like manner.

Salts.—NaA' 3 aq. S. 33 in cold water. S. (90 p.c. alcohol) 25 (Lepage, J. 1876, 618). Stellate groups of needles. Lævorotatory; $\alpha_{j} = -18^{\circ} \text{ at } 22^{\circ}.$ —BaA', aq.

Metasantonic acid C₁₅H₂₀O₄. [161°]. Formed by heating silver hydrosantonate with water. Got also by distilling santonic acid at 45 mm., by boiling santonide with KOHAq, and by the action of Na₂CO,Aq on isosantonin. Trimetric crystals (from ether). Lævorotatory. Its K and Na salts are very soluble.—AgA': pp., m. sol. water.

Methyl ether MeA'. [102°]. Chloride C₁₅H₁₉O₂Cl. [139°].

Parasantonic acid $C_{15}H_{20}O_4$. [173°], $[\alpha] = -99^\circ$ in chloroform (Carnelutti a. Nasini G. 10, 534). Formed by boiling parasantonide with NaOHAq or HClAq. Trimetric crystals, m. sol. water and ether. Converted by AcCl, Ac₂O, or PCl₅ into parasantonide. Boiling HIAq and P form (α) - and (β) - metasantonin. The parasantonates are mostly v. sol. water.—BaA'₂: slender needles (from water).

Methyl ether Me₂A'. [184°]. [a]_D = -109° in chloroform at 27°.

Ethyl ether EtA'. [178°]. $[a]_D = -100^\circ$ in chloroform at 20°.

n-Propyl ether PrA'. [113°]. $[a]_D = -91^\circ$ at 26°. Colourless prisms.

Allyl ether C_aH_aA' . [149°]. [a]_D = -92°. Lævorotatory.

Photosantonic acid C18H22Os i.e.

CH:CH.CH.CHMe.CO₂H (?). [4]₀= -125° . Formed, as ethyl ether of the lactonic acid, by exposing a solution of santonin in 65 p.c. alcohol to direct sunlight (Sestini, Rep. ital. chim. pharm. 1865). Formed also by exposing an aqueous solution of sodium santonate to sunshine. Prepared by insolution of a 1 p.c. solution of santonin in diluted HOAc for one month (Cannizzaro a. Villavecchia, B. 18, 2750, 2859). Trimetric prisms (from alcohol). Gives off H_2O at 100° , forming the lactonic acid $C_{15}H_{20}O_4$, and then melts at 155° . V. sl. sol. water, v. sol. alcohol and ether. Lævorotatory.

Salts.— (NH₄)₂A" 6aq. — BaA" aq: white amorphous pp.—CaH₂A"₂ 3aq: silky needles, sl. sol. cold water.—CaA" xaq.—Ag₂A": white pp.

Methyl ether of the lactonic acid. [57°]. Got by the action of sunshine on a solution of santonin in MeOH. Long prisms (from MeOH).

Ethyl ether of the (a)-lactonic acid $C_{17}H_{24}O_4$. [69°]. (305°). [a]_D = -118° in a 6 p.c. alcoholic solution at 14°. Got by exposing an alcoholic solution of santonin to sunlight (Sestini, J. 1876, 622). Formed also from $Ag_2C_{15}H_{20}O_5$ and EtI (V.). Formed also by the action of alcohol and H_2SO_4 on photosantonic acid. Thin tables, almost insol. cold water, v. sol. alcohol and ether. Lævorotatory. NaOH converts it into sodium photosantonate.

Ethyl ether of the (8)-lactonic acid C_{1,H_2,O_*} [155°]. [a]_D=77° in alcohol at 14°. Formed, together with the preceding, by insolation of a solution of santonin in absolute alcohol. Tables (from ether). Dextrorotatory.

Anhydrophotosantonic acid $C_{1a}H_{2e}O_4$ i.e. $C_{1s}H_{1e}(CO_2H)_2$. [133°]. [a]_p = +31.9. Obtained by saponification of its ether, which is formed by passing gaseous HCl into a solution of photosantonic acid in absolute alcohol (Villavecchia, B. 18, 2862). Crystalline solid. V. sol. alcohol and ether.—BaA": white amorphous solid, v. e. sol. water.

Di-ethyl ether Et₂A": (180°-186°). [a]_D = +20.4°. Colourless liquid.

Isophotosantonic acid C13 H22O5 i.e.

CH:CH.CH.CHMe.C(OH)₂
CH:CH.CH.CHMe.CH.CH(OH).CH₂·CO₂H^(?)
CH:CH.CH.CHMe.CH.CH(OH).CH₂·CO₂H^(?)
[a]_D = 124° in a 3 p.c. alcoholic solution at 11°.
Formed, together with photosantonic acid, by exposing a solution of santonin (1 kilo.) dissolved in HOAc (52 litres of S.G. 1·054) for several months to sunshine (Cannizzaro a. Fabris, B. 19, 2260). Thick trimetric crystals (from alcohol), sl. sol. water, m. sol. ether, v. sol. alcohol. Dextroordatory. At 100° it gives an anhydride C₁₃H₂₀O₄. Sol. alkalis and hot Na₂CO₂Aq, forming orange-red solutions.—BaA'₂ aq: white amorphous powder, v. e. sol. alcohol and water.

Acetyl derivative C₁₈H₂₁AcO₅. [183°]. [a]_D=59°. Transparent needles. Dextrorotatory.

Di-acetyl derivative *C₁₃H₂₀Ac₂O₅. [c. 166°]. Crystals, v. sl. sol. alcohol and ether. Converted into C₁₅H₂₁AcO₅ by heating with absolute alcohol at 120°.

Pyrophotosantonic acid $C_{14}H_{20}O_{2*}$ [94·5°]. Formed by heating photosantonic acid in a current of CO_2 or H_2 (Sestini a. Danesi, G. 12, 83). Got also by heating photosantonic acid with HI (Cannizzaro a. Fabris, B. 19, 2260). Crystals, sol. alcohol and ether.—Bah'_{2*} Forms, on distillation, $C_{18}H_{20}$ (222°).

SANTONIDE $C_{18}H_{18}O_3$. $[127\cdot5^\circ]$. $[\alpha]_{11}=754^\circ$ in chloroform at 20° ; 666° in alcohol at 20° . S.V.S. 206. Formed by evaporating a solution of santonic acid in HOAc and heating the residue at 180° (Cannizzaro a. Valente; Carnelutti a. Nasini, G. 10, 528; 13, 149). Trimetric crystals. Dextrorotatory. M. sol. chloroform, v. sl. sol. alcohol. Not attacked by PCl₈. Boiling KOHAq forms metasantonic acid.

Parasantonide $C_{15}H_{18}O_{2}$. [110°]. S.G. 20 1·202. [a]_D = 897° at 26°. S.V.S. 206. Formed by boiling santonic acid with HOAc, evaporating, and heating the residue at 260° (Cannizzaro, G. 8, 309; 10, 528). Trimetric crystals, v. sl. sol. alcohol, m. sol. Ac₂O. Not attacked by Ac₂O or PCl₃. Boiling KOHAq converts it into parasantonic acid.

Metasantonide. [138°]. $[a]_D = -223^\circ$ at 26°. S.V.S. 236. Prepared by heating santonin with conc. H_2SO_4 for some hours at 100° and ppg. by water (Cannizzaro a. Valente, Rend. Accad. Lincei, ii.; G. 10, 42). Crystals (from ether). Converted by Na₂CO₂Aq into metasantonic acid.

SANTONIN $C_{16}H_{16}O_{2}$ i.e. CH:CH.CH.CHMe.CO CH:CH.CH.CHMe.CH.CH $\stackrel{CH}{O}^{2}$ CO or

 216; Alms, ibid. 34, 319; 39, 190; H. Trommsdorff, A. 11, 190; Heldt, A. 63, 10, 40).

Preparation. - From Artemisia maritima (Linn.) which contains 2 p.c. The seeds (5 pts.) are treated with lime (1 pt.) and water, and extracted with alcohol. The extract is freed from alcohol by evaporation and neutralised with HCl, when santonin crystallises out. In this process a lime-compound (C₁₅H₁₈O₄)₂Ca, is first formed, and afterwards decomposed by HCl, which forms santonic acid, $C_{15}H_{20}^2O_4$, whence santonin is formed by splitting off H_2O (Busch, J. pr. [2] 35,

Properties. - Flattened hexagonal prisms, which turn yellow in light. May be sublimed. Levorotatory, the specific rotation being but little affected by temperature or concentration of the solution (Hesse; Nasini, G. 13, 139). Nearly insol. cold water, v. sol. hot alcohol and ether. Poisonous, exerting an anthelmintic action, and producing temporary colour-blindness (Lavater, *Pharm. Viertelj.* 2, 110; Wells, *J. Ph.* [3] 15, 111; Martini, *C. R.* 47, 259; 50, 545). Reacts with hydroxylamine, forming an oxim (Cannizzaro, B. 18, 2746); santonin is best administered as oxim (Coppola, C. C. 1887, 1206, 1301). Not attacked by aqueous KMnO. Resinified by boiling conc. HClAq. KOH turns santonin crimson (especially in presence of alcohol) and forms $C_{15}H_{19}KO_4$. AcCl has no action. Conc. H_2SO_4 containing FeCl₃ gives a violet colour (Lindo, C. N. 36, 222; Knapp, D. P. J. 268, 42).

Reactions .- 1. By treatment with 1 mol. of PCl₅ it gives a compound C₁₅H₁,ClO₂[125° uncor.]. With 2 mols. of PCl, it gives a compound C_{1s}H_{1s}Cl₂O, [182° uncor.] (Pawlewski, B. 18, 2900). According to Klein (B. 26, 982) the compound is $C_{13}H_{13}Cl_3O_2$.—2. Reduced by HI and P to santonous acid $C_{13}H_{20}O_3$.—3. HNO₃ (S.G. 1·123) at 95° forms CO₂, succinic, oxalic, and acetic acids, and HCy (Wagner, B. 20, 1662). — 4. Boiling baryta-water forms santonic acid.-5. Hot KOHAq forms santoninic acid. -6. Potash-fusion forms formic and propionic acids (Banfi a. Chiozza, A. 91, 112).—7. An alcoholic solution exposed for some months to sunlight forms the ethyl ether of the lactonic acid of photosantonic acid (v. supra).-8. A solution in HOAc exposed to sunshine yields photo- and isophoto- santonic acids. - 9. Forms santonol on distillation with zinc-dust. Distillation over red-hot zinc-dust gives di-methylnaphthalene (Cannizzaro, G. 12, 415).-10. Cold conc. HClAq forms an isomeride C₁₅H₁₈O₃, [260°], sl. sol. hot alcohol, insol. water (Andreocci, B. 26, 1373). $[\alpha]_D = +112^\circ$. This body is not acted upon by hydroxylamine or phenyl-hydrazine, but yields an acetyl derivative C₁₅H₁₇AcO₃ [156°]. Like santonin it is insol. Na₂CO₃Aq, but sol. NaOHAq. It is reduced by zinc-dust and HOAc to $C_{15}H_{20}O_{3}$ [175°] [a]_D = -53.3° an acid isomeric with santonous acid, which when fused with potash yields propionic acid and di-methyl-(8)naphthol.-11. Br in glacial acetic acid forms C₁₅H₁₈O₃.HOAc,Br₂. Crystals (from alcohol), decomposing above 60°. On boiling with alcohol and aniline it yields bromo-santonin C₁₅H₁₇BrO₂ [151°]. On boiling with KOHAq the acetate bromide yields santonin (Klein, B. 25, 3317).

Oxim C₁₀H₁₀O.(NOH) aq. [209°] (Klein, B.

26, 411; cf. Gucci, G. 19, 367). Needles (from dilute alcohol). Ac₂O yields $C_{15}H_{18}O_2(NOAc)$ [166°] or [170°] (K.); [203°] (G.). The oxim prepared by alkaline hydroxylamine gives with HCl and FeCl, a violet-red colour [syn-oximic acid].

Bensyl ether of the oxim [152°]. HIAq forms benzyl iodide.

Phenyl-hydraside C18H18O2(N2HPh). [220°]. Pale-yellow needles (from alcohol), not decomposed by cold acids (Cristaldi, G. 17, 526; Grassi, C. C. 1887, 1163).

Chlorosantonin C₁₈H₁₇ClO₃. Formed from santonin and chlorine-water (Heldt, A. 63, 34; Sestini, Bl. 5, 202). Crystalline. Turns yellow in light.

Di-chloro-santonin C₁₅H₁₆Cl₂O₃. Formed by passing Cl for a long time through water containing santonin in suspension (Sestini). Groups of small plates (from alcohol). Coloured orangered by alcoholic potash.

Tri-chloro-santonin C₁₈H₁₈Cl₈O₈. Γ213°1. Monoclinic prisms, not coloured by sunshine, m. sol. alcohol and ether.

(a)-Metasantonin $C_{15}H_{18}O_{5}$. [161°]. (239°). '.S. 205. [a]_D = 124° in chloroform at 20°. Formed, together with (β) -metasantonin, by boiling santonic acid or parasantonide with HIAq and P (Cannizzaro, G. 4, 446, 452; 8, 318; 10, 461). Trimetric hemihedral plates (from ether). Dextrorotatory. Not affected by light. Yields C₁₅H₁₇BrO₈ [212°] and C₁₅H₁₈Br₂O₈ [184°].

(β)-Metasantonin $C_{15}H_{19}O_3$. [136°]. S.V.S. 211. [α]_D = 124°. Formed as above. Monoclinic crystals. Yields $C_{15}H_{1}$,BrO₃ [114°] and C18H18Br,O8 [186°].

Photosantonin. A name for the ethyl ether of the (a)-lactonic acid of photosantonic acid (v. supra).

SANTONOUS ACID $C_{15}H_{20}O_{5}$ i.e. $C_{16}H_{8}Me_{2}(OH).C_{2}H_{4}.CO_{2}H$? [179°]. [α]_D = 74° in alcohol at 20°. Formed by boiling santonin with HIAq and P (Cannizzaro a. Carnelutti, B. 12, 1574; G. 12, 393; 13, 385). White needles, v. e. sol. alcohol and ether, sl. sol. cold water, sol. Na₂CO,Aq. May be distilled in vacuo. Distilled over powdered zinc in a current of hydrogen it yields di-methyl-naphthalene, dimethyl-naphthol, propylene, and a little xylene. On heating with Ba(OH), above 360° it yields CO₂, CH₄, and di-methyl-naphthol C₁₂H₁₁OH [135°], which yields C₁₂H₁₁OMe [68°]. Converted by heat into the anhydride of isosantonous acid, and, finally, into di-methyl-naphthol dihydride and propionic acid.

Salts .- NaA'. Needles, ppd. by adding ether to its alcoholic solution. V. sol. water .-BaA', -*AgA': white pp., quickly turning black.

Methyl ether MeA'. [82].

Ethyl ether EtA'. [117°]. [a]_D = 78° in chloroform or 67° in HOAc at 20°. White crystals, sol. alcohol and ether. Dextrorotatory. Converted by BzCl into C13H18BzEtO, [78°]. Na added to its ethereal solution ppts. C15H16NaEtO4, whence EtI produces C₁₅H₁₈Et₂O₃ [32⁵], which on saponification yields ethyl-santonous acid

 $C_{13}H_{10}E_1O_3$ [116°], [a]_b = +75° in alcohol. Isosantonous acid $C_{13}H_{20}O_3$. [155°]. Formed by heating santonous acid, dissolving the resulting anhydride in NaOHAq, and ppg. with HClAq. Got also by heating santonous acid (1 pt.) with Ba(OH)₂ (3 pts.) in a bath of molten lead (Cannizzaro a. Carnelutti, B. 12, 1574; G. 12, 400). Small plates, sl. sol. water, sol. alcohol and ether. Inactive to light. May be distilled in vacuo.

Ethyl ether EtA'. [125°]. Crystalline. BzCl at 100° yields C₁₅H₁₈BzEtO₃ [91°]. Na added to its ethereal solutions gives no pp., but on adding alcohol C₁₅H₁₈NaEtO₃ is thrown down as an amorphous powder, readily decomposed by water. The Na derivative is converted by EtI into C₁₅H₁₈Et₂O₃ [54°], whence alcoholic potash produces C₁₅H₁₉EtO₃ [143°], v.sol. alcohol and ether.

SAPONIFICATION. The conversion of the natural fats into soap (and glycerin) by boiling with alkalis. In a more general sense it is used to denote the resolution of compound ethers into acids and alcohols. In the widest sense it is used to denote the production of an acid from a neutral substance by hydrolysis.

SAPONIN $C_{32}H_{54}O_{18}$ (Rochleder, Z. [2] 3, 632); $C_{10}H_{20}O_{10}$ (Stütz, A. 218, 231); $C_{42}H_{52}O_{17}$ (Hesse, A. 261, 373). Struthiin. Githagin. Polygalin. Polygalic acid. Senegin. A glucoside occurring in the common soapwort (Saponaria officinalis) (Schrade, N. Journ. d. Chem. 8, 548), in the Oriental soapwort (Gypsophila Struthium) (Bley, A. 4, 283; Bussy, A. 7, 168), in quillaja bark (O. Henry a. Boutron-Charland, J. Ph. 4, 249), in the corncockle (Agrostemma Githago) (Malapert, J. Ph. [3] 10, 339; Scharling, A. 74, 351; Christophsohn, Ar. Ph. [3] 6, 432, 461), in the root of Polygala Senega (Bolley, A. 90, 211; 91, 117), and in many other plants. amorphous powder, insol. alcohol and ether, v. e. sol. water. The dilute aqueous solution froths like soap when shaken; the lather is prevented by addition of alcohol. Impure saponin produces sneezing. Saponin begins to turn black at 145° (Blyth), but does not melt or sublime when heated. Baryta forms amorphous (C₁₉H₂₀O₁₀)₂Ba(OH)₂. A solution of saponin is ppd. by lead acetate. Decomposed by boiling dilute acids into sapogenin and sugar CeH12O6 ([a]_D = 52° at 25°) (Rochleder a. Schwarz, Sitz. W. 11, 339; Overbeck, N. Br. Arch. 77, 135; Schiaparelli; cf. Crawford, Pharm. Viertelj. 6, 361; Flückiger, Ph. [3] 8, 488).

Butyryl derivative $C_{19}H_{28}(C_4H_7O)_4O_{10}$ (Stütz); $C_{22}H_{45}(C_4H_7O)_4O_{17}$ (Hesse). [68°-72°]. According to Schiaparelli (G. 13, 422; Ph. [3] 14, 801), saponin obtained from soapwort has the formula $C_{22}H_{34}O_{18}$, is lævorotatory, $[a]_D = -8$ at 20°, yields amorphous $Ba_2H_2(C_{22}H_{32}O_{18})_3$, and is split up by boiling dilute H_2SO_4 into glucose and saponetin $C_{40}H_{48}O_{18}$. Formed by hydrosaponetin $C_{41}H_{22}O_2$.

sapogenin C₁₁H₁₂O₂. Formed by hydrolysis of saponin. Concentric groups of needles (from alcohol), sol. ether, insol. water. From solution in dilute potash it is ppd. by conc. KOHAq as flocculent potassium-saponin. Potash-fusion gives acetic and butyric acids and a crystalline compound [128°].

Oxysapogenin $C_{1i}H_{22}O_{3}$ is obtained by the action of HClAq on a glucoside present in the alcoholic extract of *Herniaria* (Barth a. Herzig, M. 10, 172).

SAPPANIN C₁₂H₁₀O₄. Formed, together with resorcin and a little pyrocatechin, by fusing extract of sapan wood with NaOH (Schreder, B. 5, 572). Crystallises from water in plates (containing 2aq), nearly insol. cold water, v. sol. alcohol and ether, insol. CHCl₂. Neutral in reaction. FeCl₃ colours its aqueous solution red. NaOCl gives a grass-green colour. May be distilled. Pb(OAc)₂ gives a yellowish pp. Reduces hot Fehling's solution and ammoniacal AgNO₃. HNO₃ forms tri-nitro-resorcin (styphnic acid). Yields diphenyl on distillation with red-hot zinodust. AcCl gives C₁₂H₆Ac₄O₄, crystallising from alcohol in prisms. Br and HOAc give C₁₂H₃Br₄O₄, crystallising from dilute alcohol in needles.

SAPRINE C₅H₁₈N₂. [From σαπρός, putrid.] An alkaloid in putrefying flesh (Gautier, Bl. [2]

48, 13), v. Promaines.

SARCINE v. Hypoxanthine. SARCOLACTIC ACID v. Lactic acid.

SARCOLEMMA v. MUSCLE.

SARCOSINE v. METHYL-AMIDO-ACETIC ACID. SARCOSINIC ACID C₃H₁NO₂. [195°]. Occurs in a variety of shellac known as Sonora gum (Hertz, J. 1876, 912). White silky scales, v. sol. water, insol. alcohol and ether. Cannot be sublimed. Converted by nitrous acid into lactic acid. Forms salts with acids and bases.

It is a weaker acid than alanine. SARCOSINURIC ACID v. vol. iii. p. 260.

SARSAPARILLA v. PARILLIN.

SASSAFRAS OIL. Obtained by steam-distillation from the bark and wood of the root of Laurus sassafras of North America. Contains saffrole (q.v.) and a terpene $C_{10}H_{16}$ (156°). S.G. 2 ·835.

SATIVIC ACID v. Tetra-oxy-stearic acid. SATURATION, CAPACITY OF. When a base is added to an acid, a point is reached whereat the properties of the base are neutralised by those of the acid, and the properties of the acid are neutralised by those of the base. The compound that is produced—a salt—has neither the properties of the acid nor the properties of the base. The capacity of saturation of an acid is measured by the quantity of a base, taken in formula-weights, which must be added to the quantity of the acid expressed by its formula in order to form a normal salt. The capacity of saturation of a base is measured by the number of formula-weights of an acid which must be added to a formula-weight of the base in order to form a normal salt. The study of the capacities of saturation of acids and bases leads to the classification of acids as monobasic, dibasic, tribasic, &c., and of bases as mono-acid, di-acid, tri-acid, &c. (v. Acids, vol. i. p. 49; and Acids, Basicity of, vol. i. p. 51). The notion of equivalency also arises from the study of the capacities of saturation of acids and bases (v. vol. ii. p. 446).

When chlorine combines with potassium, one atom of the metal is saturated by one atom of the halogen; but three atoms of chlorine are required to saturate a single atom of bismuth or antimony. The notion of a definite capacity of saturation has been extended from the acids and

bases to the atoms of the elements. The development of this conception leads to the hypotheses of valency and atom-linking (v. CLASSFICATION, vol. ii. pp. 213-215; EQUIVALENCY, vol. ii. p. 446; and FORMULE, vol. ii. p. 572). M. M. P. M.

SAVINE OIL. Oleum Sabinæ. The essential oil from Juniperus Sabina contains a sesquiterpene $C_{15}H_{24}$, which has a crystalline hydrochloride $C_{15}H_{24}$ 2HCl [118°] (Wallach, A.

238, 82).

SAVORY. Essence of savory (from Satureja montana) contains 40 p.c. carvacrol (232°) and two terpenes (174°) and (183°) (Haller, C. R. 94, 132). The oil from Satureja hortensis contains 20 p.c. cymene, 50 p.c. carvacrol, a terpene (179°). S.G. ½ 855, and a very small quantity of a phenol which differs from carvacrol in not being extracted by ether from its dilute alkaline solution, and yields a phosphate [69°] (Jahns, B. 15, 816).

SCAMMONY. A purgative gum-resin composed of the dried milky juice of Convolvulus scammonia and of other species of Convolvulus. It is purified by boiling with alcohol, neutralising the alkaline filtrate by dilute H₂SO₄, filtering, and evaporating (Perret, Bl. [2] 28, 522). It

contains jalapin (q. v.).

SCANDIUM. Sc. At. w. 44. Mol. w. unknown. The oxide of a new element was isolated in 1879 by Nilson (B. 12, 554) from euxenite and gadolinite; the same oxide was obtained, about the same time, by Cleve, also from Scandinavian minerals (Bl. [2] 31, 486 [1879]). The name scandium was given to the metal of the new earth by Nilson to denote the locality of the minerals from which the oxide had been obtained. In his first memoir Nilson represented scandia as ScO₂, and determined the at. w. of Sc to be c. 170; in his second memoir (B. 13, 1439) he adopted Cleve's correction, and formulated the oxide as Sc₂O₃, and gave the at. w. of the metal as 44. The element scandium has not yet been isolated.

Occurrence. — Sc₂O₃ is found, along with yttria, ytterbia, &c., in very small quantities in a few rare minerals—gadolinite, keilhauite, yttrotitanite, and euxenite. From 4 kilos. gadolinite Cleve obtained 8 g. Sc₂O₃, and 1·2 g.

Sc₂O₃ from 3 kilos. yttrotitanite.

Preparation.—The metal has not been

isolated (v. Scandium oxide, infra).

Detection.—Sc salts give a very brilliant emission-spectrum, rich in rays. Thalén (B. 12, 555) gives the wave-lengths of the most characteristic lines as 6304, 6079, 6037 in the orange; 5526 in the yellow (very marked); 5031 in the green; 4415, 4400, 4374, 4325, 4314, and 4249 in the indigo (v. also B. A. 1884, 440).

Chemical relations.—Ten years before Nilson discovered Sc, Mendeléeff asserted that an element would be found in the group of which boron is the first member, and that this element would resemble B and Al. To this unknown element Mendeléeff gave the name ekaboron, and he stated the properties of the element and its compounds in detail. The properties of the compounds of scandium correspond with great closeness with those of Mendeléeff's ekaboron (v. tables in vol. i. p. 352; cf. vol. iii. p. 815). The properties of akaboron were arrived at by

comparing the properties of pairs of elements related to one another, as regards positions in the periodic arrangement, similarly to ekaboron and its atom-analogues. The relations expressed by the following scheme were specially studied by Mendeléeff:

Eb:Al = Ca:Mg = Ti:Si = V:P = Cr:S:Eb:B = Ca:Be = Ti:C = V:N = Cr:O = Mn:F.

These statements mean, the relations between the properties of Eb and Al are similar to the relations between the properties of Ca and Mg. &c. For a brief working out of similar relations in the case of eka-aluminium v. Gallium (vol. ii. pp. 598-9). Scandium is the second even-series member of Group III. It follows B, and is followed by Y, La, and Yb in the even series of this group. So resembles Al in many respects, but the difference between these elements is shown, among other ways, in the fact that the double sulphate of Sc and K-Sc₂(SO₄)₃.3K₂SO₄—is not an alum (v. EARTHS, METALS OF THE; vol. ii. p. 424).

Scandium, chloride of, ScCl₃(?). White hygroscopic needles separate on evaporating a solution of the oxide in HClAq to a syrup; HCl is given off on heating, and an oxychloride

remains as a white powder (Cleve, l.c.).

Scandium, oxide of, Sc.O₃. (Scandia.) Prepared by ppg. solutions of Sc salts by NH₃Aq, washing the white, gelatinous, very bulky pp. of hydrated oxide, drying at 100°, and then heating strongly; also obtained by heating to redness Sc nitrate, sulphate, or oxalate. A white, light powder, resembling MgO; sol. in hot conc. H₂SO₄Aq or HNO₃Aq. S.G. 3·864 (Nilson a. Pettersson, B. 13, 1461). S.H. from 0° to 100° = ·153 (N. a. P.).

Scandia is best obtained from euxenite. The finely-powdered mineral is fused with KHSO, the product is dissolved in cold water, and ppd. by NH₃Aq, the pp. is dissolved in HNO₃Aq, the solution is boiled for some time, filtered from any pp. which forms, and ppd. by oxalic acid; the ppd. oxalates are dried, heated strongly, washed with hot water to remove the K₂CO₈, dissolved in HNO₃Aq, and the solution is evaporated to dryness and the residue heated till it melts; the fused nitrates are treated with water, when a pp. forms, consisting of oxide of Th with small quantities of oxides of Ce, U, and Fe; the filtered solution is evaporated to dryness, the nitrates are partly decomposed by heating, and then treated with water; the residue, which contains a basic nitrate of Sc, is evaporated with HNO, Aq, and the residue is again heated for some time, treated with water, and the insoluble evaporated with HNO, Aq, and so on. This process separates didymia, yttria, and terbia, the nitrates of which earths are not so readily decomposed by heat to basic nitrates as So nitrate is. By repeating this process 68 times Nilson (B. 13, 1442) obtained a nitrate solution which showed no absorption bands, and contained only salts of Sc and Yb. The solution of Sc and Yb nitrates is evaporated with conc. H₂SO₄, saturated K₂SO₄Aq is added, the salt Sc₂(SO₄)₈.3K₂SO₄ which separates (Yb sulphate remains in solution) is dissolved in water, and ppd. by NH,Aq; the ppd. Sc,O,.xH,O is washed and dissolved in HNO,Aq, and Sc oxalate is ppd. by oxalic acid; the ppd. oxalate is washed,

dried, and decomposed by heat. The Sc.O. thus obtained contains only traces of ytterbia, which may be removed by transforming the earths into nitrates, heating for some time, treating with water, again transforming the insoluble into nitrate, and so on.

Scandium, salts of (Nilson, l.c.; Cleve, l.c.; also C. R. 89,419). A few salts of Schave been prepared; they are white solids, resembling the salts of Al. The sulphate forms a double salt with K2SO4, viz. Sc₂(SO₄)₃.3K₂SO₄, which is not an alum. The salts which have been isolated are nitrate, oxalate, selenites, sulphate, and double sulphates of NH,, K, and Na. M. M. P. M.

SCATOLE v. METHYL-INDOLE.

SCHEELE'S GREEN. Commercial name for CuHAsO₃; v. Copper arsenites, vol. i. p. 306.

SCHEIBLER'S REAGENT. Composed of sodium tungstate (100 g.), sodium phosphate (70 g.), water (500 c.c.) and a little nitric acid (J. pr. 80, 211). It ppts. alkaloids.

SCHIFF'S REACTIONS. (a) For aldehydes: a solution of a rosaniline salt, bleached by SO2, is coloured purple-red by aldehydes. urea: a purple colour with furfuraldehyde and HCl. (c) For cholesterin: a red colour with conc. H.SO4 followed by NH3.

SCHULTZE'S REAGENT. A saturated solution of sodium phosphate (4 pts.) mixed with SbCl, (1 pt.). It gives a white pp. with alkaloids

SCHULZE'S REAGENT. Dry ZnCl₂ (25 pts.), KI (8 pts.), water (8.5 pts.), and sufficient iodine to be dissolved on warming for a short time.

Gives a blue colour with cellulose. SCHWEIZER'S SOLUTION. A saturated solution of freshly ppd. Cu(OH)2 in ammonia (of 20 p.c.). It dissolves silk, cotton wool, and

SCOPARIN $C_{21}H_{22}O_{10}$. Occurs, together with sparteine, in Spartium scoparium (Stenhouse, Tr. 1851, 422; A. 78, 15). Pale-yellow brittle mass, tasteless, inodorous, and neutral. Ppd. as a jelly by adding HCl to its cold ammoniacal solution, but may be obtained from cold alcohol in crystalline stars. V. sl. sol. cold water, m. sol. hot water and alcohol. Sol. alkalis. Its alkaline solution deposits a greenish-brown resin on boiling. Bleaching-powder colours its solution dark green. Br gives a bluish-green colour. HNO, forms picric acid. Potash-fusion gives phloroglucin and protocatechuic acid

(Hlasiwetz, A. 138, 190).

SCOPOLAMINE C₁,H₂NO. Hyoscine. [53°] (Schmidt); [55°] (Hesse). $[a]_D = -13.7°$ in alcohol at 15°. An alkaloid occurring in the root of Scopolia atropoides and also in hyoscyamus seeds (E. Schmidt, Ar. Ph. 230, 207; B. 25, 2601; cf. Ladenburg, A. 206, 299; 276, 345; B. 14, 1870; 25, 2388; O. Hesse, A. 271, 100). Crystalline. Not affected by nitrous acid. It is either identical with hyoscine, or sometimes confounded with that alkaloid. Decomposed by baryta-water into atropic acid and scopoline (pseudatropine, oscine) C₂H₁₂NO₂ [110°] (242° uncor.), which yields B'HAuCl, [225°], B'₂H₂PtCl₃ [230°], and C₂H₁₂BzNO₂ [59°], which gives (C₂H₁₂BzNO₂)₂H₂PtCl₄ [184°]. — B'HBr Saq. — B'HAuCl₄. [214°].—B'HCl 2aq.—B'HI.

Acetyl derivative C₁₇H₄₂AcNO₄.

-B'HAuCl.

SCOPOLEINE C₁₇H₂₈NO₅ (?). Extracted by alcohol from the root of Scopolia japonica (Eijkman, R. T. C. 3, 178). Crystals (from

SCOPOLETIN $C_{12}H_{10}O_5$ (E.); $C_{10}H_8O_4$ (T.) i.e. $C_eH_2(OMe)(OH) < CH:CH (198° uncor.]$.

Occurs in the root of Japanese Belladonna (Scopolia japonica) (Eijkman, R. T. C. 3, 189; Takahashi, C. C. 1888, 1364). Slender needles or prisms, sl. sol. cold water, v. sol. alcohol and ether. Poisonous. Its solutions exhibit blue fluorescence. KOH and MeI yield C10H,MeO4 [144°] identical with di-methyl-æsculetin, and converted by Br into C10H8BrMeO4, whence potash forms $C_{11}H_{10}O_{3}$, an acid from which dimethoxy-coumarin $C_{10}H_{10}O_{3}$ [58°] may be obtained. Methyl-scopoletin is converted by potash-fusion, extracting with MeOH and heating with MeI into C,H2(OMe)3.CH:CH.CO2Me [102°], which may be reduced to the acid C₆H₂(OMe)₃CH₂.CH₂.CO₂H.

Acetyl derivative C10H7AcO4. [176°]. Benzoyl derivative C, H,BzO,. [158°]. SCOPOLIN C, H, O, 2aq (?). [c. 218°]. Occurs in Scopolia japonica (Eijkman, R. T. C. 3, 177). White needles, sol. hot water and alcohol, insol. ether and CHCl₃. Decomposed by heating with dilute acids into glucose and scopoletin.

SCUTELLARIN C10H3O3. [199°]. Occurs in the root of Scutellaria lanceolata (Takahashi, C. C. 1889, ii. 100). Flat yellow needles, insol. cold water, v. sol. alcohol and ether, sol. NaOHAq and Na₂CO₃Aq. Does not reduce Fehling's solution. Does not react with phenyl-hydrazine. Is not poisonous.

SCYLLITE. A neutral substance resembling inosite, occurring in cartilaginous fishes (Frerichs a. Städeler, J. pr. 73, 48). SEA SALT. Comm

Common name of sodium

chloride (q. v. in this volume).

SEA WATER v. WATER (in this volume).

SEBACIC ACID C₁₀H₁₈O₄ i.e. (CH₂)₈(CO₂H)₂

Ipomic acid. Pyrolic acid. Mol. w. 202 Mol. w. 202. [133°]. (295° at 100 mm.) (Krafft, B. 22, 816). S. 1 at 17°; 2 at 100° (Neison a. Bayne, J. 1874, 625). H.C.p. 1,295,668 (Louguinine, C. R. 107, 597; 1,296,800 (Stohmann, J. pr. [2] 40, 202). H.F. 264,200 (S.).

Formation.-1. Dry distillation of oleic acid or of fats containing ole (Thénard, Ann. Chim. 39, 193; Dumas a. Péligot, A. Ch. 57, 332; Redtenbacher, A. 35, 188).—2. By distilling castor oil or ricinoleïc acid with KOH (Bouis, A. 80, 303; 97, 34.—3. By the action of HNO₃ on spermaceti, stearic acid (Arppe, A. 124, 98), convolvulic acid, jalapin, jalapic acid (Mayer, A. 83, 143; 95, 160).—4. One of the products of the distillation of crude fatty acids in a current of superheated steam (Cahours, C. R. 94, 610).-5. The ethyl ether is formed by electrolysis of CO,Et.CH,.CH,.CH,.CO,K, the yield being 20 p.c. of the theoretical amount (Crum Brown a. Walker, A. 261, 120).—6. By oxidation of hendecenoic acid by HNO, (Becker, B. 11, 1414).-7. By reduction of tetracetylene dicarboxylic acid C_s(CO₂H), by sodium-amalgam (Baeyer, B. 18, 2271).

Preparation.—Castor oil is heated with conc. KOHAq for some hours at 40°, and the resulting hard crust crushed, dried, and heated in an iron vessel till the mass begins to turn brown. The product is shaken with cold water and the solution ppd. by HCl (Witt, B. 7, 219).

Properties.—White needles or plates, sl. sol. cold water, v. sol. alcohol and ether. Reddens

litmus.

Reactions. - 1. HNO, forms succinic, pimelic, and adipic acids (Carlet, J. Ph. [3] 24, 176; Wirz, A. 104, 280; Arppe, Z. 1865, 295; cf. Schlieper, A. 70, 121).—2. KMnO, yields succinic, glutaric, and adipic acids (Carette, C. R. 101, 1498).—3. Distillation with lime yields decinene and heptoic aldehyde (Petersen). Distillation with BaO forms octane. — 4. MeONa at 300° forms ennoic acid (Mai, B. 22, 2133).—5. Glycerin at 200° forms (C3H3(OH)2)2A" which solidifies at -40° (Berthelot, A. Ch. [3] 41, 293).—6. Br at 165° forms C₁₀H₁₆Br₂O₄ [115°], which yields Na₂A", KHA", BaA" 2aq, CaA" 2aq, PbA", Ag₂A", and Me₂A" [50°] (Claus B. 20, 2882).

Salts.-K2A': crystalline powder, v. sol. water.-KHA": minute needles, v. sol. water (Neison, C. J. 27, 301). -Na,A": minute crystals. -NaHA": crystalline powder. -CaA" (dried at 100°): thin scales, sl. sol. water.—CaH,A" white pp., sl. sol. water.—BaH₂A''₂: needles (from water), v. sol. water.—BaA'': crystalline crusts, v. sol. water.—SrA".—MgA": white crusts, v. sol. water.—MgH₂A"₂: needles, v. sol. water.—ZnA": white pp., sl. sol. water.—NiA".—CoA".—CoA" 2aq.—PbA".—CuA": emeraldgreen pp.-HgA": insoluble pp.-Hg2A": amorphous powder.—AlH₃A"₃: white powder, v. sol. hot water.—Ag₄A": white powder, v. sol. HNO₃. -Aniline salt (PhNH2),H2A". [134°]. very plates, sol. warm water. May be sublimed (Gehring, C. R. 104, 1451).

[38°]. ether Me,A". Methyl (Neison, C. J. 29, 314; cf. Carlet, C. R. 37, 130).

Pearly crystals (from ether).

Mono-ethyl ether EtHA". Solidifies at

10° as snow-white needles.

Ethyl ether Et, A". [c. 5°]. (308° cor.). S.G. \(\frac{15}{15}\) 9682; \(\frac{25}{25}\) 9605. M.M. 14·496 at 14° (Perkin, C. J. 45, 518).

Butyl ether (C4H0)2A". (344°). S.G. 9.942; 15.933. Formed by the action of gaseous HCl on a mixture of sebacic acid and butyl alcohol at 150° (Gehring, C. R. 104, 1289, 1625). Oil, miscible with alcohol and ether. Chlorine in sunlight forms (C,Cl,)2A", crystallising in hexagonal prisms [172°].

Isoamyl ether (C,H,,),A". (above 360°). S.G. 12 '951. Oil. Alcoholic NaOH forms (CaH11)HA", which solidifies at 10°, forming a erystalline mass, splitting up at 325° into sebacic acid and $(C_5H_{11})_2A''$. Chlorine in excess forms, in 160 hours, $(C_5Cl_{11})_2A''$ [179°], crystallising in volutile trimetric manner.

volatile trimetric prisms, 81. sol. alcohol.

Chloride C_aH_{1a}(COCl)₂. (220° at 75 mm.).

Formed from sebacic acid (40 g.) and PCl_a(85 g.)
(Auger, A. Ch. [6] 22, 361). Thick liquid, decomposed by heat. Benzene and AlCl₃ give $C_aH_{1e}(CO.C_uH_b)_2$ [89°] and $C_eH_b.CO.C_aH_{1e}.CO_gH$ [79°].

Anhydride C₈H₁₆<a>CO O. [78°]. Formed by heating the Na salt with the chloride (Auger, A. Ch. [6] 22, 862). Minute yellowish crystals (from benzene).

Amide C.H. (CO.NH.). Formed from the ethers and NH. Aq (Rowney, A. 82, 123; Carlet, C. R. 37, 128). Groups of minute needles. m. sol. hot water.

Amic acid C₈H₁₆(CO₂H)(CONH₂). Formed from EtHA" and NH, Aq, and got also by heating (NH₁)₂A" (Rowney, C. J. 4, 334; Kraut, J. 1863, 358). Crystalline, v. sol. alcohol and hot water.

Anilide C.H. (CONPhH). [198°] (Pellizzari, A. 232, 147). (above 360°). Formed by heating sebacic acid (1 pt.) with aniline (1 pt.) at 215° (Gehring, C. R. 104, 1451; cf. Maillot, C. R. (2007). C. R. 87, 737). Scales (from alcohol), v. sl. sol. ether.

Nitro-anilide C₈H₁₆(CO.NHC₆H₄.NO₂)₂. [116°]. Formed by the action of HNO, on the anilide (Gehring, C. R. 104, 1716). Slender needles (from water). Tastes sweet. May be sublimed.

References. - PERCHLOR- and OXY-SEBACIC ACID.

SEBACIN v. DECINENE.

SEBACOXYL-AMIDO-BENZOIC ACID CARBOXY-PHENYL-SEBACAMIC ACID.

SEBACYL-DIBENZAMIC ACID v. Sebacyl derivative of m-Amido-Benzoic Acid.

SELENATES. Salts of selenic acid (H2SeO4), M21SeO4 and M1HSeO4. These salts are formed by dissolving metallic oxides, hydroxides, or carbonates in H₂SeO₄Aq; the more insoluble selenates are formed from the alkali salts by double decomposition. The selenates generally resemble, and on the whole are isomorphous with, the sulphates (von Gerichten, A. 168, 214). The selenates generally crystallise with less water than the sulphates under the same conditions of temperature (Topsöe, Ar. Sc. 45, 76). Selenates are generally undecomposed at red heat; heated in H, or with Na₂CO₃, or C, they give selenides; boiled with HClAq, they give off Cl and leave SeO₂Aq. For the crystallographic measurements of selenates, v. Topsöe (l.c.); Topsöe a. Christiansen (A. Ch. [4] 31); and Fabre (C. R. 105, 114). For measurements of the specific volumes of selenates, v. Pettersson (B. 9, 1559, 1676) and Schröder (J. pr. [2] 22, **à**32).

Aluminium selenates. The normal salt. Al₂(SeO₄)₃, is a white solid; by dissolving AlO₃H₃ in H₂SeO₄Aq, and evaporating. Yields various basic salts by heating with AlO₂H₃ or by partially ppg. the normal salt in solution by NH₃Aq. Forms alums with alkali metal selenates. For accounts of these alums v. Wohlwill (A. 114, 162); Weber (P. 108, 615); Pettersson (B. 9, 1559, 1676); Fabre (C. R. 105, 114). For the alums formed by $\mathrm{Al}_2(\mathrm{SeO}_4)$, and sulphates of alkali metals, and by selenates of alkali metals with Al₂(SO₄)₃ and Cr₂(SO₄)₃, v. von Gerichten (B. 6, 162).

Ammonium selenates. The normal salt. (NH₄),SeO₄, is formed by neutralising H₂SeO₄Aq by NH, Aq and evaporating; it seems to exist in two different crystalline forms (v. von Lang, W. A. B. 45 (2nd part), 109; Topsõe, ibid. 66, 5). When heated, it gives off NH, and leaves the acid salt (NH₄)HSeO, (Cameror a. Davy, C. N. 38, 133). This acid salt is also formed by adding excess of H.SeO.Aq to NH.Aq (Topsoe, Lc.).

y y

Barium selenate BaSeO4. A white salt, obtained by decomposing an alkali selenate by BaCl, Aq, filtering, and washing. Insol. water and HNO, Aq; boiled with HClAq gives BaSeO, which dissolves. S.G. 4.67 at 22° (Schafarik, J. 1863. 15; Michel, C. R. 106, 878).

Beryllium selenate BeSeO₄. 4aq. Rhombic

crystals (Topsöe, W. A. B. 66, 5).

Cadmium selenate CdSeO, 2aq. By dissolving CdO.xH₂O in H₂SeO, Aq, and crystallising; lustrous, rhombic plates; S.G. 3-632 (von Hauer, J. pr. 80, 214). Forms double salts with NH, and K (Topsöe, l.c.; von H., l.c.; von Gerichten, B. 6, 162).

Calcium selenate CaSeO .. 2aq. Resembles CaSO₄; obtained by ppg. N₂SeO₄Aq of Ca(NO₃),Aq, dissolving in cold water, and ppg. by heating (von Hauer, l.c.). Also formed by heating (von Hauer, l.c.). Also formed by

C. R. 106, 878).

Chromium selenate. Not isolated. Double salts, which are alums, are known, of the composition Cr₂(SeO₄)₂.M₂SeO₄. 24aq (Wohlwill, A. 114, 162; Fabre, C. R. 105, 114; Pettersson, B. 6, 1466). Alums are also known of the composition Cr₂(SeO₄)₃.M₂SO₄. 24aq (von G., l.c.).

Cobalt selenate CoSeO. 7aq. Hydrated salts have been obtained with 6aq and 5aq (v. Mitscherlich, P. 11, 327; Topsöe, W. A. B. 66, 5). Forms double salls with K₂SeO₄, and (NH₄)₂SeO₄

(T., l.c.; von G., l.c.; von H., l.c.).

Copper selenate CuSeO . 5aq. Blue, transparent, lustrous crystals, isomorphous with the sulphate. By dissolving CuO₂H₂ in H₂SeO₄Aq (Mitscherlich, P. 11, 330). Forms double salts with selenates of NH, K (Topsöe, l.c.; von G., l.c.); also with Mg and Zn selenates (Wohlwill, l.c.).

Didymium selenate Di₂(SeO₄)₃ (Cleve, Bl. [2] 43, 362). Also with 5aq (C., l.c.); and with 6aq (Frerichs a. Smith, A. 191, 350). Forms double salts with NH, and K selenates (C., l.c.).

Iron selenate FeSeO . 7aq (Ferrous selenate). By evaporating, in H, a solution of Fe, or FeCO₂, in H₂SeO₄Aq; resembles FeSO₄, 7aq (Topsöe, *l.c.*). Obtained by Wohlwill (A. 114, 189) with 5aq. Forms a double salt with K₂SeO₄ (von G., l.c.).

Lanthanum selenate La₂(SeO₄), 12aq (F. a.

S., l.c.).

Lead selenate PbSeO4. A white powder; by ppg. K₂SeO₄Aq by Pb(NO₂)₂Aq (Schafarik, J. 1863. 1*5*).

Magnesium selenate MgSeO₄. 6aq. monoclinic crystals; isomorphous with the sulphate; S.G. 1.928 (Topsöe, i.c.). Forms double salts with NH, and K selenates (T., l.c.; von G., l.c.).

Nickel selenate NiSeO4. aq. By evaporating solution of NiCO, in H.SeO.Aq (Mitscherlich, l.c.; Wohlwill, l.c.). Forms double salts with selenates of NH, and K (von G., l.c.; von Hauer, J. pr. 80, 214).

Mercury selenates. 1. Mercuric selenate, HgSeO. By digesting HgO with H,SeO.Aq, or by adding excess of H,SeO.Aq to mercuric acetate, and evaporating; a yellowish-white solid. Decomposed by water, giving a basic salt HgSeO, 2HgO (Cameron a. Davy, Trans. I. 28, 187). 2. Mercurous selenate, Hg,SeO4; by ppg. HgNO, Aq by K, SeO, Aq (C. a. D., lc.). With

NH, Aq forms (NHg2)2SeO4. 2aq.

Potassium selenates. The normal salt, K₂SeO₄, is obtained by fusing Se, K₂SeO₂, or native lead selenide, with KNO, dissolving in hot water, allowing to cool partly, pouring off from excess of KNO, which separates, and collecting the crystals which separate from the mother-liquor. Equally sol. hot and cold resembles K₂SO₄. The acid salt KHSeO₄ is obtained like, and closely resembles, KHSO₄ (M., l.c.). K₂SeO₄ forms many double salts; with MSO₄, M = Cd, Co, Cu, Fe, Mn, Hg, and Zn (von Gerichten, B. 6, 162); with Al₂(SeO₄), and Cr2(SeO4)3, the compounds being alums (von G., i.c.; Pettersson, B. 9,1559, 1676; Schröder, J. pr. [2] 22, 432; Fabre, C. R. 105, 114).

Silver selenate Ag₂SeO₄. Similar to, and isomorphous with, Ag₂SO₄ (Mitscherlich, *l.c.*). Sodium selenate Na₂SeO₄. Prepared like

K₂SeO₄; separates with 10aq when crystallised below 40° (M., l.c.).

Strontium selenate SrSeO4. Obtained by fusing SrCl2 with NaCl and an alkali selenate; S.G. 4.23. Exactly resembles celestine in crystallographic and optical properties (Michel, C.R.

106, 878).

Thallium selenate Tl₂SeO₄. By dissolving Tl2CO, in H2SeO, Aq, and evaporating. Forms white, prismatic needles, isomorphous with K2SeO4; sl. sol. cold water; insol. alcohol or ether (Kuhlmann, Bl. 1864 [1] 330; Oettinger, Zeit. Chem. Pharm. 1864. 440). Forms double salts, which are alums, with $Al_2(SeO_4)_3$ and with $Cr_2(SeO_4)_3$ (Fabre, C. R. 105, 114).

Zinc selenate ZnSeO. 7aq. Various hydrates are known (v. Mitscherlich, P. 11, 326; 12, 144; Topsöe, l.c.). Forms a double salt with Tl₂SeO₄ (Werther, Bl. 1865 [1] 60); also with K₂SO₄

(von Gerichten, B. 6, 162).
THIOSELENATES M¹₂SeSO₃. (Selenothiosulphates. Selenosulphates.) Solutions of the normal alkali sulphites dissolve Se, forming these salts. The acid has not been isolated. The salts correspond with the thiosulphates, S

being replaced by Se.

Potassium thioselenate K, SeSO, aq. Formed by mixing K₂SeAq with SO₂Aq (Cloëz, Bl. 3, 112; [2] 4, 419). Better prepared by digesting K₂SO₃Aq with excess of Se, filtering the cold liquid from Se, and concentrating without heat; crystals of K₂SO₂ mixed with K₂SeS₂O₂ (v. infra) separate, and K₂SeSO₂ crystallises from the mother-liquor (Cloëz, *l.c.*; Rathke, *J. pr.* 95, 1). White, lustrous, six-sided tablets; deliquescent in moist air; effloresce over H2SO4. Decomposed rather easily by heat, giving K polyselenides; cold water separates Se, and forms K₂SeS₂O₆ (v. infra); acids evolve SO₂ and separate Se; BaCl, Aq ppts. BaSO, and Se from an aqueous solution of the salt; an ammoniacal solution of Ag₂O forms Ag₂Se and K₂SO₄Aq on heating

DITHIO - TRISELENATES MI2SeS2O. (Selenotrithionates.) These salts may be regarded as derived from trithionates (Mi.S.O.) by replacing S by Se; as the salts M.S.O. are called trithionates, so the salts M₁Se₂O₂ might be called triselenates if they were isolated, and the salts M₁S₂SeO₂ may be called dithio-triselenates. Potassium dithio-triselenate K₂SeS₂O₄. This salt is formed in the preparation of potassium thioselenate (v. supra); if KHSO₂ is present in the K₂SO₃ used considerable quantities of K₂SeS₂O₄ are produced (Rathke, J. pr. 95, 1). It is also obtained by mixing K₂SeSO₃ with excess of K₄SO₃ and conc. H₄SeO₄Aq. Forms lustrous, greenish-yellow, monoclinic prisms, isomorphous with K₂S₃O₆; stable in air; sol. water without change (Rathke, J. pr. [2] 1, 33). Acids evolve SO₂ and ppt. Se, on warming; ammoniacal solution of Ag₂O ppts. Ag₂Se, forming also K₂SO₄Aq and H₂SO₄Aq.

Dithio-triselenic acid, H₂SeS₂O₆, is said to be present in the solution obtained by the action of excess of SO₂ on SeO₂Aq (Schulze, J. pr. [2] 32, 390).

M. M. P. M.

SELENHYDRIC ACID H₂Se; v. Hydrogen selenide, vol. ii. p. 724.

SELENIC ACID H SeO. Mol. w. not determined. S.G. 2.9508 at 15°; S.G. of superfused acid, 2.6083 at 15° (Cameron a. Macallan, Pr. 46, 13). Melts at 58°; the molten acid may be cooled below 5° without solidifying (C. a. M., Lc.). H.F. [Se,O',Aq]=76,660; [SeO²Aq,O]=20,500 (Th. 2, 279).

Formation.—1. By passing Cl into H₂SeO₄Aq, and evaporating.—2. By fusing Se or SeO₂ with KNO₃, ppg. BaSeO₄ from solution of the fused mass, decomposing by K₂CO₃, filtering from BaCO₃, neutralising any K₂CO₃ present by HNO₄Aq, ppg. PbSeO₄, and decomposing this salt by H₂S (von Gerichten, A. 168, 214).—3. By heating Se with HNO₃Aq, boiling with HClAq (to reduce any H₂SeO₄ to H₂SeO₄), ppg. traces of H₂SO₄ by adding a little Ba(NO₃)₂Aq, filtering from BaSO₄, neutralising by Na₂CO₃, evaporating from BaSO₄, neutralising by Na₂CO₃, evaporating to dryness, heating to dull redness, boiling with HNO₃Aq, ppg. PbSeO₄ by addition of Pb(NO₃)₂Aq, and decomposing by H₂S (Fabian, A. Suppl. 1, 241).—4. By fusing SeO₂ with excess of KNO₃, boiling with HNO₄Aq, ppg. CaSeO₄ by addition of Ca(NO₃)₂Aq, decomposing the CaSeO₄ by addition of CdC₂O₄, and decomposing the CdSeO₄ so obtained by H₂S (von Hauer, J. pr. 80, 214).—5. By adding AgNO₃ or Ag₂CO₃ to a solution of pure SeO₂ in water, digesting the ppd. Ag₂SeO₃ with water and a slight excess of Br, filtering from AgBr, and evaporating the filtrate, which contains H₂SeO₄ (Thomsen, B. 2, 598).—6. By oxidising H₂SeO₄Aq by K₂CrO₄Aq, PbO₂, or MnO₂ (Wohlwill, A. 114, 169, 176).

Preparation.—SeO, is heated till sublimation begins; it is then dissolved in water, and BaOAq is added so long as a few drops, when filtered, continue to give a pp. with BaOAq; the filtered liquid is now free from H₂SO, and H₃SeO₄, it is evaporated to dryness, and the residual SeO₂ is heated till it sublimes (Thomsen, B. 2, 598). The pure SeO₂ thus obtained is dissolved in water, excess of AgNO₄Aq is added, the ppd. Ag₂SeO₃ is washed with water and shaken with water and Br—at first Br is added, and then BrAq till the liquid shows a faint orange colour; the liquid filtered from AgBr contains only H₂SeO₄, it is evaporated and treated as described hereafter (T., l.c.). Another method consists in neutralising the solution of pure SeO₂ by K₂CO₃, evaporating to dryness, fusing with a

little KNO₃ for some time, dissolving in water, slightly acidifying with HNO₃Aq, boiling for some time to decompose any nitrite present, allowing to cool, adding Pb(NO₃)₂Aq, washing the ppd. PbSeO₄, suspending this salt in water, and decomposing it by a rapid stream of H₂S; the filtered liquid is then evaporated (von Gerichten, A. 168, 214).

The solution of H. SeO, is concentrated by evaporation in an open dish on a water-bath so long as water passes off; an acid containing c. 83-84 p.c. H₂SeO₄ is thus obtained; this acid is placed in a small flask connected with a short wide tube containing solid KOH, which is connected with a large U tube also containing solid KOH; the U tube is connected with the receiver of an airpump, or with a vessel in connection with a good water-pump. The air is pumped out of the apparatus, and when a good vacuum is obtained the small flask is heated to 100° so long as acid distils over (this is known by the KOH effervescing slightly, as ordinary KOH contains K.CO.); the temperature is then raised to 180° (an oilbath being used to heat the flask), and heating at this temperature is continued so long as acid distils off; as soon as the potash ceases to be acted on the heating is stopped and the flask is at once cooled; a very good vacuum must be maintained throughout the process. H.SeO. crystallises as the flask is cooled. A small quantity of H₂SeO₄Aq should be evaporated thus at a time (Cameron a. Macallan, Pr. 46, 13).

Properties.—A white crystalline solid, crystallising in long hexagonal prisms, and melting at 58° to a colourless oily liquid. The liquid acid may be cooled below 5° without solidifying if it is not stirred; a crystal of H₂SeO₄ dropped into the liquid causes it to solidify at once, the temperature rising to 58°. The liquid acid is specifically lighter than the solid (v. data at beginning of article) (C. a. M., l.c.). H₂SeO₄ is very hygroscopic; it mixes with water, with production of heat.

The following table is taken from Cameron

and Macallan (l.c.) :-

8.G. H₂ScO₄Aq at 15° S.G. H,SeO.Aq at 15° P.c. P.c. H.ScO. H,SeO. 99.73 2.6083 90 2.384899.5 2.6051 89 2:3568 99.0 2.5975 88 2.329187 2.3061 98.5 2.586386 2.2795 98.0 2.5767 85 2.255897.5 2.5695 2.225884 97 2.56012.5388 83 2.1946 96 2.5163 82 2.1757 95 81 94 2.4925 2.1479 93 2.4596 80 2.1216 92 2.4322 79 2.0922

Reactions.—1. With water much heat is produced, and hydrates are probably formed (v. infra, Hydrates of setenic acid). H.SeO, acts as a dehydrating agent, similarly to H.SO. (C. a. M., l.c.).—2. Heated in vacuo begins to decompose into O, H.O, and SeO, at c. 200°; the water formed dilutes the remaining acid, which then distils over. When dilute H.SeO.Aq is heated at ordinary pressure water distils off till temperature rises to 205°, when the acid has the

73.5

1.9675

2.4081

91

composition H₂SeO₄.H₂O; traces of H₂SeO₄ then distil over; at 260° white fumes of H2SeO, come off. When H2SeO4 is strongly heated at the ordinary pressure, some of it is decomposed to SeO₂, O, and H₂O; the remaining acid is diluted by the water till a dilute acid distils off (C. a. M., l.c.).—3. Dilute H₂SeO₄Aq dissolves many metals, with evolution of H; warm conc. acid dissolves copper and gold, with partial reduction to H.SeO. -4. Digestion with hydrochloric acid reduces H₂SeO₂Aq to H₂SeO₃Aq, with evolution of Cl; the acid is not reduced by sulphur dioxide nor by sulphuretted hydrogen .-5. Sulphur reacts with liquid H2SeO, at c. 60°, forming a deep-blue, very unstable substance (? SeSO₃) which decomposes to SO₂ and H₂SeO₃ (C. a. M., l.c.). - 6. Selenion dissolves in H₂SeO₄, giving a green unstable substance—? Se₂O₃, (C. a. M., l.c.).—7. Tellurium dissolves in H₂SeO₄, forming a reddish body, probably TeSeQ₃ (C. a. M.).—8. Heated with phosphoric anhydride, SeO2 is given off; but at a lower temperature crystals were obtained which C. a. M. think were SeO,.

HYDRATES OF SELENIC ACID. The monohydrate H, SeO, H,O was obtained by C. a. M. (l.c.) by diluting H₂SeO₄ to 88.96 p.c., freezing at -23°, melting and re-crystallising by freezing several times. This hydrate forms long needles, melting at 25°, and remaining liquid even 50° below its melting-point; Pickering (priv. comm.) gives melting-point as 16.6°. S.G. of solid hydrate = 2.6273 at 15°; S.G. of superfused hydrate = 2.3557 at 15°. C. a. M. did not succeed in obtaining other hydrates as solids. For salts of H₂SeO₄ v. Selenates, p. 433.

M. M. P. M. **SELENIDES.** Se combines with most metals when heated with them. Se also combines with many non-metals, e.g. with B, Br, Cl, F, H, I, C, P, and S. Many metallic selenides are formed by heating the elements together (Uelsmann, A. 116, 122; Schneider, A. 97, 192; Little, A. 112, 211); also by the interaction of Se and metallic salts (Schneider, P. 127, 642; l'otilitzin, B. 12, 697); sometimes by the interaction of Se and metallic sulphides at high temperatures (P., l.c.). For thermal data regarding the formation of many metallic selenides v. Fabre (C. R. 102, 1469; 103, 269, 345). Selenides of the alkali metals are sol. water; they are formed by saturating the hydroxides in water by H₂Se (Fabre, C. R. 102, 613), by reducing the selenites by C (Wöhler a. Dean, A. 97, 1), by reducing selenates by H at 200° (Fabre, C. R. 102, 1469), and by heating the elements together (Uelsmann, A. 116, 122). The selenides of metals other than the alkali metals are insol. M. M. P. M.

SELENION. Se. (Selenium.) At. w. 78.8. Mol. w. 157.6 (v. infra). M.P. 217° for crystalline variety (Hittorf, P. 84, 214; Draper a. Moss, C. R. 33, 1; v. Action of heat on selenton). Boils at 676°-683° (Carnelley a. Williams, C. J. 35, 563). S.G. c. 4.7 crystalline; c. 4.3 amorphous (for more details v. description of varieties of Se, infra). V.D. 111 at 860°, 92 at 1040°, 82 at 1420° (Deville a. Troost, C. R. 49, 239; 56, 891). S.H. crystalline -20° to +7° = 0732, crystalline 20° to $98^{\circ} = .0762$, amorphous -27° to $+8^{\circ}$ - 0746 amorphous 19° to 87° - 1036 (Regnault. A. Ch. [3] 46, 257). C.E. vitreous at 40° = 0000368 (Fizeau, C. R. 68, 1125; cf. Spring. B. 14, 2580). For E.C. v. infra. $\mu_{A} = 2.654$, $\mu_{a} = 2.692$, $\mu_{D} = 2.98$ (v. Sirks, P. 143, 429). For emission and absorption spectra v. B. A. **1884.** 440. H.C. $[Se,O^2] = 57,080$; $[Se,O^2,Aq] = 56,160$; $[Se,O^3,Aq] = 76,660$ (*Th.* 2, 274).

Historical.—In 1817 Berzelius (S. 23, 309) obtained a new element from the crystals formed in the leaden chamber of a sulphuric acid works at Gripsholm in Sweden; as the new element was found to resemble tellurium (discovered in 1798 by Klapstock), Berzelius gave it the name selenion ($\sigma \in \lambda h \nu \eta = \text{the moon}$). As Se is distinctly a non-metallic element, and as the termination um is characteristic of the names of metals, it seems altogether improper to use the name selenium rather than selenion.

Occurrence.—In small quantities uncombined with other elements; in combination with various metals, such as Bi, Cu, Pb, Hg, Ag, generally along with sulphur. Most selenion-containing metallic sulphides are very poor in Se; but certain selenides from the Argentine Republic contain from 29 to 48.5 p.c. Se, combined with Cu, Pb, and Ag. Minerals containing Se are fairly widely distributed, but occur only in small quantities (Stromeyer, S. 43, 452; MacIvor, C. N. 56, 251; Nordström, B. 12, 1723; Pisani, C. R. 88, 391; Hensler a. Klinger, B. 18, 2556; Stelzner, J. 1874. 1234). When sulphides which contain Se are roasted, the Se condenses in the flues. The deposits that form in sulphuric acid chambers often contain Se; and many specimens of the acid also contain this element. Commercial HClAq sometimes contains Se, derived from the H₂SO₄ used in making the HClAq (Kemper, J. 1860. 84; Nilson, B. 7, 1719; Lamy, C. R. 74, 1285; Scheurer-Kestner, C. R. 74, 1286; Personne, C. R. 74, 1199; Drinkwater, An. 8, 63).

-1. By Formation .passing SO, into H_2SeO_3Aq ; $H_2SeO_3Aq + 2SO_2 + H_2O$

=2H₂SO₄+Se.—2. By allowing the solution of an alkali selenide to stand in air (Böttger, J. pr. 71, 512).—3. By adding acid to solution of a thio-selenate (Böttger, J. pr. 94, 439), or to KCNSeAq (Oppenheim, J. pr. 71, 266).—4. By the action of such metals as Zn or Fe on H₂SeO₃Aq made strongly acid by HCl or H₂SO₄.

5. By heating (NH₄), SeO₃ or (NH₄), SeO₄.

Preparation.—1. From the flue-dust formed in roasting sulphides containing Se. The fluedust is washed with water, dried, and distilled; this method is applicable to dusts rich in Se. Dusts poorer in Se are washed, the heavier particles are lixiviated with dilute HClAq and then with water, dried, fused with pearlash and soda, and washed with water: the solution of alkali selenide thus obtained is allowed to stand in the air, and the Se which ppts. is washed, dried, and distilled (Böttger, J. pr. 71, 512). Or the washed dust is made into a thin paste with equal volumes of H2SO, and H2O, heated to boiling, with addition from time to time of a little HNO, or KClO, till the red colour of the liquid is destroyed, conc. HClAq is added equal to half the volume of the liquid, and the whole is evaporated to one-fourth its bulk, whereby Cl is evolved and H_SeO, is reduced to H_SeO,; SO, is passed into this solution, the ppd. Se is

SELENION.

washed, dried, and distilled (Wöhler, G. A. 69, 264).-2. From the deposit in the leaden chambers of sulphuric acid works. The washed deposit is boiled with conc. Na2SO, Aq till it is black (from formation of PbS), and the liquid, which contains Na SeO, is filtered into dil. HClAq, whereby Se is ppd. This process is repeated several times (Böttger, J. pr. 94, 439). Or the washed deposit is digested at 80°-100° with fairly conc. KCNAq until the red colour of the liquid has changed to grey, the liquid is filtered, the residue washed with boiling water till HClAq ceases to produce a reddish opalescence in the washings, and excess of HClAq is added to the filtrate and washings, whereby Se is ppd. (KCNAq + Se = KCNSeAq;KCNSeAq + HClAq = KClAq + HCNAq + Se). The ppd. Se is dissolved in HNOsAq, the solution is evaporated (slowly, to prevent loss of SeO2) to dryness, the SeO2 is sublimed in a tube of hard glass in a stream of dry dust-free air, the sublimate is dissolved in water, HClAq is added, and Se is ppd. by SO₂ (Oppenheim, J. pr. 71, 266). Se, as prepared by reducing cold H₂SeO₂Aq, is

a red amorphous powder; an amorphous vitreous variety is obtained by melting ppd. Se and cooling very rapidly by pouring on to a porcelain plate (Hittorf, P. 84, 214; Rammelsberg, P. 152, 151; Regnault, A. Ch. [3] 46, 257); a black crystalline variety of Se is prepared by allowing conc. K, SeAq or Na, SeAq to stand in air until a crust forms on the surface (H., l.c.; R., l.c.), also by crystallising any form of Se from Se₂Cl₂ (Rathke, A. 152, 181); a red crystalline variety is prepared by slowly and repeatedly heating a solution of Se in CS, in a sealed tube to somewhat under 100°, and allowing to cool (R., l.c.; Rathke, A. 152, 181), also by subliming amorphous Se (Bornträger, D. P. J. 242, 55); a grey metal-like variety is prepared by very slowly heating amorphous or red crystalline Se to 96°-97° (H., l.c.; R., l.c.; Fabre, C. R. 103, 53). According to Schutze (J. pr. [2] 32, 390) a colloidal soluble form of Se is obtained by reducing H₂SeO₃Aq by SO₂; no acid must be present. The colloidal Se remains in solution, colouring the liquid dark red. The solution is not changed by boiling but on adding an acid or a salt Se is ppd. in red flocks.

Properties.—Se exists in several, probably in five, forms.

I. Amorphous selenion; sol. in CS. (i.) Red-powdery amorphous Se forms sealing-waxred flocks, or when dried at a low temperature a dark-red powder; when formed by reduction of hot H.SeO, Aq it appears as a black powder (Hittorf, P. 84, 214). S.G. at 20° = 4.3 (Rammelsberg, P. 152, 151; Schaffgotsch, P. 90, 66); 4.2 (Rathke, A. 152, 181). This variety is a non-conductor of electricity (Hittorf, l.c.). It is sol. CS.; S. at b.p. of CS₁(46:6°) = 1, at 0° = 016 (Mitscherlich, J. 1855. 314); according to Rammelsberg (l.c.) the solubility in CS₂ varies much. For S.H. v. beginning of this article. Petersen (Z. P. C. 8, 601) gives H.C. of this variety $[Se,O^2] = 57,250$, and at. vol. 18.4; the change from this form to any other form of Se is attended with contraction and the production of a small quantity of heat. For action of heat v. infra.

(ii.) Amorphous vitreous Se forms a black lustrous solid, appearing red in thin layers. S.G. 4.282 at 20° (Schaffgotsch, J. pr. 43, 308; 60, 312). Sol. in CS₂. No fixed melting-point; when heated it softens at 60°, and becomes gradually less viscous till at 250° it is quite liquid; if this liquid is rapidly cooled, the vitreous variety is re-formed (Hittorf, Lc.; Draper a. Moss, C. N. 33, 1). Heated very slowly to 96°-97° this variety changes to metallic Se, with production of heat (R., l.c.; Regnault, A. Ch. [3] 46, 257; Fabre, C. R. 103, 53).

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II. Crystalline selenion; insol. in CS_x —(i.) Black crystalline Se. Small, microscopic, glittering leaflets. S.G. 4.8 (R., l.c.); 4.76 to 4.78 at 15° (Mitscherlich, J. 1855. 314). Insol. CS_2 . H.C. [Se,O'] = 55,820; at. vol. = 16.5; change from amorphous Se to this form is accompanied by slight production of heat and contraction (Petersen, Z. P. C. 8, 601).

(ii.) Red crystalline Se. Small, thin, transparent, lustrous crystals; in thicker layers appears black and opaque. S.G. 4.46 to 4.5 at 15° (R., l.c.; M., l.c.; Rathke, l.c.). Melts at 217° (H., l.c.; D. a. M., l.c.). Insol. in CS₂. Petersen (l.c.) gives H.C. $[Se, O^2] = 56,200$; and at. vol. This variety is a conductor of electricity, the conductivity increasing with increase of temp. and also by exposure to light. (For details, v. infra.) When heated to c. 110° this variety becomes black. The crystals are monoclinic, ratio of axes = 1.62:1:1.6 (M., l.c.; Regnault, l.c.); isomorphous with monoclinic S (R., l.c.). For S.H. v. beginning of this article.

(iii.) Grey metallic Se. Lustrous, grey, metal-like, granular solid, resembling grey pigiron; somewhat malleable (Regnault, lc.). sol. in CS. S.G. 4.4 to 4.5 (R., l.c.). Melts at 217° (H., l.c.); on cooling, the vitreous variety is chiefly formed, but if the cooling is stopped at 210° after a time there is a sudden rise to 217°, and the whole solidifies in the metallic form. All varieties of Se are sol. Se₂Cl₂ and

SeEt.

Action of heat on selenion. Any one of the crystalline varieties of Se melts at 217°; when the molten Se is allowed to cool slowly it gradually becomes less soft, a thermometer placed in the cooling substance continuously falls and shows no point of rest (Hittorf, l.c.; Regnault, l.c.); the final product is crystalline Se, but if the cooling is caused to take place rapidly the amorphous vitreous variety is formed; if the cooling is stopped at 210°, and that temperature is maintained for some time, there is a sudden rise to 217° and the Se again solidifies in the crystalline (metallic) form. When amorphous Se is heated it softens at c. 60°, and becomes gradually less viscous until at 250° it is quite liquid; if the heating is very gradual, there is a sudden evolution of heat at 96°-97° (Regnault, l.c., found temperature rise to 200°-230°), and the metallic variety is produced. Se boils at 676°-683°, forming a reddish. yellow vapour, the V.D. of which does not become constant until 600°-700° above the b.p.

Effects of light and heat on the electrical conductivity of crystalline selenion. Amorphous Se does not conduct electricity; but when heated rapidly it begins to conduct at 165°-175°, and conductivity increases until it attains a

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maximum at c. 217° (Draper a. Moss, C. N. 33, 1). Crystalline Se conducts, the conductivity increasing proportionally with increase of température to 217°, after which it decreases and reaches a minimum at 250° (Hittorf, P. 84, 214; D. a. M., l.c.). If Se is heated for a considerable time to 200° and is then allowed to cool, conductivity decreases as temperature rises and increases as temperature falls (W. Siemens, P. 159, 117). This observation seems to show that at c. 200° a variety of Se is formed which behaves like a metal, and that on cooling this form remains mixed with the ordinary crystalline variety, so that the conductivity depends on the relative quantities of the two forms, and therefore on the temperature (cf. Kalischer, W. 31, 101). The electrical conductivity of crystalline Se increases enormously when the Se is exposed to light (Sale, Pr. 21, 283; Smith, B. 6, 204; Rosse, P. M. [4] 47, 161; Forssmann, W. 2, 513). Adams and Day (Pr. 24, 163) found that the resistance of Se varied directly as the square root of the intensity of the source of light; also that light caused an electric current in Se from the less to the more illuminated part (Pr. 25, 113; cf. Bidwell, C. N. 52, 191).

Se may be sublimed and distilled in vacuo, or in an indifferent gas (Schuller, J. 1884. 1550); heated in air it is burnt to SeO₂. Se dissolves in conc. H₂SO₄, forming SeSO₂, and is ppd. again on adding water; Se is sol. in alkali solutions, forming alkali selenides; also in KCNAq, in alkaline sulphite solutions, in Se,Cl, Se, Br, and SeEt, the amorphous varieties are al. sol. CS,

Se resembles S and Te in its chemical character; it is somewhat less negative than S and more negative than Te; it shows distinct analogies with Cr, Mo, W, and U (v. Oxygen GROUP OF ELEMENTS, vol. iii. p. 705; and CHRO-MIUM GROUP OF ELEMENTS, vol. ii. p. 168).

The atomic wt. of Se has been determined (1) by analysing SeCl₄ (Berzelius, P. 8, 21 [1818]); (2) by analysing K₂SeO₄ (Mitscherlich a. Nitzsch, P. 9, 627 [1827]; (3) by oxidising Se to SeO₂, by reducing SeO₂ to Se, and by analysing BaSeO₃ (Sace, A. Ch. [3] 21, 119 [1847]); (4) by analysis of HgSe (Erdmann a. Marchand, J. pr. 55, 202 [1849]); (5) by converting Se into SeCl. (Dumas, A. Ch. [3] 55, 186 [1859]; (6) by reducing SeO₂ to Se and by analysing Ag₂SeO₃ (Pettersson a. Ekman, B. 9, 1210 [1876]; (7) by determining S.H. of Se (Regnault, A. Ch. [3] 46, 257); (8) by determining V.D. of, and analysing, SeH₂, SeO₂, Se₂Cl₂, SeCl₄, and SeBrCl₅.

The molecular wt. of Se has been found by determining the V.D. of Se at 860°, c. 1000°, and c. 1400°; the values at 1400° agree with the

formula Se.

The atom of Se is divalent in the gaseous molecule SeH2, and tetravalent in the gaseous

molecules SeCl, and SeBrCl,

Reactions and Combinations.—1. Water and ozone produce H₂SeO₄ (Mailfert, C. R. 94, 860, 1186). Water alone does not react at 160° (Cross a. Higgin, C. J. 35, 249).—2. Conc. sulphuric acid dissolves Se on warming, forming H₂SeO₃, and some H₂SeO₄, and giving off SO₂. Fuming sulphuric acid forms a greenish solution containing SeSO₂ (v. Thio-oxide, p. 441); son boiling H₂SeO₂ is formed and SO₂ evolved

(Divers a. Shimosé, B. 18, 1209).—3. Hot conc. nitric acid oxidises Se to HaSeO, with evolution of NO. Hydrochloric acid does not react with Se.-4. Se dissolves in alkali solutions, forming alkali selenides; fusion with alkalis produces the same compounds .- 5. Fusion with nitre or potassium chlorate forms K2SeO4.-6. Potassium cyanide solution dissolves Se to KCNSeAq. 7. Solutions of alkali or alkaline earth sulphites dissolve Se, forming thioselenates (Rathke a. Zschiesche, J. pr. 92, 141; Uelsmann, A. 116, 123; v. also Thioselenates, p. 434).—8. Se vapour passed over heated lead fluoride forms Se fluoride (Knox, Pr. Irish Acad. 1841. 299).-9. Se combines with oxygen to form SeO2, when heated in O or air.—10. Se combines with chlorine, bromine, and iodine at the ordinary temperature.—11. Heated tolow redness with hydrogen, H.Se is formed (Uelsmann, A. 116, 122).-12. Phosphorus combines at the ordinary temperature.-13. Se combines with arsenic and antimony, by heating the elements together .-14. Selenides of most metals are formed by heating Se with metals (v. Selenides, p. 436). Sometimes the combination is effected without heat, under the influence of light (v. Bidwell, P. M. [5] 20, 178; Moss, C. N. 33, 203).—15. Se dissolves in molten sulphur, but no definite compounds are formed in this way.—16. With sulphuric anhydride Se combines to form SeSO, (v. Thio-oxide, p. 441).

Selenion, acids of. Scienhydric acid H.Se, vol. ii. p. 724; selenic acid H_.SeO_., p. 435; selenious acid H_.SeO_., p. 445; Selenocyanhydric acid HCNSe, vol. ii. p. 348; also Selenocyanic acid, this vol. p. 443; salts of thioselenic acids, H₂SeSO₂ and H₂SeS₂O₆, are known, p. 434; and salts of selenosamic acids, Se(NH,)OM1 and 2SeO(NH₂).OH.OM¹, have been isolated, p. 446.

Selenion, antimonide of; v. vol. i. p. 283,

ANTIMONY, Combinations of, No. 6. Selenion, arsenides and arsenosulphides of;

v. vol. i. p. 303, Arsenic, Combinations of, No. 7. Selenion, boride of. A yellow-grey solid is formed by strongly heating amorphous B in a stream of H₂Se; the solid is decomposed by water with rapid evolution of H₂Se (Sabatier, C. R. 112, 865). As S. gives the formula B₂S₃ to the boron sulphide formed by passing H2S over B, it is probable that the Se compound has the composition B2Se3.

Selenion, bromides of. Two compounds of Se and Br are known, Se, Br, and SeBr,; both are formed by the direct union of the elements. Both are decomposed by heat, but they are more stable towards heat than the corresponding S compounds (Evans a. Ramsay, C. J. 45, 62). As the V.D. of neither has been determined, the formulæ are not necessarily molecular; but from the analogy of Se₂Cl₂ and SeCl₄, which formulæ represent the compositions of gaseous molecules of these chlorides, it is probable that the mol. wts. of the bromides correspond with the formulæ Se₂Br₂ and SeBr₄.

SELENIOUS BROMIDE Se, Br. (Selenion mono-bromide.) Formula probably molecular, from analogy of Se, Cl.. To 15-9 parts powdered Se covered with three times its weight of dry CS, 16 parts Br are gradually added; after a time the CS, is distilled off; and Se,Br, remains as a thin oily liquid, having a dark reddish-brown colour (appearing black in a thickish layer), and an unpleasant smell resembling that of S₂Cl₂ (Serullas, A. Ch. [2] 35, 349; Schneider, P. 128, 827). A thin layer of Se₂Br₂ transmits rose-red light; the absorption spectrum has been examined by Gernez (C. R. 74, 1190). S.G. of Se₂Br₂ = 3·604 at 15°. Bris given off on heating, then a little SeBr₄, and at 225°-230° most of the Se₂Br₂ distils over, and finally some Se is obtained (Schneider, l.c.). Se₂Br₂ dissolves in all proportions in CS₂, less easily in CHCl₃, EtBr, and EtI. It sinks in water, and then decomposes to H₂SeO₃Aq, HBrAq, and Se. Absolute alcohol forms SeBr₄ (which remains dissolved) and Se. Se₂Br₂ dissolves Se (c. 22 p.c.), on adding CS₂ the Se separates. Br is absorbed by Se₂Br₂, with formation of SeBr₄. A solution of Se₂Br₂ in CS₂ shaken with dry AgCN forms a little Se cyanide (Schneider, P. 123, 634).

SELENIC BROMIDE SeBr. (Selenion tetrabromide.) Formula probably molecular, from analogy of SeCl₄. To a solution of 1 part Se₂Br₂ in CS₂ 1½ parts Br are gradually added, the SeBr, which separates is allowed to stand in contact with the CS_2 for some days, whereby it becomes crystalline, and the CS_2 is distilled off; the compound obstinately retains some CS2. By heating to 75°-80° Br is given off, and the SeBr₄, containing Se₂Br₂, sublimes in black, lustrous, six-sided leaflets, mixed with a little pure SeBr, in dark-orange crystals, and Se2Br2 remains (Schneider, P. 129, 450). SeBr, is also obtained by dissolving 1 part powdered Se in 10-12 parts Br, pressing the crystalline mass between filter-paper, and removing excess of Br by gently warming. SeBr₄ dissolves in CS₂, CHCl₃, and EtBr; it dissolves with decomposition in alcohol, also in HClAq. SeBr, is very hygroscopic; in moist air it gives Br and Se,Br,; in much water it forms H,SeO,Aq and HBrAq. Combines with SO, to form SeCl. SO, (v. Selenion thio-oxychloride, p. 441). Forms compounds with KBr and NH, Br, K2SeBr, and (NH₁)₂SeBr₆ (v. Muthmann a. Schäfer, B. 26, 1008).

Selenion, bromochlorides of. Two compounds of Se, Br, and Cl were obtained by Evans and Ramsay (C. J. 45, 68) by the interactions of Se₂Cl₂ and Br, and Se₂Br₂ and Cl. The bromochlorides thus formed are SeBrCl₃ and SeBr₃Cl.

SELENION TRIBROMOCHLORIDE SeBr₃Cl. Orange, hygroscopic crystals formed by adding 2·1 parts Br, in CS₂, to 1 part Se₂Cl₂, crystallising from CS₂, and gently warming in a current of air to remove adhering CS₂ (Se₂Cl₂ + 3Br₂ = 2SeBr₃Cl). Decomposes when heated, giving off Br; as V.D. at 200° was c. half that calculated for SeBr₃Cl, dissociation is probably complete at that temperature.

SELENION BROMOTRICHLORIDE SeBrCl₂. Mol. w. 264·66. V.D. at 179° and 754 mm. pressure = 181·5; dissociation begins at c. 190°-200°. Yellow-brown crystals; obtained by passing Cl into Se₂Br₂ in CS₂, washing the pp. with CS₂, and drying in a current of warm air. Gives off Br when heated above c. 190°, and yields sublimate of SeBr₄.

Selenion, carbide of, v. Carbon selenide, vol. i. p. 693.

Selenion, chlorides of. Two chlorides are formed by passing Cl over heated Se; using an excess of Se the product is Se₂Cl₂, and with an excess of Cl the product is SeCl₄. Attempts to form SeCl₂, corresponding with SCl₂, have been unsuccessful.

Selenious chloride Se₂Cl₂. (Selenion monochloride. Selenion selenochloride.) Mol. w. 228·34. V.D. 110 at 200° – 250° (Evans a. Ramsay, C. J. 45, 62; Chabrié, Bl. [3] 2, 803). S.G. 2·906 at $17\cdot5^{\circ}$ (Divers a. Shimosé, C. J. 45, 198). Boiling begins at c. 145° , but no definite b.p. can be given (E. a. R., l.c.). H.F. [Se²,Cl²] = 22,150 (Th. 2, 405).

Formation.—1. By passing Cl over excess of heated Se.—2. By heating SeCl₄ with Se at 100° (Berzelius, P. 9, 225).—3. By passing HCl into a solution of Se in fuming H₂SO₄ (Divers a. Shimosé, C. J. 45, 194, 198).

Preparation.—HCl gas is passed into a solution of Se in fuming H_2SO_4 till a deep-red liquid is deposited and the mother-liquor is yellowish-brown; the mother-liquor is then poured off, the red liquid is dissolved in fresh fuming H_2SO_4 and HCl is passed in; this process is repeated again; finally, the red liquid is digested, in a closed vessel, with dry KCl (to remove H_2SO_4) and decanted (D. a. S., l.c.). The reactions are probably (1) $2SeSO_3 + 2HCl = SeSO_3.SeCl_2 + H_2SO_3$; (2) $SeSO_3.SeCl_2 + HCl = Se_2Cl_2 + SO_2.Cl.OH$ (D. a. S.).

Properties, Reactions, and Combinations.—Se₂Cl₂ is a heavy, deep-red liquid, smelling something like S₂Cl₂; S.G. 2.906 at 17.5°; somewhat volatile at ordinary temperatures, giving off orange-coloured vapour at 100°, beginning to boil at c. 145°, but exhibiting no fixed b.p. (E. a. R., l.c.). V.D. determined at c. 200° gave results agreeing with Se₂Cl₂. For absorption spectrum of vapour v. Gernez, C. R. 74, 1190. Se₂Cl₂ is easily sol. CS₂, CHCl₃, C₆H₆, and CCl₄. It is slowly decomposed by H₂O, EtOH, and Et₂O; a solution in CS₂ shaken with water gives H₂SeO₃Aq, HClAq, and Se. Warm Se₂Cl₂ dissolves Se freely; on cooling the Se is partly ppd. in the metal-like form (Rathke, A. 152, 181). Se₂Cl₂ is decomposed by shaking with Hg or Ag (D. a. S., l.c.); with P it gives PCl₃ and Se (Baudrimont, A. Ch. [4] 2, 5). Se₂Cl₂ combines with PCl₃ to form a yellow substance (B., l.c.).

SELENIC CHLORIDE SeCl₄. (Selenion tetrachloride.) Mol. w. 220·28. V.D. at $180^{\circ} = 110$ (Evans a. Ramsay, C. J. 45, 65). H.F. [Se,Cl⁴] = 46,160 (Th. 2, 405).

Formation.—1. By heating Se, or Se₂Cl₂, in excess of Cl.—2. By heating SeO₂ with PCl₃ (v. Preparation).—3. By the action of SOCl₂ on SeOCl₂.

Preparation.—To 13 parts PCl₃ in a long-necked flask 7 parts SeO₂ are added attle by little, the reaction being allowed to cease after each addition before more SeO₂ is added (SeO₂+PCl₃=SeOCl₂+POCl₃); the mass becomes liquid and then suddenly solidifies (3SeOCl₂+2POCl₃=3SeCl₄+P₂O₃); excess of POCl₃ is now removed by heating the flask while a rapid stream of dry CO₂ is passed through it; a wide tube is then fitted by a cork into the neck of the flask, and the flask is placed in a tray containing hot sand (if the flask is

allowed to cool the condensing P2Os may cause

it to crack) (Michaelis, J. Z. 6, 79)

Properties, Reactions, and Combinations. A slightly yellow crystalline solid; volatilises when heated, without melting, in yellow vapours. Evans a. Ramsay (C. J. 45, 65) found V.D. at 180° to be 110, agreeing with the formula SeCl4; dissociation to Se₂Cl₂ and Cl began at c. 210° and at 350° c. 66 p.c. was decomposed. Clausnizer (B. 11, 2007) got the value 57 for V.D. at 218°, at which temperature E. a. R.'s value was c. 86; at 368° Chabrié (Bl. [3] 2, 803) obtained the number 57, which agrees fairly with the results of E. a. R. SeCl, is nearly insol. CS,. In moist air SeOCl, and HCl are produced (Weber, P. 108, 615); water forms H₂SeO₃Aq and HClAq. H₂S produces HCl and Se sulphide; SO₂ has no action. NH₃Aq produces Se, N, and H at ordinary temperatures, and N selenide at a low temperature (Verneuil, Bl. [2] 38, 548). Dry NH₃ combines with SeCl₄ at -20°, forming a very unstable compound, probably SeCl, 4NH, (V., l.c.). With PCl, the compound SeCl, 2PCl, is formed (Baudrimont, A. Ch. [4] 2, 5).

Selenion, chlorobromides of, v. Selenion

BROMOCHLORIDES, p. 439.

Selenion, cyanide of; v. Cyanogen selenide,

vol. ii. p. 358.

Selenion, fluoride of. No definite compound of Se and F has been isolated. According to Knox (Pr. Irish Acad. 1841. 299) a fluoride of Se sublimes when Se vapour is passed over

molten PbF₂.
Selenion, haloid compounds of. Se combines with the halogens to form two types of compounds, Se2X2 and SeX4 (the fluorides have not been examined); the formulæ are probably molecular, as the V.D.s of Se₂Cl₂ and SeCl₄ have been determined. The compounds Se₂X₃, where X = Cl or Br, are liquids at ordinary temperatures; the other compounds are solids. All the compounds are fairly easily decomposed by heat, but less easily than the corresponding S haloid compounds.

Selenion, hydride of; v. Hydrogen selenide,

vol. ii. p. 724.

Selenion, hydroxides of. The hydroxides of Se, SeO(OH)₂, and SeO₂(OH)₂ are acids; v. Selenious acid, p. 445, and Selenic acid, p. 435.

Selenion, iodides of. Two compounds of Se and I are known, Se, I, and SeI, corresponding

with the bromides and chlorides.

SELENIOUS IODIDE Se₂I₂. (Selenion monoiodide.) Formula probably molecular, from analogy of Se₂Cl₂. Formed by heating the elements together in the ratio Se:I(1:1:61); also by heating Se₂Br₂ and EtI at 100° in a sealed tube (Schneider, P. 129, 627). Crystalline, steel-grey, somewhat metal-like solid; melting to a dark-brown liquid at 68°-70°. Decomposed to Se and I by continued heating at 100°; also by solvents of I. Water forms H₂SeO₂Aq, HIAq, and Se.

SELENIC IODIDE Sel, (Selenion tetra-iodide.) Formula probably molecular, from analogy of SeCl. Formed by heating together Se and I in the ratio Se:41(1:6:44); also by mixing SeBr₄ and EtI in the ratio SeBr₄:4EtI, the SeI₄ separates after a little. Best prepared by adding HIAq to conc. H₂SeO₂Aq as long as a pp. is produced; the pp. is washed, pressed, and dried over H2SO4 (Schneider, l.c.). A dark-grey solid, melting at 75°-80° to a blackish-brown liquid. Gives up all I when heated to c. 100°, or treated with solvents of I; slowly decomposed by water. Attempts to form an iodochloride of selenion were unsuccessful (v. Evans a. Ramsay, C. J. 45, 70).

Selenion, nitride of; v. Nitrogen selenide,

vol. iii. p. 570.

Selenion, oxides of. Only one oxide of Se has been isolated with certainty; this oxide is SeO. Berzelius thought that an oxide SeO was formed, along with SeO2, by heating Se in air or O, by heating Se and SeO, and by the partial oxidation of Se sulphide by HNO. According to Chabrié (A. Ch. [6] 20, 202) Se takes up O approximately corresponding with that required to form SeO, when heated in air to 180°. Attempts to prepare SeO, have failed. By passing gaseous SeO2 and O over hot spongy Pt, von Gerichten (A. 168, 214) obtained a white sublimate that dissolved in water with a hissing sound, forming H,SeO, but Cameron a. Macallan (Pr. 46, 32) failed to confirm this result; C. a. M. (l.c.) did not get SeO, by passing O over hot Pt sponge mixed with Se, nor by the action of ozone on SeO2, nor by heating selenate of Sb, Bi, Fe, Pb, Hg, or Ag. By heating dry H₂SeO₄ with P₂O₅ in a closed flask they got a trace of a crystalline solid, analyses of which pointed to the composition SeO₃. A solution of Se in H2SeO4 perhaps contains Se2O3 (v. Selenic ACID, Reactions, No. 6, p. 436).

SELENION DIOXIDE SeO₂. (Selenious oxide. Selenious anhydride.) Mol. w. not determined. S.G. 3.9538 at 15° (Cameron a. Macallan, Pr. 46, 27; Clausnizer, A. 196, 265). [Se,O²] = 57,250 from amorphous Se; 55,820 from black crystalline Se; 56,200 from red crystalline Se (Petersen, Z. P. C. 8, 601). Melts at 340° in a sealed tube; at ordinary pressure volatilises without melting, slowly at 250°, rapidly at 280° (Chabrié, A. Ch. [6] 20, 202; C. a. M., l.c.). For absorption-spectrum of SeO₂ vapour v.

Gernez (C. R. 74, 803).

Prevaration .- 1. Se is placed in the lower part of a V-tube with one limb longer than the other, and the longer limb drawn to a fine opening; the shorter limb is connected with a gasholder, from which a stream of dry O is passed into the V-tube; the tube is heated till the Se burns; the SeO₂ condenses in the longer limb of the tube. -2. Se is dissolved in conc. HNO, Aq, the solution is evaporated to dryness, best in a retort (solidification occurs suddenly, with evolution of heat); the dry H, SeO, is heated in the retort till sublimation begins, then let cool, and dissolved in water; the solution is ppd. by BaOAq; the solution, after filtering, is evaporated to dryness, and the residue is heated, when SeO₂ sublimes; traces of H₂SeO₄ and H₂SO₄ are thus removed, excess of BaO remains in the residue (Thomsen, B. 2, 598; cf. Wohlwill, A. 114, 176; Fischer, P. 67, 412).

Properties and Reactions .- Long, white, lustrous, four-sided needles (for m.p. &c. v. beginning of this article). 1. Slowly withdraws water from moist air, forming H2SeO2; dissolves in water, forming same acid, with disappearance of

heat, $[SeO^2,Aq] = -920$ (Th. 2, 405).—2. Sl. sol. boiling acetic axhydride, crystallising unchanged; heated to 180°-200° oxidation of the Ac.O occurs (to CO₂, &c.) and Se separates (Hinsberg, A. 260, 40).—3. Dissolves in aqueous acetic acid, H.SeO, separating on cooling (H., l.c.).-4. Dissolves in alcohol of 96 p.c., and is re-obtained on evaporation over H2SO, (H., L.c.). 5. Reacts with phosphoric chloride to give SeCl, P₂O₃, and POCl₃; heated with phosphoryl chloride gives SeOCl₂ and P₂O₃ (Michaelis, J. Z. 6, 79). -6. Heated with sodium chloride yields SeOCl, and Na SeO, (Cameron a. Macallan, Pr. 46, 35). -7. Heated with selenion tetrachloride in a closed tube produces SeOCl₂; with SeBr, probably forms SeOBr₂ (Schneider, P. 129, 450). 8. Decomposed by dry ammonia to H₂O, Se, and N; when NH, is passed into SeO₂ in absolute alcohol, SeO(NH₂).ONH₄ is formed (v. Selenos-AMIC ACIDS, SALTS OF, p. 446). -9. Heated with dry HCN and Ac₂O, Se cyanide is probably formed (Hinsberg, A. 260, 40).—10. Hydriodic acid reduces SeO₂ to Se and H₂O, even at 10°.—11. SeO₂ and sulphur dioxide do not react at any temperature (Schulze, J. pr. [2] 32, 390).-12. The compounds K₂SeBr₆ and (NH₄)₂SeBr₆ are produced by dissolving SeO₂ in hydrobromic acid and adding potassium or ammonium bromide (v. Muthmann a. Schäfer, B. 26, 1008)

Combinations.—1. According to Ditte (A. Ch. [5] 10, 82), SeO₂ combines with dry hydrogen chloride to form two compounds, one of which, SeO₂.2HCl, is liquid, and the other, SeO₂.4HCl, is solid, at the ordinary temperature. hydrogen bromide SeO, 4HBr is formed, and when this is kept very cold and HBr is passed over it, D. says that SeO, 5HBr is produced. Hydrogen fluoride is said also to combine with SeO2, but the product has not been examined. - 2. With alkali halides the compounds MX.2SeO, 2H,O are formed (Muthmann a. Schäfer, B. 26, 1008).—3. Hinsberg (A. 260, 40) described a compound with alcohol, SeO₂.C₂H₆O, obtained by evaporating SeO2 in absolute alcohol over CaCl2.-4. Large white crystals of the compound SeO₂.SO₃ are obtained by warming SeO₂ and pure sulphuric anhydride to 100°, distilling off excess of SO₃ at 60°-70°, and allowing to cool. At 100°

the compound gives up SO₃ (Weber, B. 19, 3185). SELENION THIO-OXIDE SeSO₃. (Selenion sulphoxide. Selenion-sulphur trioxide.) Mol. w. not determined. Magnus (P. 10, 491) noticed that Se dissolved in fuming H₂SO₄ to produce a green solution; Weber (P. 156, 531) separated and analysed the green compound, and gave it the formula SeSO₃. Divers a. Shimosé (C. J. 45, 201) further examined this substance. Powdered Se is placed in liquid SO₃; heat is given off, and a dark-green liquid is formed; after a time the whole solidifies; after warming to 30°-40° for a few minutes the liquid part is drained off, and the adhering SO₃ is removed by the use of a Sprengel pump (for details v. D. a. S. on TeSO₃, C. J. 43, 324). SeSO₃ is a green solid, which may be heated to c. 35° without change; decomposition begins at c. 40°, and at c. 120° SO₂ and O are rapidly evolved and Se separates; by heating slowly in vacuo it changes to a yellow powder, which D. a. S. think to be a modification of the thio-oxide. SeSO₃ is sol. conc. H₂SO₄; on adding much water Se is ppd.,

and H₂SeO₂ and H₂SO₂ go into solution. HCl reacts with SeSO₃ in H₂SO₄ to form Se₂Cl₂ and SO₂.Cl.OH (v. Selenious chloride, Preparation, p. 439). Schulz-Sellack (B. 4, 113) observed that the vapours of SO₂ react with Se to form a yellow powder.

Selenion, oyacids of, and their salts. The oxyacids of So are selenious acid H₂SeO₃ (v. p. 445) and selenic acid H₂SeO₄ (v. p. 435); salts of the thioselenic acids H₂SeSO₃ and H₂SeSO₄ have been isolated (v. p. 434); and salts of the selenosamic acids SeO(NH₂).OH and 2SeO(NH₂).(OH)₂ are known (v. p. 439).

Selenion, oxybromide of. By melting together SeO₂ and SeBr₄ Schneider (P. 129, 450) obtained needles which were probably SeOBr₂, corresponding with SeOCl₂ formed in a similar

Selenion, oxychlorides of. The oxychloride of Se, ScOCl₂, corresponds with sulphuryl chloride SOCl₂; an oxychloride containing Se and S, and corresponding with S₂O₂Cl₄, has also been isolated.

SELENYL CHLORIDE SeOCl₂. (Selenion oxychloride.) Mol. w. not determined.

Preparation.—1. Equivalent weights of SeCl₄ and SeO₂ are heated in a sealed tube to 150°, and the product is purified by repeated distillation (Weber, P. 118, 615).—2. SeO₂ and PCl₃ are brought together in the ratio SeO₂:PCl₃; SeOOl₂ and POCl₃ are produced, but the heat evolved suffices to cause a secondary reaction, which yields SeCl₄ and P₂O₃, so that very little SeOCl₂ can be obtained by this method (Michaelis, J. Z. 6, 79; cf. Selenio chloride, Preparation, p. 439).—3. By distilling SeO₂ and NaCl in the ratio SeO₂:NaCl (Cameron a. Macallan, Pr. 46, 35).

Properties.—A yellowish liquid, fuming in the air, boiling at 179.5°, and solidifying below 0° to colourless crystals, which melt at 10° (M., l.c.).

Reactions and Combinations (M., l.c.).—
1. Water decomposes SeOCl₂, forming H₂SeO₂Aq and HOlAq.—2. Ammonia produces SeO₂, Se, N, NH₄Cl.—3. Sulphuryl chloride produces SeOl₄ and SO₂.—4. Phosphoryl chloride forms SeOl₄ and P₂O₅ (v. Preparation of selenic chloride, p. 439).—5. With phosphorous chloride POOl₅, SeOl₄ and Se₂Ol₂ are produced.—6. SeOCl₂ combines with stannic chloride to form SSeOCl₂-SnOl₄; also with titanic chloride and antimonic chloride to form 2SeOCl₂-TiOl₄ and 2SeOCl₂-SbCl₅ (Weber, P. 125, 325).

SELENION, THIO-OXYCHLORIDE OF, SeSO,Cl., (Pyrothioselenyl chloride. Selenion sulphooxytetrachloride.) Mol. w. not determined.

oxytetrachloride.) Mol. w. not determined. Formation.—1. By the interaction of SO₂ and SeCl₁ (H. Rose, P. 44, 315).—2. By dissolving SeCl₂ in Nordhausen sulphuric acid (Clausnizer, B. 11, 2007).—3. By heating SO₂Cl₂ and SeOCl₂ in a sealed tube to 170°–180° (Č., l.c.).—4. By the interaction of SO₂OH.Cl and SeO₂, SeOCl₂, or SeCl₄ (C., l.c.).

Preparation.—SeCl, and SO.OH.Cl are warmed together in the ratio SeCl,:2(SO₂.OH.Cl) till the SeCl, is quite dissolved:

 $SeCl_1 + SO_2OH.Cl = SO_2Cl.OSeCl_1 + Cl.$ (Excess of $SO_2OH.Cl$ is needed to hasten the reaction and to serve as a solvent.) The reddish solution solidifies on cooling; the white needles are separated by pressing between porous plates, and

dried over H2SO4.

Properties and Reactions. - Small white needles; unchanged in vacuo or on gentle warming; deliquesces in moist air; melts at 165° and boils at 183°; V.D. at 209° = 48.5, for $SeSO_3Cl_4 = 150$; calculated dissociation has therefore occurred, perhaps thus: $2SeSO_3Cl_4 = 2SO_3 + Se_2Cl_2 + 3Cl_2$ (C., l.c.). Decomposed by water, at once, to H2SO, Aq, H2SeO, Aq, and HClAq.

Selenion, phosphides of; v. PHOSPHORUS

selenides, p. 144.

Selenion, sulphides of. The only compound of Se and S that has been isolated is SeS. Divers a. Shimidzu (C. J. 47, 446) say that this is the only compound of the two elements that exists, but Rathke (B. 18, 1534) considers that others exist, although none but SeS has been obtained in a state of purity. Se and S may be melted together in all proportions, but no compound is produced (Bettendorff a. von Rath, P. 139, 329; Rathke, P. 141, 590). Isomorphous mixtures of the two elements crystallise from solutions of Se and S in CS₂ (B. a. von R., l.c.). The pp. formed by passing H2S into H2SeO3 contains Se and S in the ratio SeS, but it is a mixture of SeS with S (H. Rose, P. 107, 186; D. a. S., l.c.; Rathke, B. 18, 1534). The pp. containing Se and S in the ratio Se₂S obtained by passing H₂Se into H₂SO₃ is merely a mixture (D. a. S., l.c.).

Selenion monosulphide SeS. Mol. w. not determined. Very dilute H₂SeO₃Aq, cooled to 0°-5°, is saturated with H₂S, which is first passed through a flask filled with pieces of ice; the yellow pp. which forms is washed, dried in vacuo, moistened with CS2, and allowed to stand for some days till the mass becomes crystalline, the CS2 withdrawing S; CS2 is added to the crystals, after a few minutes the CS₂ is decanted, and the residue is washed with C.H. and then with alcohol. SeS forms small, orangeyellow, lustrous tablets; S.G. 3.056 at 0°, 3.035 at 52°; S.H. 1274. On heating it melts, gives off S vapour, and then vapour of S and Se. Sol. CS₂, but not crystallisable therefrom by evaporation; insol. water and ether (Ditte, C. R. 73, 625, 660).

Selenion, sulphoxide of; v. Selenion thio-

oxide, under Selenion oxides, p. 441.

Selenion, sulphoxychloride of; v. Selenion thio-oxychloride, under Selenion oxychlorides; p. 441.

Selenion, thio-oxide of; v. Selenion oxides,

p. 441.

Selenion, thio-oxychloride of; v. Selenion ichlorides, p. 441. M. M. P. M. oxychlorides, p. 441. M. M. P. M. SELENION, ORGANIC COMPOUNDS OF.

Methyl selenide or Selenion methide Me.Sc. Mol. w. 109. (58°). Formed by distilling a solution of barium methyl sulphate with K2Se (Wöhler a. Dean, A. 97, 5) or of KMeSO, with Na₂Se (C. L. Jackson, A. 179, 1). Heavy oil with nasty smell. Burns with bluish flame. PtCl, forms (Me₂Se)₂PtCl₄, crystallising from alcohol in yellow plates. Conc. HNO₂ forms (Me SeO)HNO₂ [91°], whence HClAq produces Me SeCl₂ [60°], while HBrAq gives Me₂SeBr₂ [82°] and KIAq ppts. Me₂SeI₂.

Mathylocodida Me SeI

Methylo-iodide Me,SeI. Tri-methyl-monium iodide. Yields (Me,SeCl),PtCl, selenonium iodide.

crystallising from water in dark-red octahedra (Jackson, B. 7, 1277).

Di-methyl diselenide Me, Se. Formed from KMeSO, and K2Se, (W. a. D.; Rathke, A. 152, 211). Heavy reddish-yellow oil, with unpleasant odour, sol. HNO, Aq.

Methane selinic acid CH₃.SeO₂H. [122°]. Formed by oxidising Me₂Se₂ with HNO₃ (W. a. D.). Deliquescent crystals, v. sol. alcohol. Gives with HCl crystalline CH.SeO2Cl.-AgA': prisms.

Ethyl selenide Et₂Se. (108°). (Rathke; Pieverling, A. 185, 331). Oil, without unpleasant odour. HCl added to its solution in dilute HNO₃ ppts. Et₂SeCl₂, whence NH₃Aq forms (Et₂Se)₂OCl₂ (Joy, A. 86, 85).

Ethyl iodide Et. Sel. Crystalline. Converted by moist Ag₂O into a strongly alkaline base which forms crystalline salts. - (Et, SeCl), ZnCl

thin plates.—(Ét₃SeCl)₂PtCl₄.

Di-ethyl diselenide Et₂Se₂. (186°). Liquid with disgusting odour. HCl added to its solution in dilute HNO, gives monoclinic crystals of C2H8SeO2Cl.

Ethane selinic acid Et.SO.H. Yields with

HCl monoclinic crystals of C₂H_sSeO₂Cl.

Ethyl selenhydrate EtSeH. Selenionmercaptan. Formed from KSeH and Ca(EtSO4)2 (Siemens, A. 61, 360). Heavy oil, boiling much below 100° and having an unpleasant odour. Reacts with HgO.

Ethyl selenite SeO(OEt)₂. (184°). S.G. 1.49. Formed from Ag₂SeO₃ and EtI, and 16 1.49. got also by heating SeOCl2 with NaOEt and dry ether at 180° (Michaelis, A. 241, 158). Thick liquid, miscible with water and alcohol. The chloride SeO(OEt)Cl [10°] (175°), formed from SeOCl₂ and alcohol, crystallises in needles.

Selenide of carbon CSe. Formed in small quantity by passing a mixture of CCl₄ and H.Se through a red-hot tube (Rathke, A. 152, 199). Pungent yellow liquid. Converted by alcoholic potash into EtO.CSe2K, crystallising in yellow needles, v. sol. water and alcohol.

Se(CH2.CO2H)2. Selenido - di - acetic acid Formed from ammonium chloro-acetate and alcoholic (NH₄)₂Se (Ulrich, B. 8, 773). Monotables. - *(NH₄),A": insol. alcohol - *CuA": bluish-green pp.

Amide.—Formed from ClCH2.CONH2 and alcoholic (NH₄)₂Se. Prisms, m. sol. cold water.

Di-phenyl selenide SePh₂. S.G. 20 1.450. V.D. 8.17 (calc. 8.09). Formed, together with selenophenol and Se₂Ph₃.C₆H₄Cl (240° 250° in vacuo), from SeCl, and benzene in presence of AlCl, (Chabrié, C. R. 109, 183; Bl. [2] 50, 183; [3] 2, 788). Oil. Boils at 227° under a few cm. pressure. Bromine forms Se(C, H, Br), [112°], crystallising from alcohol in colourless hexagonal plates. H2O2, in presence of HCl and a current of air, forms Se(C,H,Cl).C,H,OH [145°] converted by HNO, into a crystalline product

Phenyl selenhydrate C,H,SeH. phenol. [60°]. Formed from benzene, SeCl., and AlCl, (Chabrié, C. R. 109, 183; Bl. [3] 2,

788). Sol. alcohol, insol. water.

Tetra-methyl-di-amido-di phenyl-selenide Se(C_cH,NMe₂): [124°]. Formed from SeOCl₂ and di-methyl-aniline in ether (Godchaux, B. 24, 765). Yellowish needles, m. sol. cold alcohol and ether.—B"H,SO. [55°]. Colourless needles, v. e. sol. water.—B"2C_eH₂(NO₂)₃OH. [135°]. Yellow plates.

Tetra - ethyl - di - amido - di - phenyl - selenide $Se(C_{e}H_{\downarrow}NEt_{2})_{2}$. Formed from SeOCl2 [83°]. and PhNEt, in ether. Silky needles, v. e. sol. hot ether and alcohol.—B"H₂Cl₂. [73°].— B"2C,H2(NO2),OH. [135°]. Yellow plates and

needles.

Di-phenyl-selenone SeOPh₂. (230° at 65 mm.). S.G. 19:61:48. Formed, together with the compound PhSeO(C₆H₄Cl) [94°], from SeOCl₂, benzene, and Amber-yellow liquid. Not attacked by cold HNO₂. Bromine-water yields SeO(C₂H₄Br)₂ [120°]. H₂O₂, HCl, and a current of air form SeO(C₂H₄Cl)₂ [159°].

Phenyl seleno-carbimide Ph.NC.Se. Formed by shaking Na2Se with the chloride of phenylcarbamine, PhNCCl2, and ether for a day, filtering, distilling with steam, and drying the oil in vacuo over H₂80₄ (Stolte, B. 19, 2350). Pale yellowish-red oil, v. sol. alcohol and ether.

Phenyl-seleno-urea NHPh.CSe.NH₂. [182°]. Formed by passing NH, into Ph.NC.Se dissolved in ether, and also by the action of H₂Se on phenyl-cyanamide in benzene (Stolte, B. 19, 1579, 2350). Crystals (from alcohol).

Di-phenyl-seleno-urea CSe(NHPh)₂. [186°]. Formed from aniline and phenyl seleno-carbimide (Stolte, B. 19, 2351). Crystals (from alcohol), sl. sol. ether.

Selenaldine C₆H₁₃NSe₂. Formed by passing SeH₂ into an aqueous solution of aldehydeammonia in absence of air (Wöhler a. Liebig, A. 61, 11). Colourless crystals, with slight odour, sl. sol. water, v. sol. alcohol and ether. Its solutions are turned brown by air.

Amido-selenazole C3H4N2Se i.e. Se $\overline{C(NH_2)} \gg N$. Selenazylamine. [121°]. CH:CH Formed by warming seleno-urea with di-chlorodi-ethyl oxide and aqueous NaOEt (G. Hofmann, A. 250, 308). White needles, v. sol. alcohol and ether, m. sol. benzene and water. Decomposed by prolonged boiling with water.—B'2H,PtCl6: orange-red pp.

Acetyl derivative C.H.AcN.Se. [c. 210°].

Needles.

Oxy-amido-selenazole Se $.C(NH_2)$ N. Selenohydantoin. [190°]. Formed by boiling selenourea with chloro-acetic acid and alcohol (G. Hofmann, A. 250, 312). Thick crystals, m. sol. hot water and alcohol, insol. ether. -B'HCl: needles.

CH:C(OH) > N. Di-oxy-selenasole [147°]. Formed by boiling oxy-amido-selenazole with NaOHAq (Hofmann). Prisms, v. sol. alcohol, m. sol. water.

Se $C(NH_2) \gg N$. Amido-methyl-selenazole CH:CMe [80°]. Formed from seleno-urea and chloroacetone (G. Hofmann, A. 250, 305). Needles, v. sol. alcohol and ether. Alkaline in reaction. -B'HCl: prisms (from water).-B'₂H₂PtCl₆: orange-red crystalline pp.

Acetyl derivative C.H.AcN.Se. [122°]. Needles (from benzene), sl. sol. water.

Amido-methyl-selenazole carboxylic acid Se.C(NH₂) N. [195°]. Formed by heat-CO,H.C :CMe ing seleno-urea with chloro-acetoacetic ether (G. Hofmann, A. 250, 309). Needles, m. sol. water, v. sol. alcohol and ether.—HA'HCl: plates, v. e. sol. water.

Acetyl derivative C, H, AcN, SeO, [220°].

Needles, almost insol. hot water.

Amido-phenyl-selenazole C.H.N.Se Se $C(NH_2) \gg N$. [132°]. Formed from bromo-CH:CPh acetophenone and seleno-urea (G. Hofmann, A. 250, 307). Needles or prisms, insol. water, sol. alcohol and acids.—B'HBr: yellowish plates, sl. sol. cold water.

Di-phenyl-selenazole C₁₅H₁₁NSe i.e.

Se .CPh N. [99°]. Formed from ω-bromoacetophenone and seleno-benzamide (G. Hofmann, A. 250, 317). Plates, v. sol. alcohol and ether. Its salts are decomposed by water .--B'2H2PtCl6: pale-yellow pp.

Phenyl-methyl-selenazole $C_{10}H_{9}NSe$ Se .CPh N. (283°) at 737 mm. Formed from PhCSe.NH₂ [126°] and chloro-acetone (G. Hofmann, A. 250, 316). Yellow oil.—B'2H2PtCla:

pale-yellow needles.

Phenyl-methyl-selenazole carboxylic acid C, H, SeNO, i.e. [207°]. Formed by saponifying its ether, which is got from seleno-benzamide and chloro-acetoacetic ether (Hofmann, A. 250, 318). Small needles (from benzene), m. sol. alcohol.—AgA'.

Ethyl ether EtA'. [124°]. Plates. Seleno-benzoic acid. Amide C,H₃.CSe.NH₂. [126°]. Formed by passing H2Se into an alcoholic solution of benzonitrile made slightly alkaline with NH₃ (Dechend, B. 7, 1273; G. Hofmann, A. 250, 314). Golden needles (from water or Reacts with a-chloro-ketones, forming ether). selenazoles; e.g. X.CO.CHCl.X + CPh(SeH):NH= H₂O + $\overset{\text{Se}}{\text{CX:CX}} > \text{NHCl}$.

Seleno-benzoic aldehyde v. Benzylidene

Selenocyanic acid HSeCN. The K salt is got by fusing K, FeCy, (3 pts.) with Se (1 pt.) (Crookes, A. 78, 177; cf. Berzelius, S. J. 31, 60). The free acid is obtained, in solution, by decomposing the lead salt by H2S. It is decomposed by acids into Se and HCy. FeCl, does not give a red colour, but ppts. Se. Iodine added to a solution of KSeCN ppts. KC₂N₂Se₄. Air containing a little chlorine passed over a 10 p.c. solution of KSeCN forms a red crystalline pp. KSe₄C₂N₃ aq changing to yellow Se₃Cy₂, which separates from chloroform in yellow plates, and is decomposed by water into HCy, SeO₂, and Se and yields in vacuo at 108° a crystalline sublimate of Se,Cy2, decomposed by hot water into HCy, SeO2, and Se (Verneuil, C. R. 103, 144). Alcohol converts KSe C, N, into Se and KSe, C, N, which forms brown crystals, sol. alcohol.

Salts.—KA'. Formed as above, and also by dissolving Se in aqueous KCy (Wöhler; Schiellerup, A. 109, 125; G. Hofmann, A. 250, Very deliquescent needles, alkaline to litmus.—NH,A' (Cameron a. Davy, C. N. 44, 68).—KA'HgCl₂: white pp.—KA'HgBr₂.— KA'HgL₂.—KHg₂A'₃: prisms.—KHgA'₃.—Hg₂A'₂: olive-green pp. — HgA'₂: felt-like mass of yellowish crystals.—AgA'₂HgCl₂—PbA'₂: lemonyellow needles (from water).—K₂PtA'₆. Dark-yellow needles (from water).—AgA'. Satiny red scales (Clarke, B. 11, 1325).—AgA'. Satiny crystals, blackened by light. Converted by ICy into volatile crystalline SeCy2 (Linnemann, A. 120, 36).

Methyl ether MeA'. (158°). Yellow oil (Stolte, B. 19, 1577)

Allyl ether C.H.A'. Oil.

Methylene ether CH.A'₂. [132°]. Formed from MeI and alcoholic KSeCN (Proskauer, B. 7, 1281). Rhombohedra (from alcohol). HNO, forms CH2(SeO3H)2.

Ethylene ether C₂H₄A'₂. [128°]. HNO, forms, on boiling, deliquescent C₂H₄(SeO₃H)₂, which yields BaA", sol. hot water. KOH yields

diethylene tetraselenide [131°].

Propylene ether CH₃.CH(SeCN).CH₂(SeCN). [66°]. Formed by heating KSeCN with propylene bromide and alcohol (Hagelberg, B. 23, 1091). White crys-

Trimethylene ether $CH_2(SeCN).CH_2.CH_2(SeCN).$ [51°]. Converted by KOH into trimethylene diselenide [55°].

Benzyl ether v. vol. i. p. 502. Selenocyano-acetic acid CH₂(SeCN).CO₂H. [85°]. Formed from chloro-acetic acid and potassium selenocyanide (G. Hofmann, A. 250, 300). Yellowish needles (from ether and chloroform), v. e. sol. water and alcohol.—BaA'a: amorphous.

Selenocyano-acetoacetic ether

CH3.CO.CH(SeCN).CO2Et. Formed from chloroacetoacetic ether and KSeCN (H.). Liquid.

Selenocyano-acetone CH₃.CO.CH₂.SeCN. Formed from chloro-acetone and KSeCN (H.). Yellowish oil. Gives an oily phenyl-hydrazide and an oxim. Not more sol. alkalis than water.

Selenocyano-acetophenone

C_zH_z.CO.CH_zSeCN. [85°]. Formed by boiling ω bromo-acetophenone with alcohol and KSeCN (Hofmann). Crystalline.

Selenocyano-ethyl-phthalimide

C.H.O.: N.CH. CH. SeCN. [125°]. Formed by heating bromo-ethyl-phthalimide with KSeCy and alcohol (Coblenz, B. 24, 2133). Crystalline mass. Converted by boiling dilute caustic potash into $Se_2(C_2H_4.NH.CO.C_6H_4.CO_2H)_2$ [119°], which forms yellowish crystals and is converted by HCl at 180° into Se₂(C₂H₄NH₂)₂, which yields B"2HCl [188°] and B"2C₂H₂N₂O, [178°], both being crystalline.

 γ -Selenocyano-propyl-phthalimide

C₈H₂O₂:N.CH₂.CH₂.CH₂.SeCN. [102°]. Formed by heating bromo-propyl-phthalimide with KSeCN and alcohol (Coblenz, B. 24, 2134). Converted by boiling dilute (10 p.c.) KOH into crystalline Se₂(C₃H_e,NH.CO.C_eH_e,CO₂H)₂ [84°], whence HCl at 180° forms Se₂(C₃H_eNH₂)₂, which yields B"2HCl [170°] and B"2C₂H₂N₂O, [165°].

•Selencoyano-o-toluic acid. Nitrile

CH₂(SeCN), C₀H₄, CN. o-Cyano-bensyl seleno-cyanide. [121°]. Formed by warming the ni-trile CH₂Cl.C₀H₄, CN with KSeCN (Drory, B. 24, 2564). Long brownish needles (from alcohol)

H₂SO₄ into Se₂(CH₂·C₂H₄·CN), [110°]. HClAq at 100° forms Se and o-oyano-benzyl chloride. Selenocyanuric acid H₃Se₃C₂N₃. Formed

from Cl₃Cy₃ and Na₂Se (Stolte, B. 19, 1578).

Methyl ether. [174°].

Seleno-phthalimidine C_eH₄ < C(NH) > Se.
Formed from CH₂(SeCN).C_eH₄.CN and conc.
H₂SO₄ (Drory, B. 24, 2566). Sol. alcohol.

Reactions.-1. NaOH forms the compound $\mathrm{Se_2(CH_2.C_6H_4.CN)_2}$, crystallising in needles [110°], sol. alcohol and CS2, and converted by HCl at 160° into Se and phthalide.—2. When steam is blown through a mixture of selenophthalimidine, alcohol, and KOH there is formed selenophthalide $C_eH_4 < \stackrel{CO}{CH_2} > Se$, crystallising in broad plates [58°], insol. water, sol. alcohol.—3. Alcoholic potash and MeI form CH₂(SeMe).C₆H₄.CN (180°-200°).

Salts.—B'HCl: needles, slightly sol. water Orange powder.and alcohol.—B',H,PtCl, B'C₆H₂N₂O₇.—B'HBr. [264°]. Needles: sl. sol. water.—B'HI₂ (?) [223°]. Formed by heating CH₂(SeCy).C₆H₁.CN with HIAq at 100°. Darkviolet needles. - Sulphate [145°-150°].

 $C_{\sigma}H_{4} < \stackrel{N}{\sim} Se.$ [76°]. Formed Piaselenole from o-phenylene-diamine and an aqueous solution of selenious acid (Hinsberg, B. 22, 2897). Needles, v. sol. alcohol and ether, sl. sol. water. Its salts are yellow and decomposed by much water with separation of the base. Iodine forms a green periodide.

Amido-piaselenole C₈H₂(NH₂):N₂Se. [150°]. Formed by adding H₂SeO, to a cool solution of tri-amido-benzene hydrochloride and NaOAc (Hinsberg, B. 22, 2898). Brownish-red needles, sl. sol. water, m. sol. alcohol. SnCl, separates Se.—B'HCl: reddish-brown needles, sl. sol.

Oxy-piaselenole. Ethyl derivative C_aH₃(OEt):N₂Se. [104°]. This is formed from $C_0H_3(OEt)(NH_2)_2$ and aqueous H_2SeO_3 . Needles forming a yellowish-red solution in H280.

Methyl-piaselenole CH .CH:C.N Se. [73°]. (267° uncor.). Formed from tolylene-diamine and aqueous SeO, at 80° (Hinsberg, B. 22, Needles, sl. sol. cold water. yields tolylene-o-diamine and Se. Does not react with nitrous acid or with Ac2O .- B'2H2PtCl2.

Chloro-methyl-piaselenole C₆H₂ClMe:N₂Se. [150°]. Formed by warming o-tolylene-diamine with SeO₂ and conc. HClAq (Hinsberg, B. 23, 1395). Small needles, volatile with steam.

Naphthopiaselenole $C_eH_4 < \stackrel{CH:C.N}{CH:C.N} > Se.$

[129°]. Formed from (aß)-naphthylene-diamine sulphate, aqueous NaOAc, and SeO, (Hinsberg, B. 22, 866). Needles, sl. sol. water, m. sol. alcohol, v. sol. ether. Conc. H2SO, gives an intense yellow colour. SnCl, and HCl give Se and naphthylene-diamine.

Seleno-urea CSe(NH₂)₂. [c. 200°]. S. 11 at 19°. S. (alcohol) 3 at 18°. Formed by passing H.Se into a 2 p.c. solution of cyanamide in ether (Verneuil, C. R. 99, 1154; 100, 1296; Bl. [2] 43, 58, 583). White needles, v. sol. Bl. [2] 43, 58, 583). White needles, v. sol. hot water, sl. sol. alcohol and ether. Its solutions are decomposed by light, especially or prisms (from benzene). Converted by dilute in presence of an alkali, with separation of selenium. In presence of air and HClAq it yields (CSeN₂H₄)₃O2HCl, which forms brown crystals with violet reflex, converted by barytawater into Se, seleno-urea, and cyanamide. The compound (CSeN₂H₄)₃O2HBr may be formed in like manner. When the 'oxy-triselenourea' hydrochloride is allowed to stand in the acid liquid in which it is formed, it changes into yellow crystalline (CSeN₂H₄)₂Cl₂. HIAq converts selenourea into (CSeN₂H₄)₂I₂. H₂SO₄ added to a saturated solution of selenourea forms scarlet crystals converted by heating with alcohol (100 pts.) and H₂SO₄ (4 pts.) at 50° into (CSeN₂H₄)₂OH₂SO₄, crystallising in pearly plates.

Ethylene- ψ -seleno-urea $C_3H_6N_2Se$ i.e. CH_2 -Se C:NH. Formed by evaporating a solution of bromo-ethyl-amine hydrobromide (8 g.) with KSeCy (5 g.) to dryness on a waterbath (Baringer, B. 23, 1003). Unstable. Converted by bromine-water into ethylene- ψ -urea.—B'HBr. [170°]. Needles.—B' $_2H_2$ PtCl $_6$.—B' $_6H_3N_3O_7$. [220°]. Needles.

Propylene- $\dot{\psi}$ -selenourea C₄H₈N₂Se. Obtained in like manner from β -bromo-propyl-amine.— B'C₆H₃N₈O₇. [110°].—B'₂H₂PtCl₈. Light-yellow

plates.

Trimethylene - ψ - selenourea C₄H₈N₂Se. Formed from γ - bromo-propyl-amine and KSeCy.—B'HBr. [135°]. White crystals (from alcohol) soon turning red. —B'C₈H₃N₃O₇. [c. 53°]. Small concentric needles.

Seleno-isovaleric aldehyde $C_5H_{10}Se$. [57°]. Crystalline (Schröder, B. 4, 402).

Di-methyl selenophene Se CMe: CH.

Selenoxene. (154° uncor.). Formed by heating acetonyl-acetone with phosphorus pentaselenide P₂Se₃ at 180°. Colourless mobile liquid of slight smell. With isatin and conc. H₂SO₄ it gives a carmine-red colouration. By Laubenheimer's reaction a reddish-brown colour is produced. Benzoyl-formic acid also produces a reddish-brown colouration (Pagl. B. 18, 2955)

reddish-brown colouration (Paal, B. 18, 2255).

SELENIOUS ACID H₂SeO₃. Mol. w. not determined. This compound is the acid of the

anhydride SeO₂.

Preparation.—Se is dissolved in hot conc. HNO₂Aq, the solution is evaporated to dryness, the residue is dissolved in water, BaOAq is added to ppt. traces of H₂SeO₄, the filtrate is evaporated to dryness in a retort, and the residue is heated till SeO₂ sublimes (Thomsen, B. 2, 598); the pure SeO₂ thus obtained is dissolved in c. ½ its weight of hot water, on cooling large crystals of H₂SeO₃ separate (Weber, P. 118, 479). To prevent formation of H₂SeO₄ in the oxidation of Se, Divers a. Shimosé (C. J. 47, 439) recommend to dissolve Se in a large excess of conc. H₂SO₄ instead of using HNO₃.

Properties.—Large white crystals, resembling nitre; effloresces in air; withdraws water from moist air; S.G. 3.0066 at 15.7° (Clausnizer, B. 11, 2009). Poisonous. The acid crystallises unchanged from solution in aqueous acetic acid, or in aqueous C_eH_e (Chabrié, A. Ch. [6] 20, 202). H.F. [Se,O²,Aq] = 56,160 (Th. 2, 274). H₂SeO₃ is a dibasic acid; this is shown by the existence of two salts M¹HSeO₃ and M¹₂SeO₃ (v. Selenites), and also by the heat of neutralisation, which is given by Thomsen (Th. 1, 172) as follows:

[H²SeO³Aq, NaOHAq] = 14,770, [H²SeO³Aq] 2NaOHAq] = 27,020, [H²SeO³Aq, 4NaOHAq, = 27,480. As no derivative of H₂SeO₃ exists corresponding with C₂H₂SO₂OH, and as only one ethyl selenite SeO(OEt)₂ can be obtained, and as the compound SeO.Cl.OEt has been isolated, it appears that the acid is SeO(OH)₂ and not SeO₂OH.H (Michaelis a. Landmann, B. 13, 656; A. 241, 150).

Reactions.—1. Heat causes H.SeO, to decompose to SeO, and H.O.—2. Many oxidisable bodies remove O from H.SeO.. For instance, sulphur dioxide passed into H.SeO, Aq ppts. Se; the reduction is complete only when the compounds interact in the ratio H.SeO,:2SO, (Schulze, J. pr. [2] 32, 390); the presence of HCl hastens the reaction (v. Michaelis, J. Z. 6, 88; Rathke, A. 152, 194). When SO, and H.SeO, interact in other proportions than H.SeO,:2SO, thioselenic acids are formed (cf. p. 435). Hydrogen sulphide ppts. Se and S in the ratio Sc:2S (Divers a. Shimidzu, C. J. 47, 441). Various metals, such as Cu, Fe, Ag, or Zn, also reduce H.SeO,Aq is partly reduced by exposure to air containing dust. M. M. P. M.

SELENITES. Salts of selenious acid (H₂SeO₃), M₂¹SeO₃, and M¹HSeO₃. The alkali salts are formed by adding alkalis to H₂SeO₃Aq and evaporating; most of the other selenites are formed from the alkali salts by double decomposition.

Selenites give selenides, or metallic oxide and Se, when heated with charcoal. Heated with NH₄Cl, out of air, they give a sublimate of Se. Most selenites, except those of the alkalis, are insol. in water.

Many selenites combine with SeO₂ to form acid salts; many basic selenites are also known.

Boulzoureano (Bl. [2] 48, 209) found that many selenites can be obtained in well-formed crystals by adding Na,SeO₃Aq to solutions of metallic salts, and heating the pp. so formed with very dilute H₂SeO₃Aq in sealed tubes at 200°; also by dissolving metallic carbonates in dilute H₂SeO₃Aq, adding water, and heating in sealed tubes at 200°. The older investigations of selenites were by Berzelius (v. Lehrbuch) and Muspratt (C. J. 2, 52). Boulzoureano (l.c.; also A. Ch. [6] 18, 289) has made a more thorough investigation of these salts. Nilson (Bl. [2] 21, 253; 23, 260, 353, 494) has also prepared and described many selenites.

It has been thought sufficient to give the names of the metals selenites of which have been analysed. Abstracts of B.'s papers will be found in C. J. 54, 220; 60, 262; and an abstract of Nilson's paper will be found in the 3rd Swpp. of the first edition of this DICTIONARY, p. 1786.—Al, Am, Sb, Ba, Be, Bi, Cd, Ca, Ce, Cr, Co, Cu, Di, In, Fe, La, Pb, Li, Mg, Mn, Hg (cf. Cameron a. Davy, Trans. I. 28, 146), Ni, K, Ag, Na, Sr, Tl, Th, U, Zn. M. M. P. M.

SELENO-ANTIMONATES v. Antimonates, vol. i. p. 286.

SELENOCYANHYDRIC ACID v. vol. ii. p. 348.

SELENOGYANIDES v. vol. ii. p. 348. SELENOMETAHYPOPHOSPHITES RIPSe and Ru(PSe)₂ v. Diphosphorus selenide, this vol. p. 145.

R₂¹¹P₂Se₇ v. Phosphoric selenide, p. 145.

SELENOPYROPHOSPHITES R41P2Se, and R₂^{II}P₂Se₅ v. Phosphorous selenide, p. 145.

SELENOSAMIC ACIDS, SALTS OF. Selenosamic acid SeO(NH2)OH has not been isolated, but the NH, salt of this acid, and also the NH, salt of disclenosamic acid, 2SeO(NH2).(OH)2,

have been obtained.

Ammonium selenosamate SeO(NH2)ONH, is formed by the action of dry NH, on SeO2 dissolved in alcohol. It is a white, crystalline, deliquescent salt; very unstable, easily giving off NH, and forming disclenosamate. Water partially converts this salt into (NH₄), SeO₃.

Ammonium diselenosamate 2SeO(NH₂).OH.ONH₄ is formed by boiling an alcoholic solution of the selenosamate, or by letting the solution stand over H2SO, in vacuo. It crystallises from alcohol in large, deliquescent prisms (v. Cameron a. Macallan, Pr. 44, 112).

M. M. P. M. SELENOTHIO-OXIDE SeSO, v. Selenion thio-oxide, under Selenion oxides, p. 441. SELENOTHIO-OXYCHLORIDE SeSO₃Cl₄ v.

SELENION THIO-OXYCHLORIDE, p. 441.

M.SeSO. SELENOTHIOSULPHATES Thioselenates, under Selenates, p. 434. SELENOTRITHIONATES M₂SeS₂O

Dithiotriselenates, under Selenates, p. 434. SELENOTRITHIONIC ACID H2SeS2O.

Dithiotriselenic acid, under Selenates, p. 435. SELENYL BROMIDE and CHLORIDE v.

Selenion oxygnomide and Oxychloride, p.

SEMINOSE v. MANNOSE.

SENEGIN v. SAPONIN.

SENNA v. CATHARTIC ACID.

SENNITE is identical with PINITE, the methyl ether of dextrorotatory inosite.

SEPTYL. A name for HEPTYL.

SEQUOIENE C₁₃H₁₀. [105°]. (290°-300°). V.D. 5·66 (obs.). Occurs, together with C₁₀H₁₆ (155°), S.G. 15.852, $[a]_D = 24°$, and oily $C_{18}H_{20}O_{3}$ (229°), in the leaves of Sequoia gigantea (Lunge a. Steinkauler, B. 13, 1656). White plates with characteristic smell, volatile with steam; sol. alcohol, ether, benzene, and chloroform.

SERICIC ACID C15H30N4O7. Formed by the action of conc. baryta-water on silk (Champion, C. R. 72, 330). Deliquescent, translucent mass.

-BaA'₂.-PbA'₂: pp.

SERIN C₃H,NO₃. a-Amido-hydracrylic acid?

S. 3 at 10°; 5 at 20°. Formed by boiling silk with dilute H₂SO₄ (Crämer, J. pr. 96, 76; Baumann, B. 15, 1735; Melikoff, C. C. 1881, 354). Monoclinic crystals, insol. alcohol and ether. Converted by nitrous acid into glyceric acid.— CuA'₂.—HA'HCl: needles, v. e. sol. water.— HA'HNO₃: minute needles.

SERINE v. PROTEÏDS.

SERUM ALBUMEN v. PROTEÏDS. SERUM CASEIN v. PROTEÏDS.

SERUM GLOBULIN v. PROTEÏDS.

SESAME OIL. S.G. 20 9191. $\mu_D = 1.474$ at 20° (Long, Am. 10, 392). A fatty oil obtained from the seeds of Sesamum orientale. It contains glycerides of oleic and linoleic acids (Pohl, J. pr. 63, 400; Hazura a. Grüssner, M. 10, 247). Gives a raspberry-red colour on warming with HClAq and cane-sugar (Zipperer, Chem.

SELENOPYROPHOSPHATES R. P.Se, and Zeit. 11, 1600). The oil contains 05 p.c. of sesamin C₁₈H₁₈O₅ [118]. S. (alcohol) 27 at 20°; 8.07 at 78°. S.G. 1.305. H₂SO₄ containing HNO, colours sesamin green and afterwards bright red (Tocher, Ph. [3] 23, 700).

SEXTYL. Another name for HEXYL.

SHEA BUTTER, obtained in West Africa from the nuts of Bassia Parkii, contains olein and stearin (Oudemans, J. pr. 89, 205).

SHELLAC v. LAC.

SHIKIMIC ACID C,H10O, i.e.

 $CH(OH) < \frac{CH(OH).CH}{CH_2.CH(OH)} > C.CO_2H.$ [184°]. S.G. 14 1.599. $\mu_{\infty} = 1.303$ at 14°. $[\alpha]_D = -186^\circ$ in a 7 p.c. solution. S. (ether) .015 at 23°; S. (alcohol) 2.25. Occurs in the fruit of Illicium religiosum (Japanese, shikimi) and Chinese star-anise (Eykman, R. T. C. 4, 49; 5, 299; B. 24, 1278). Crystalline powder composed of minute needles, sl. sol. alcohol and CHCl, nearly insol. ether. Lævorotatory. Yields poxy-benzoic acid when heated with HClAq.-NHA': crystals.

Dihydride

 $CH(OH) < CH(OH) \cdot CH_2 > CH.CO_2H.$ [175°]. S.G. 2 1.47. $\mu_{\infty} = 1.324$ at 19°. $[a]_D = -18^\circ$. S. 14 in the cold. Formed by reducing the acid

with sodium-amalgam in presence of HClAq. Monoclinic crystals. Converted by HCl into benzoic acid.

Dibromide

CH(OH) < CH(OH).CHBr > CBr.CO₂H. [c. 188°]

 $\mu_{\infty} = 1.295$. [a]_D = -58° . Formed by adding Br to a cold solution of the acid. Tetrahedra. Converted by moist Ag₂O into C₇H₂BrO₅ [235°], S. 6·3 at 17°, $[\alpha]_D = +22^\circ$, crystallising in hexagonal needles, sl. sol. ether, converted by baryta into oxyshikimic acid [156°].

SHIKIMIPICRIN $C_{10}O_{3}(?)$. [200°]. S. 6.2 in the cold. Occurs in the fruit of Illicium religiosum (Eykman, R. T. C. 4, 53). Large

crystals, m. sol. alcohol, sl. sol. ether.

SHIKIMOLE v. SAFROLE.

SIEGBURGITE. A fossil resin found near Siegburg. It contains styracin, metastyrene, and some free cinnamic acid (Klinger a. Pitschke, B. 17, 2742).

SIKIMINE. [175°]. A poisonous substance in the seeds of *Illicium religiosum* (Eykman, Ph. [3] 11, 1046). Prisms, sl. sol. water, ether,

and CHCl,, v. sol. alcohol and HOAc.

SILICA SiO₂. (Silicon dioxide. anhydride.) Mol. w. not known. S.G. quarts 2.647 to 2.652, tridymite 2.282 to 2.326, amorphous 2.20 at 15.6° (Schaffgotsch, P. 68, 147). S.H. quarts 1737 at ord. temp., 305 at c. 400°, constant from 400° to over 1000° (Pionchon, C. R. 106, 1344). H.F. $[Si,O^2] = 219,240$ from amorphous Si; 211,120 from crystalline Si (Troost a. Hautefeuille, C. R. 70, 252).

Occurrence.—Compounds of SiO2 are very widely distributed in large quantities; quartz, flint, chalcedony, jasper, opal, &c., are nearly pure SiO2. Silica is the chief constituent of sandstones; felspar, agate, and a vast number of other minerals contain silica. Silica (or silicates) is also found in many plants.

Preparation.—(i) Amorphous silics.—
1. Quartz is finely powdered, best by heating to

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redness and plunging into cold water, and heated with c. 6 times its weight of Na₂CO₃ in a Pt crucible till the whole fuses quietly; the cold mass is treated with water, the solution, when clear, is poured slowly into excess of fairly conc. HClAq, the liquid (after filtration, if necessary) is evaporated to dryness in a Pt dish, the residue is moistened with conc. HClAq, warmed, and allowed to stand for an hour or two; this treatment with acid is repeated two or three times; water is then added and, after warming and standing, the solution is filtered off, the residual SiO₂ is washed with hot water, dried, and heated to redness.—2. SiF, (v. p. 459) is passed into water, the gelatinous SiO, which is ppd. is pressed between linen, thoroughly washed, dried, and heated to redness.

(ii.) Crystalline silica.-1. Moist ppd. SiO2-e.g. the pp. got by passing SiF, into water-is dissolved in boiling NaOHAq, and the solution is heated in a sealed tube; crystals of tridymite (v. infra, Properties) separate below 180°, and crystals of quartz separate above 180° (Maschke, P. 145, 549).—2. Small tubes of hard glass are half-filled with solutions of dialysed silicic acid (v. Silica, hydrates of, infra) containing c. 10 p.c. SiO₂, closed by fusion, and heated to 250° for some months; small crystals of quartz are thus obtained (Kroustchoff, C. R. 104, 602).—3. Any hydrate of SiO₂ yields crystalline silica by heating in a forge-fire (Fremy, A. Ch. [3] 38, 327).—4. Crystalline SiO₂ is formed by fusing silicates with microcosmic salt (G. Rose, J. pr. 101, 228), or with borax (G. Rose, B. 2, 388). For other methods of preparing crystalline silica v. Rammelsberg (B. 5, 1006), Hautefeuille (C. R. 86, 1133; 90, 830), Sénarmont (A. Ch. 42, 129), Friedel a. Sarasin (Bl. [2] 31, 481), and Daubrée (Ann. M. [5] 12, 297)

Properties.—The mol. w. of silica is unknown, but there is reason to think it is a considerable multiple of that indicated by the formula SiO₂. By applying the cryoscopic method to an aqueous solution of colloidal silicic acid (v. Metasilicic acid, infra) Sabanéeff (J. R. 21, 515; Abstract in C. J. 58, 1215) obtained numbers pointing to silica being a very high polymeride of SiO₂. The great infusibility and non-volatility of silica, especially when this compound is compared with CO₂, make it probable that the molecular formula of silica is $nSiO_2$, where n has a large value (v. Mendeléeff's Principles of Chemistry, vol. ii. p. 93).

Orystalline silica. SiO₂ crystallises as quarts in colourless, transparent, hexagonal, six-sided prisms having the axes-ratio 1:1.0999 (S.G. c. 2.65), and as tridymite in hexagonal forms with the axes-ratio 1:1.631 (S.G. c. 2.3). Crystalline SiO₂ is insol. or almost insol. in alkali solutions; it is attacked by HFAq slowly and without notable rise in temperature.

Amorphous silica is a white, loose, tasteless powder; S.G. 2.2. It dissolves in hot alkali solutions; after continued heating to a high temperature it is less sol. or nearly insol., owing to change into the crystalline form. Dissolves in HFAq, with intumescence and considerable rise of temperature.

Silica is fusible in the O-H flame to a

colourless glass; when fused it may be drawn into extremely fine threads. Cramer (Zeit. filr angewand. Chem.) finds that quartz is volatile at a temperature whereat Pt boils (v. Abstract in C. J. 64, ii. 164). SiO₂ is insol. in acids, except in HFAq. It is slightly volatilised in steam (v. Jeffreys, A. 39, 255). Silica is insoluble in molten microcosmic salt (NaNH₄HPO₄).

Reactions.-1. Fused with alkalis, alkali silicates are formed; with alkali carbonates CO₂ is evolved.—2. Decomposes most solid salts by fusion with them with formation of silicates (cf. Silicates, p. 448). According to Mills a. Meanwell (C. J. 39, 533) SiO₂ does not react with pure Na₂SO₄ or K₂SO₄ even at a very high temperature.—3. SiO₂ is decomposed by potassium at a red heat, forming K silicate and silicide; it is also reduced at red heat by calcium, strontium, and magnesium (Bunsen a. Matstrontium, and magnesium (Bunsen a. Matthiessen, A. 94, 111; Phipson, Pr. 13, 217; Parkinson, C. J. [2] 5, 128; Winkler, B. 23, 2642).—4. Heated with carbon and certain metals, SiO₂ is reduced, with formation of silicides of the metals; Ir, Pd, Pt, and Ru react in this way (Boussingault, C. R. 82, 591; cf. Warren, C. N. 60, 5). Au and Ag reduce SiO₂ only when strongly heated therewith in presence of when strongly heated therewith in presence of Na_2SiF_a and Na (W., l.c.).—5. Dry SiO_2 (prepared from SiF_4) which has not been strongly heated reacts with molten metaphosphoric acid to form SiO₂.P₂O₅ and SiO₂.2P₂O₅.4H₂O (Hautefeuille a. Margottet, C. R. 96, 1052; 104, 56; v. Silico-PHOSPHORIC OXIDES, p. 462).—6. Mixed with finely divided carbon and heated in hydrogen sulphide, SiS₂ is formed (v. Silicon sulphide, p. 462).—7. A mixture of SiO₂ and carbon heated in chlorine or hydrogen chloride gives SiCl₄ and CO (v. SILICON TETRACHLORIDE, p. 458); when bromine or hydrogen bromide is used, SiBr, is formed (v. Silicon tetrabromide, p. 457).—8. With hydrogen fluoride, SiF₄ is obtained (v. SILICON TETRAFLUORIDE, p. 459).—9. Heated to low redness in vapour of boron trichloride, SiCl, and B_2O_3 are formed (Troost a. Hautefeuille, A. Ch. [5] 7, 476).

SILICA, HYDRATES OF. (Silicic acids.) The hydrates of SiO₂ act as weak acids. The compositions of these hydrates are very uncertain. The silicates nSiO₂.mRO are derivable from hydrates of SiO₂, some of which have been isolated while others are hypothetical (v. SILICATES, p. 448).

METASILICIC ACID H₂SiO₃ = SiO₂·H₂O. (Silica monohydrate.) Obtained by the action of moist air on Si(OEt)₄; Si(OEt)₄+3H₂O = SiO₂·H₂O+4EtOH (Ebelmen, J. pr. 37, 359). Also by dialysing a mixture of Na silicate and excess of HClAq until the liquid in the dialyser is free from chlorides, concentrating by boiling in a flask, allowing to gelatinise, and drying the jelly-like mass over H₂SO₄ for two days (Graham, T. 1861. 183). SiO₂·H₂O is described by Graham as a transparent, glassy, lustrous solid; insol. water.

ORTHOSILICIC ACID H₄SiO₄ = SiO₂.2H₂O. (Silica dihydrate.) This hydrate of SiO₂ has not been isolated with certainty; it is supposed to exist in the solution obtained by dialysing a solution of K or Na silicate with excess of HClAq; but as the compositions of the silicates of K and Na are very doubtful, the argument

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based on the supposition that these silicates are ortho- salts—and, therefore, will yield ortho- acid—is not very trustworthy. Various esters of the form Si(OX), exist where X is Me, Et, &c. The solution of SiO_2xH_2O , when x is supposed to = 2, obtained by dialysing K or Na silicate solution with excess of HClAq till the liquid in the dialyser is free from chlorides, is described by Graham (l.c.) as a clear liquid, which may be boiled in a flask till it contains c. 13 p.c. SiO2 without separation of any solid. The liquid soon becomes opalescent on keeping, and after a time sets to a transparent, colourless jelly, which is insol. water, and, dried over H₂SO₄, gives SiO₂·H₂O (v. Metasilicic acid, p. 447). The solution of SiO, xH,O is coagulated by a minute trace of an alkali or alkaline earth carbonate, also by CO₂, but not by dil. H₂SO₄Aq, HNO₅Aq, or H.C₂H₃O₄Aq. The solution reddens litmus (cf. Doveri, A. 64, 256; Liebig, A. 94, 373; Karsten, P. 6, 357; Fuchs, A. 82, 119; Struckmann, A. 94, 341; Rose, P. 108, 20; Kühn, J. pr. 59, 1; Fremy, A. Ch. [3] 38, 314; Maschke, J. pr. 68, 234).

TRISILICIC DIHYDRATE 3SiO.2H₂O = H₄Si₃O₈. By decomposing K or Na silicate solution by CO₂, or SiF₄ by H₂O, and drying the gelatinous pp. so formed in vacuo, Fremy (l.c.) obtained a hydrate containing 16·2 to 16·8 p.c. water (8SiO₂·2H₂O requires 16·67 p.c. water). Several naturally occurring silicates may be regarded as derived from this hydrate (v. Silicates,

infra).

Disilicic monohydrate $2\mathrm{SiO}_2.\mathrm{H}_2\mathrm{O} = \mathrm{H}_2\mathrm{Si}_2\mathrm{O}_3$. The gelatinous pp. formed by decomposing SiF_4 by water, after drying for six weeks in air at $20^\circ-25^\circ$, was found by Merz (J. pr. 99, 1771) to contain from 13·1 to 13·5 p.c. water (the formula

given requires 13:05 p.c. water).

OTHER HYDRATES. By drying the gelatinous pps. obtained by decomposing SiF, by water, or by decomposing alkali silicate solutions by weak acids and washing (v. Kühn, J. pr. 59, 1), solids are obtained of varying and uncertain composition (v. Carnelley a. Walker, C. J. 53, 80); hydrates of fair definiteness agreeing approximately with the formulæ 3SiO_xH_xO = H_xSi_xO_x, 9SiO_x2H_xO = H_xSi_xO_x, and 6SiO_xH_xO = H_xSi_xO_x, have been obtained (Fuchs, A. 82, 119; Gottlieb, J. pr. [2] 6, 185; Merz, J. pr. 99, 1771). For methods of preparing gelatinous pps. of SiO_xxH_xO v. Struckmann, A. 94, 337; Kuhlmann, A. 41, 231; Liebig, A. 94, 374; Guyton-Morveau, A. Ch. [2] 31, 246; Hautefeuille a. Margottet, C. R. 96, 1052. Concerning the absorption of water by SiO_x v. van Bemmelen, B. 11, 2232; 13, 1466.

Thomsen (Th. 1, 215) has examined the thermal phenomena accompanying the interaction of NaOHAq and SiO₂Aq; the SiO₂Aq was prepared by adding exactly an equivalent of HCl in dilute solution to Na silicate solution of known composition. He gives the following tables:—

m	[mNaOHAq,SiO*Aq]	\boldsymbol{x}	[NaOHAq,xSiO'Aq]
ł	2652	Ŧ	1353
ř	3241	Ţ	2615
4	8555	3	3548
ĭ	4316	Ī	4316
4	4731	8	533 2
2	5230	2	6483
4	5412	8	705A

There seems to be no fixed point of neutral-

isation. Putting [NaOHAq,xSiO²Aq] = $\frac{x.O}{x+n}$

where C and n are constants having the values C=13,410 and $n=2\cdot13$, the calculated values agree very fairly with the observed so far as data go; if the formula is supposed to be accurate then the heat of neutralisation of SiO₂Aq, when SiO₂ is taken as $=\infty$ in the reaction NaOHAq + SiO₂Aq, is =13,410, which is about the thermal value of the interaction of an acid with one molecular weight of NaOH. The affinity of SiO₂Aq is almost nil. M. M. P. M.

SILICATES. Compounds of SiO, with basic radicles. The basic radicles of silicates are generally oxygen-containing groups; sometimes they contain negative elements other than O. Silica, like other feebly acidic oxides, combines with bases in many proportions, forming compounds which may be represented as normal salts, and also compounds of the type of basic and acidic salts. The general composition of silicates is given by the formula nSiO₂.mR, where R = basic radicle. Most of the silicates occurring in rocks vary in composition between the limits SiO24MO and 2SiO2MO, where M = a mono- or di-valent metal, and the limits $SiO_2.2M_2O_3$ and $6SiO_2.M_2O_3$ where M=a trivalent The metals most commonly found in metal. these silicates are Al, K, Na, Ca, Mg, Fe, and Mn. A fair number of natural silicates contain Cl or F as part of the basic radicle. Few pure silicates occur in rocks: the siliceous minerals are generally mixtures of isomorphous silicates; hence it is not possible to deduce satisfactory formulæ for naturally occurring silicates solely from the results of analyses of minerals.

Mendeléeff (Principles of Chemistry, 2, 111) compares the siliceous minerals to alloys. Several silicates which occur as mineral species have beer formed artificially; the study of the alteration-products of natural silicates, and the comparison of silicates with carbon compounds (v. e.g. Mendeléeff, l.c., note p. 111) throw light on the constitution of these bodies.

The most important memoirs on the constitution and chemical relations of the silicates are to be found in Groth's Tabellarische Uebersicht der Mineralien nach ihren Krystallographischchemischen Beziehungen (3rd ed.), pp. 89 et seq., and in a series of papers by F. W. Clarke (Am. 10, 120; Am. S. [3] 38, 384; 40, 303; 43, 190), (v. also Clarke in U.S. Geological Survey Bulletins, No. 90 [1892]; also A. C. J. May 1893).

Groth and Clarke both look on the natural silicates as derived from a few fairly simple types. They are very stable compounds; most of them are formed at high temperatures, their number is not very great, and a few typical forms containing a small number of common metals are very widely distributed. Among the commoner silicates are SiO_{2.2}MO, 2SiO_{2.3}MO, 3SiO_{2.2}MO, where M is very frequently an alkaline earth metal; 3SiO_{2.2}R₂O_{3.2} 2SiO_{2.2}R₂O_{3.3} 6SiO_{2.2}R₂O_{3.4}MO, and 3SiO_{2.2}R₂O_{3.3}MO, where R is Al or Fe, and M is K, Na, or an alkaline earth.

The consideration of the probable constitution of the typical silicates may be approached by seeking to derive these salts from the silicio acids nSiO2.mH2O. Metasilicic acid, SiO2.H2O, has been isolated; the formula may be written H_2SiO_3 or $OSi(OH)_2$. Orthosilicic acid, $SiO_2.2H_2O=H_4SiO_4=Si(OH)_4$, also probably exists. A few other hydrates of silica have been isolated, but their compositions are not quite definitely established; $2SiO_2.H_2O = H_2Si_2O_5$ and 3SiO₂.2H₂O = H₄Si₂O₈, seem to be fairly certain.

Silicates derivable from metasilicic acid O:Si(OH)₂. The typical empirical formulæ of normal salts of this class are SiO₂.M₂O, $3SiO_2.M_2O_3$, $4SiO_2.R_2O_2.M_2O$, SiO2.MO, 4SiO₂.R₂O₃.MO. The atomic ratio of Si to O in these salts is 1:3; this ratio is maintained in the acid salts. Silicates of the forms SiO₂.2MO and SiO₂.M₂O₃ may be represented as basic salts of this class; the atomic ratios of Si to O in these are 1:4 and 1:5. Examples of normal salts of this class are sodium silicate O:Si(ONa)₂, wollastonite O:Si.O₂Ca, and leucite (O:Si.O₂)₄Al₂K₂. The basic salts may be regarded as derived from O:Si(OH)2 by replacing each H by a monovalent basic radicle, such as AlO or ZnOH; for instance, siliceous calamine SiO_{2.2}ZnO.H₂O may be formulated O:Si(O.ZnOH)₂, and kyanite SiO_{2.}Al₂O₃ is probably O:Si(OAl:O)₂. If only one H is replaced in O:Si(OH), the product will be an acid salt of the metasilicate class; gyrolite, for instance, 3SiO₂.2CaO.H₂O may be represented as O:Si(OH)O.Ca.O.Si.O.Ca.O.(HO)Si:O.

Silicates derivable from hypothetical metadisilicic acid

$$2 \left(O.Si(OH)_2\right) - HOH = HO.Si.O.Si.OH.$$
 The

typical empirical formulæ of normal salts of this class are 2SiO2.M2O, 2SiO2.MO, 6SiO2.R2O8, and 8SiO2.R2O3.M2O. The atomic ratio of Si to O in these salts is 2:5; this ratio is maintained in the acid salts. Silicates of the form 2SiO22MO.H2O, wherein the atomic ratio of Si to O is 2:7, may be regarded as basic salts of Petalite 8SiO2.Al2O3.Li2O is prothis class. bably an example of a normal salt of the metadisilicate class; thus

Picrosmine 2SiO2.2MgO.H2O may perhaps be a basic salt of this class, (HOMg)O.Si.O.Si.O(MgOH); but it may also be 0

regarded as an acid orthodisilicate (v. infra). One of the soluble glasses, 4SiO2.Na2O.H2O, may be written as an acid salt of this class,

HO.Si.O.Si.ONa.

Silicates derivable from ortho-silicic acid Si(OH). The typical empiri-cal formulæ of normal salts of this class are SiO, 2MO, SiO, 2MO, 3SiO, 2MO, and SSiO, MO, 3M,O. The atomic ratio of Si to O in these salts is 1:4; this ratio is maintained in the acid salts. Salts of the forms SiO2.4MO.2H2O and SiO2.2M2O2, atomic ratio of Si to 0 = 1:8, may be formulated as basic ortho- as present in orthoclase also. Thus Vol. IV.

Olivine SiO2.2MgO, and zirconite silicates. SiO...ZrO, are likely normal salts of this class;

 $Mg < {\stackrel{O}{\bigcirc}} Si < {\stackrel{O}{\bigcirc}} Mg$ and $Si(O_4Zr)$. 3SiO₂.2Al₂O₂, is also a normal orthosilicate; AlOg:Si.OOg:Si:OgO.Si:OgAl.

Andalusite SiO2.Al2O3 is probably partly a basic salt and partly a normal salt of this class; Si(O₃Al).O.AlO. Dioptase SiO₂.CuO.H₂O may be formulated as an acid salt of the orthosilicate class; Si(OH)2.O2Cu.

Silicates derivable from hypothetical orthodisilicic acid $2Si(OH)_4 - H_2O = (HO)_3Si.O.Si(OH)_3$. The typical empirical formulæ of normal salts of this class are $2SiO_2.8M_2O$, $2SiO_2.8MO$, $2SiO_2.M_2O_3$, and $4SiO_2.M_2O_3.3M_2O$. The atomic ratio of Si to O in the normal and acid salts is 2:7. Serpentine 2SiO₂.3MgO is probably a normal salt of this class, (MgO₂)Si.O. Si(O₂Mg). Okenite O.Mg.O

2SiO2.CaO.2H2O seems to be an acid salt of this class, HO.Si.O.Si(OH), picrosmine O,Ca

2SiO₂.2MgO.H₂O may also be an acid orthodisilicate, MgO2.(HO)Si.O.Si(OH).O2Mg; but it may be formulated as a basic metadisilicate (v. supra).

Silicates derivable from hypothetical orthotrisilicic acid. By removing 2H2O from 3Si(OH)4 the acid

(HO)₂Si.O.Si.O.Si(OH)₃ would remain. Some $(OH)_2$

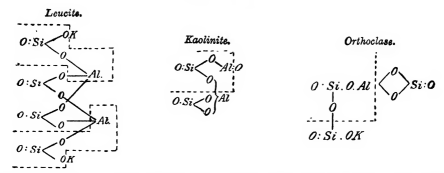
natural silicates may be looked on as salts of this hypothetical acid; meerschaum, for instance, 3SiO₂.2MgO.2H₃O, may have the formula MgOq.(HO)Si.O.Si.O.Si(OH).OqMg. Groth (l.c.), $(OH)_2$

however, thinks it is not necessary to call in a hypothetical trisilicic acid to express the composition of silicates.

Some silicates which do not fit into any of the classes already mentioned may be looked on as derived from more than one of the acids simultaneously. Others may be represented as salts of hypothetical acids formed by eliminating water from one or other of the acids that have been mentioned. Orthoclase, 6SiO₂.Al₂O₃.K₂O₃ may be taken as an example of a silicate derived, according to Groth (l.c.), from two acids, metasilicio and metadisilicio:

this formula is in keeping with the facts (1) that felspars of the orthoclase class are derived from silicates, such as leucite, which are metasilicates, and (2) that orthoclase readily yields kaolinite 2SiO, Al₂O₃, which may be a basic metadisilicate. Writing leucite as a metasilicate, and kaolinite as a basic metadisilicate, it is seen that both contain the group O:Si<0-A1-, a group which is represented

G G



The group common to the three silicates is indicated by the dotted lines. If HHO were removed from orthodisilicic acid the product

Some of the natural silicates may be represented as derived from one or other of these acids. Acids containing OH yield Cl derivatives wherein OH is replaced by Cl, generally by interaction with PCl. As Cl derivatives of some of the hypothetical silicic acids are known, the existence of these derivatives is an argument in favour of representing certain silicates as salts of acids which have not themselves been isolated. Thus, besides SiCl, which corresponds with orthosilicic acid Si(OH), and SiCl₂·SH, which is derived from the hypothetical orthothicsilicic acid, there exists Si₂OCl₂ corresponding with ortho-disilicic acid Si₂O(OH)₈, Si₄O₂Cl₁ corresponding with orthothetrasilicic acid Si₄O₃(OH)₁₀, Si₄O₄Cl₂ corresponding with the acid Si₄O₄(OH)₈, and Si₄O₅Cl₆ corresponding with the acid Si₄O₅(OH)₆ (v. supra).

O:Si.O.Si.O.Si.O.Si:O7

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 $Si_4O_7(OH)_2$

Clarke (l.c.), recognising the two main divisions of ortho-silicates and meta-silicates, seeks to derive the various members of each class as

far as possible from the typical normal aluminium salts by substituting metals or basic groups, wholly or partially, for Al.

Orthosilicates. Clarke starts with normal Al orthosilicate, 3SiO₂2Al₂O₃ = Al₄(SiO₄)₂, which forms the mineral species xenolite; allied to xenolite are fibrolite SiO₂Al₂O₃ = Al₂SiO₃, topax SiO₂Al₂O₂F₂ = Al₂SiO₄F₂, muscovite 6SiO₂.3Al₂O₃.K₂O.2H₂O = 2Al₂KH₂(SiO₄)₃, paragonite, corresponding with muscovite but containing Na in place of K, and eucryptite 2SiO₂.Al₂O₃.Li₂O = 2AlLiSiO₄. By trebling the formulæ given for fibrolite and topax, using the simplest formula given for muscovite and paragonite, and trebling the simplest formula given for eucryptite, Clarke arrives at the following comparable formulæ for the six silicates:—

Xenolite.	${\it Fibrolite}.$
$\mathbf{A}1 \begin{cases} \mathbf{O}.\mathbf{Si}:\mathbf{O_3}:\mathbf{A}1 \\ \mathbf{O}.\mathbf{Si}:\mathbf{O_3}:\mathbf{A}1 \\ \mathbf{O}.\mathbf{Si}:\mathbf{O_3}:\mathbf{A}1 \end{cases}$	$\mathbf{Al} \begin{cases} \text{O.Si:O}_{\bullet}:(\text{AlO})_{\bullet} \\ \text{O.Si:O}_{\bullet}:\text{Al} \\ \text{O.Si:O}_{\bullet}:\text{Al} \end{cases}$
$\begin{array}{c} Topaz.\\ \text{O.Si:O_3:(AlF_2)_0}\\ \text{O.Si:O_3:Al}\\ \text{O.Si:O_3:Al} \end{array}$	$\begin{array}{c} \textit{Muscovite}.\\ \text{O.Si:O}_{i}\text{:KH}_{2}\\ \text{O.Si:O}_{i}\text{:Al}\\ \text{O.Si:O}_{i}\text{:Al} \end{array}$
$\begin{array}{c} Paragonite.\\ \text{O.Si:O_i:NaH_2}\\ \text{O.Si:O_i:Al}\\ \text{O.Si:O_i:Al} \end{array}$	$\mathbf{Al} \left\{ \begin{matrix} \mathbf{C.Si:O_s;Li_s} \\ \mathbf{O.Si:O_s;Al} \\ \mathbf{O.Si:O_s;Al} \\ \mathbf{O.Si:O_s;Al} \end{matrix} \right.$

These silicates are represented as derived from normal Al orthosilicate by substituting either metals, metals and hydrogen, or basic groups, for Al. If such substitution is carried further a series of silicates may be obtained, for some of which Clarke suggests the following formulæ:—

Dumortierite.	Grossularite.
$\mathbf{Al} \begin{cases} \mathbf{O.Si:O_s:(AlO)_s} \\ \mathbf{O.Si:O_s:(AlO)_s} \\ \mathbf{O.Si:O_s:Al} \end{cases}$	$\mathbf{Al} \left\{ \begin{matrix} \mathbf{O.Si:O_3} \\ \mathbf{O.Si:O_3} \\ \mathbf{O.Si:O_3:Al} \end{matrix} \right\}$
Prehnite.	Natrolite.
$\mathbf{Al} \begin{cases} \mathbf{O.Sii:O_3:CaH} \\ \mathbf{O.Sii:O_3:CaH} \\ \mathbf{O.Sii:O_3:Al} \end{cases}$	$\mathbf{Al} \left\{ \begin{matrix} \mathrm{O.Si:O_s:NaH_s} \\ \mathrm{O.Si:O_s:NaH_s} \\ \mathrm{O.Si:O_s:Al} \end{matrix} \right.$

Fibrolite alters by hydration, giving westanite 4SiO₂.8Al₂O₃.H₂O = 2Al₃HSi₂O₃, kaolinite 2SiO₂.Al₂O₃.2H₂O = Al₂Si₂H₁O₉, and montmorillonite 4SiO₂.Al₂O₃.7H₂O = 2AlSi₂H₁O₃. Clarke suggests the following formulæ for these products of alteration of fibrolite:—

If it is supposed that two groups Al(O.SiiO₃)₃ combine with elimination of one of the O.SiO₃ groups (-SiO₂+2O), then there remains the

collocation of atoms,

Al

O.SiO₂
O.SiO₂
O.SiO₃
O.SiO₃
O.SiO₃
O.SiO₃

a number of silicates, especially iron and magnesian micas, may be derived. Thus—

Nosean.

O.Si:O₃:Na₂
O.Si:O₃:Al
O₂:Si:O₂:Na₂
Al
O.Si:O₃:Al
SO₄.Na

Clarke (Am. 10, 126) suggests that ultramarine may be analogous to these silicates; (O.Si:O_s:Na_s

The metasilicates are considered by Clarke (l.c.) in a way similar to that whereby he has elucidated the orthosilicates. The starting-point is Al metasilicate, $Al_x(SiO_3)_s$. This salt is not itself known. The formula may be

(1)
$$0:Si:O_2$$
 $O:Si:O_2$ $O:Si:O_2$

Inasmuch as many metasilicates of the form AlM(SiO₃)₂, where M=Li, K, Na, H, may be simply derived from the first, but not from the second, formula given for Al₂(SiO₃)₃, thus O:Si:O₂:Al.O.Si:O
OM, Clarke thinks that the first

formula may be taken to represent Al₂(SiO₃)₃. Several other metasilicates may be formulated as derived from the Al salt by substituting metals or basic groups for one of the AlSiO₃ groups; or many metasilicates containing divalent metals may bem ore simply derived from the normal form M^{II}SiO₃, by doubling this formula and then replacing one M^{II} in the formula M^{II}M^{II}(SiO₃)₂ by other metals or by basic groups.

The species albite and orthoclase, AlMSi,O,, are looked on by Clarke as derived from the

Al salt of the acid H₄Si₂O₈ (-3SiO₂·2H₂O), which acid has probably been isolated, in the same way as many orthosilicates are derived by him from the normal Al salt Al₄(SiO₄)₃. The triclinic felspars are generally thought to consist of mixtures ranging between the limits marked by albite AlNaSi₃O₈, and anorthite Al₂Ca(SiO₄)₂; if these formulæ are trebled and written thus

 $\begin{array}{c} Albite.\\ \text{O.Si}_3\text{O},\text{iNa}_8\\ \text{O.Si}_n\text{O},\text{iAl} & \text{both} & \text{silicates} & \text{are} & \text{repre-}\\ \text{O.Si}_2\text{O},\text{iAl} & \text{o.Si}_2\text{O},\text{iAl} & \text{o.Si}_3\text{O},\text{iAl} & \text{o.Si}_3\text{O},\text{o.Si}_3\text{O},\text{o.Si}_3\text{O},\text{o.Si}_3\text{O},\text{o.Si}_3\text{O},\text{o.Si}_3\text{O},\text{o.Si}_3\text{O},\text{o.Si}_3\text{O},\text{o.Si}_3\text{O},\text{o.Si}_3\text{O},\text{o.$

sented by strictly comparable formulæ. Clarke also draws attention to the fact that the four silicates muscovite 6SiO₂,3Al₂O₃,K,O.H₂O, garnet 3SiO₂,Al₂O₃,3CaO, orthoclase AlKSi₃O₈, and tourmaline 4SiO₂,Al₂O₃,2M₂O₃(or 6MO)B₂O₃, are very frequently associated in granite veins. By halving the formula for muscovite and writing it Al₃KH₂(SiO₄)₃, writing the formula for garnet Al₂Ca₃(SiO₄)₃, trebling the formula for rothoclase and writing it Al₃K₃(Si₃O₈)₃, halving the formula for tourmaline and writing it AlR₂(BO₂)(SiO₄)₂, where R is a metal or group equivalent to Al, very suggestive relationships are disclosed between the four silicates; thus

$$\begin{array}{lll} \textit{Muscovite.} & \textit{Garnet.} \\ \text{Al} \left\{ \begin{matrix} \text{O.Si:O_s:KH_2} \\ \text{O.Si:O_s:Al} \\ \text{O.Si:O_s:Al} \end{matrix} \right. & \text{Al} \left\{ \begin{matrix} \text{O.Si:O_s} \\ \text{O.Si:O_s:Al} \\ \text{O.Si:O_s:Al} \end{matrix} \right\} \\ \textit{Cos.io.p.:Al} \\ \textbf{Al} \left\{ \begin{matrix} \text{O.Si:O_s:R} \\ \text{O.Si:O_s:R} \\ \text{O.Si:O_s:R} \\ \text{O.Si:O_s:R} \end{matrix} \right. & \text{Al} \left\{ \begin{matrix} \text{O.Si_o.p.:K_s} \\ \text{O.Si_o.p.:Al} \\ \text{O.Si_o.p.:Al} \\ \text{O.Si_o.p.:Al} \end{matrix} \right. \end{array}$$

In Am. S. for November 1889 ([3] 38, 384), and A. C. J. May 1893, Clarke has applied the substitution method, as sketched above, to the mica group of silicates, and in the same journal for March 1892 (Am. S. [3] 43, 190) he has applied the same method to the chlorite group.

The micas all fall within the limits of comindicated by the two formulæ position Al₃R¹₅(SiO₄)₅ and AlR₉¹(Si₂O₈)₅. The first formula is derived from Al₄(SiO₄)₅ by substituting Al by one trivalent or three monovalent basic radicles; the second formula is derived from the hypothetical salt Al₄(Si₃O₈)₃ by substituting one, two, or three Al atoms by equivalent basic radicles. When F is present it is regarded as forming part of one or other of the monovalent groups Mg.F or Al:F2. If the ratio of O to Si is greater than 40:Si(SiO,) the excess of O is looked on as combined with metal to form a basic group, such as Al:O; if the ratio of O to Si is less than 40:Si the group Si₃O₃ is supposed to be present. The micas are all capable of being represented as isomorphous mixtures of various constituents, the compositions of which fall within the limiting formulæ.

The chlorites are treated by Clarke (l.c.) similarly to the micas. In Am. S. [3] 43, 198 he gives the following general formulæ which summarise the composition of the chief micas

and chlorites :-

Normal orthosilicate Al₄(SiO₄)₃.

Micas. Muscovite Ala (SiO,) R Normal Biolite Al2(SiO4), Ra. Normal Phlogopite Al (SiO4) Rg. Clintonite RII: O2: Al(SiO4)R3.

Normal orthosilicate Mg, (SiO4)20

Chlorites. $\begin{array}{ll} \textit{Aphrosiderite} \;\; \mathbf{Mg_3(SiO_4)_2R_2}. \\ \textit{Orthochlorites} \;\; \left\{ \begin{array}{ll} \mathbf{Mg_2(SiO_4)_2R_4}. \\ \mathbf{Mg(SiO_4)_2R_6}. \end{array} \right. \end{array}$ Amesite O:Mg2(SiO4)R2.

Preparation of silicates.—Alkali silicates are formed by dissolving moist amorphous SiO, in boiling solutions of alkalis or alkali carbonates, also by fusing SiO₂ with alkalis or with alkali salts of acids which volatilise by heat. Many other silicates are obtainable by double decompositions between alkali silicates (which are sol. water) and metallic salt solutions; also by heating various metallic oxides or carbonates

with SiO₂.

Some naturally occurring silicates have been prepared by the following methods:-1. Emerald Be₂SiO₃, olivine Mg₂SiO₄, and enstatite MgSiO₂, by heating SiO₂ with BeO or MgO mixed with boric acid in a porcelain-oven, till the boric acid volatilised (Ebelmen, A. Ch. [3] 22, 211).—2. Anorthite Al₂Ca(SiO₄)₂, labradorite 3SiO₂.Al₂O₃.K₂O, and oligoclase 9SiO₂.2Al₂O₃.2K₂O, by strongly heating SiO₂ with K2CO, in boric acid, or by adding CaO to molten boric acid and SiO₂ at a very high temperature (E., C. R. 32, 710; Fouquet a. Lévy, C. R. 90, 620).—8. Zirconite ZrSiO₄, and kyanite (AlO)2SiOs, by heating to redness ZrO2 in vapour of H2SiF6, or SiO2 in vapour of AlF2 (Deville a. Caron, A. Ch. [4] 5, 113; Fremy a. Feil, C. R. 85, 1032).—4. Enstatite MgSiO,, and leucite Al2K2(SiO2), by heating Mg, or Al mixed with KOH, to low redness in H and SiCl, vapour (St. Meunier, C. R. 90, 349, 1009).—5. Orthoclase AlKSi₂O₅, albite AlNaSi₂O₅, petalite AlLi(Si₂O₅)₂, by fusing a mixture of Al₂O₄, SiO₂, and tungstate or vanadate of K, Na, or Li; with excess of Al₂O₂ leucite Al₂K₂(SiO₃), is obtained (Hautefeuille, C. R. 90, 878, 541).-6. Orthoclase AlKSi,O,, by heating AlO, and SiO2, in the ratio AlO,:6SiO2, with KVO3 (Hautefeuille a. Perrey, C. R. 107, 786).—7. Phenacite Be3(SiO4) mixed with a lithia felspar, quarts, and tridymits, was obtained by heating for fifteen days at 600°-700° a mixture of 4.8 g. SiO₂, 1.5 g. BeO, 20 g. LiVO₃, and 1.5 g. Li₂CO₃ (H. a. P., C. R. 106, 1800).—8. Crystals of emerald were prepared (H. a. P., C. R. 106, 1800) by heating a mixture of 12.506 g. SiO₂, 3.58 g. Al₂O₃, 2.64 g. BeO, 6 g. Cr₂O₃, and 92 g. acid Li molybdate (ratio = Li₂O:2½MOO₃); the mixture of 12.503 in a Dt cryoible which were ture was placed in a Pt crucible, which was heated in a muffle to low redness for twenty-four hours, when the temperature was raised to 800° and kept thereat for fourteen days. The product was washed with water.

Properties and Reactions of Silicates.—All silicates except those of the alkalis are insol. water; most of them are also insol. dil. acids. For the action of water on certain silicates when the mixtures were rapidly rotated in cylindrical

vessels v. Daubrée (C. R. 64, 339). Many silicates are decomposed by evaporation with HClAq or HNO, Aq, with separation of SiO2.xH2O, which is changed to SiO₂ by heating somewhat above 100°. Some of those silicates which are not thus decomposed are acted on, with separation of SiO₂.xH₂O, by evaporating with 8 parts conc. H₂SO₄ and 3 parts water. Almost all silicates are decomposed by heating with conc. HClAq or H₂SO₄ and a little water in sealed tubes to c. 200°. Heating with conc. HFAq, or in HF gas, decomposes silicates with formation of SiF4; when a silicate is mixed with 3 parts AmF or 5 parts powdered CaF2, and the mixture well moistened with H₂SO₄, and heated in a Pt dish, all the Si is given off as SiF4. Fusion of a finely-powdered silicate with 4-6 parts mixed K2CO3 and Na2CO3 gives K and Na silicate (which is sol. water), while the metal remains as oxide. Silicates which contain F generally lose HF or silicofluoride when strongly heated; any alkali or alkaline metal, or Fe or Al, which was present in the silicate in combination with F remains as oxide (Rammelsberg, W. 7, 146). Clarke a. Schneider (Am. S. [3] 40, 303, 405, 452) have studied quantitatively the interactions of various natural silicates with HCl gas and also with HClAq; their results make it probable that in the magnesian silicates gaseous HCl attacks only the Mg which is present as the basic monovalent radicle Mg.OH.

The following account of individual silicates does not refer to silicates which occur in rocks, but only to those salts which have been prepared in the laboratory; the chief naturally occurring silicates that have been prepared artificially are mentioned under the heading

Preparation of silicates (supra).

Barium silicates. The normal meta- salt BaSiO_{3.}xH₂O is formed by ppg. dil. Na₂SiO₃Aq (v. Sodium silicates, infra) by dil. BaCl₂Aq or Ba(NO₃)₂Aq; not quite insol. hot water, easily sol. dil. HClAq; retains water at 100° (Ammon, J. 1862. 138). Lefort (J. Ph. [3] 39, 31) says that BaO.3SiO₂. 3aq (? BaH₂Si₃O₃. 2aq) is formed by ppg. a dil. Ba salt solution by a dil. solution

of 9SiO,.2Na,O.

Calcium silicates. The compounds 2SiO₂.CaO, 3SiO,.CaO, 3SiO.2CaO. SiO, 2CaO, are said to be produced by very strongly heating mixtures of quartz and marble in the proper proportions. By adding CaCl₂Aq to solution of \$SiO₂.K₂O, a gelatinous pp. is obtained which becomes crystalline after a time; dried at 100°, the solid is said to be 3SiO₂.CaO. 2aq (? CaH₂Si₃O₅.aq). The normal meta-salt CaSiO₅ is said to be formed by ppg. Na₂SiO₅Aq by dil. CaCl₂Aq (v. A., l.c.; L., l.c.). Calcium silicates form the bases of most hydraulic cements (v. Cements in Dictionary of APPLIED CHEMISTRY).

Cerium silicate. The normal meta-salt Ce(SiO₃)₂ is said to be formed by fusing Ce oxychloride with SiO₂ and CaCl₂ or NaCl (Didier, C. R. 101, 882).

Cobalt silicate. The normal ortho- salt Co_2SiO_4 is formed, according to Bourgeois (C. R. 108, 1177), by very strongly heating CoO with CoCl, and a large excess of SiO₂.

Nickel silicate. Normal ortho-salt Ni₂SiO₄

formed like the Co salt (B., l.c.).

The gelatinous pp. Magnesium silicate. obtained by adding MgCl₂Aq to solution of SSiO₂.K₂O is said to be 2SiO₂.MgO.2H₂O (? MgH₂(SiO₃)₂, aq acid meta-salt) (Heldt, J. pr. 94, 129, 157). The normal meta-salt MgSiO₃.xH₂O is obtained, according to Ammon (l.c.), by ppg. Na₂SiO₃Aq by dil. MgCl₂Aq.
Potassium silicates. The normal meta- salt

K₂SiO₃ is prepared by fusing 1 part SiO₂ with 21 parts dry K2CO3. If this product is dissolved in a little water, and alcohol is added, a gelatinous pp. is formed, which, if slightly washed and squeezed, is said to be 4SiO2.K2O; but Fremy (J. 1856. 353) gives the composition 9SiO_{2.2}K₂O aq to this pp., and this is confirmed

by Lefort (J. 1861. 205).

The name potash water-glass is given to the mixture of K silicates obtained by fusing together quartz and pearlash in a proportion about equal to 4SiO2.K2O. Water-glass is a clear transparent solid, like glass; it is completely sol. hot water; a conc. solution, with c. 28 p.c. silicate, is syrupy, somewhat turbid, and has S.G. 1.25. The solution is decomposed by acids, even by CO2, with separation of S_1O_2 xH_2O ; alkaline carbonates and chlorides, especially AmCl, ppt. SiO2xH2O; BaO, SrO, CaO, Al₂O₃, and PbO decompose the solution, forming a double silicate with the whole of the silica and a portion of the K. The solution yields pps. with solutions of most of the salts of earth metals and heavy metals. Water-glass is also formed

by boiling SiO₂ with KOHAq under pressure.
Sodium silicates. The normal meta-salt Na2SiO3 is prepared by fusing 1 part SiO2 with 13 parts dry Na₂CO₃. The salt is obtained with 8H₂O, in monoclinic crystals, by dissolving SiO₂ in an equivalent quantity of NaOH in solution, decanting, evaporating, as much as possible out of contact with air, till Na2CO3 begins to settle out, cooling to -22° , stirring till the whole solidifies, straining from motherliquor, dissolving in a little water, and allowing to stand (Ammon, J. 1862. 138; cf. Ordway, Am. S. [2] 40, 186). By adding freshly ppd. AlO₂H₂ to a boiling solution of Na₂SiO₂ and NaOH, the double salt Na₂SiO₃.Al₄(SiO₄)₃ is obtained; and the salt Na, SiO, Al, (SiO,) is ppd. by adding Na2SiO3Aq to a hot solution of alum (A., l.c.). Mixtures of Na silicates are known as soda water-glass (cf. supra, Potash water-glass). For effect of time on composition of solutions of Na silicates v. Kohlrausch (Z. P. C. 12, 773).

Strontium silicate. The normal meta-salt SrSiO₃ is said to be formed by adding

Na₂SiO₃Aq to dil. SrCl₂Aq (Ammon, l.c.).
Thorium silicates. The normal ortho-salt ThSiO, and the normal meta-salt Th(SiOs)2 are formed, according to Troost a. Ouvrard (C. R. 105, 225), by fusing ThO₂ with SiO₂ and CaCl₃ and treating with dil. HClAq. ThSiO₄ is not isomorphous with ZrSiO₄ (T. a. O., *l.c.*).

Zinc silicates. The normal meta-salt ZnSiO, is obtained in rhombic crystals by ppg. ZnSO, Aq by Na₂SiO₂Aq, and strongly heating the dried pp. with boric acid for several days (H. Traube, B.

26, 2735).

A systematic arrangement of the naturally occurring silicates will be found in Groth's Tabell. Uebersichte, pp. 103-151 [3rd ed.].

M. M. P. M.

SILICIC ACIDS v. SILICA, HYDRATES OF. p. 447)

SILICIDES. Binary compounds of Si with metals. These compounds are formed (1) by heating amorphous Si with metallic chlorides in presence of such a reducer as Na; (2) by heating K2SiF6 with metals; (3) by electrolysing a mixture of an alkali fluoride with a metallic oxide; (4) by heating metals with Si (v. Aluminium, CERIUM, COPPER, IRON, MAGNESIUM, MANGANESE, NICKEL, PLATINUM, SILICIDES OF). M. M. P. M.

SILICO-BORATES. Certain minerals consist of compounds of silicates and borates; the name silico-borate is sometimes applied to such

0:\$i.0 O:Si.O Ca.CaB₂O₄. aq, compounds. **Datholite**

and botryolite, the same formula with 2aq, are examples of borosilicates. Tourmalines Al(BO₂)(SiO₄)₂R₁ are also silico-borates (R₁ may be replaced by its equivalent of R11).

gen bromide. Tribromo-silico-methane.) drogen bromide. Obtained, not quite pure, by Buff and Wöhler (A. 104, 99) by leading HBr gas over heated Si, distilling the product, shaking with Hg, and distilling again. Gattermann (B. 22, 193) obtained pure SiHBr, as follows. Crude Mg tained pure SiHBr, as follows. Crude Mg silicide, prepared as described under Silicon, Preparation, No. 1 (p. 456), is freed from MgO by powdering, adding little by little to HClAq (1:2), shaking with this acid for some hours, washing till the wash-water is neutral, and drying thoroughly. The Si thus obtained is heated in a tube in a stream of HBr, the tube being connected with a well-cooled receiver; the details and precautions described under Silicon TETRACHLORIDE, Preparation, No. 2 (p. 458) must be observed. The HBr required may be made by the action of H2SO4Aq on KBr; G. used a cold mixture of 45 c.c. conc. H₂SO, and 15 c.c. water, and let this drop on to 75 g. KBr; he dried the gas by H₂SO₄. The liquid is KBr; he dried the gas by H₂SO. The liquid is distilled, and then fractionated till a portion boils at 115°-117°; most of the remainder boils at c. 153° and is SiBr. A colourless liquid, boiling 115°-117°; S.G. 2.7 at ordinary tem-Fumes much in air, and perature (G., l.c.). takes fire, probably because the heat produced in the decomposition of a portion by the moisture of the air suffices to start rapid oxidation Ice-cold water forms silico-formic (G., l.c.). anhydride (q. v., p. 455). Combines with PH, when compressed, to form a solid, unstable compound (Besson, C. R. 112, 530).

M. M. P. M. SiHCl_s. (Silicon SILICO-CHLOROFORM hydrogen chloride. Trichloro-silico-methane.) Mol. w. 135.41. V.D. 67.

Formation.—1. Dry HCl gas is passed over Si heated below redness; the product is condensed in a flask in a freezing mixture and fractionated (Friedel a. Ladenburg, A. 143, 118; cf. Buff a. Wöhler, A. 104, 94).—2. Dry HCl gas is passed over siliceous pig-iron containing 15 p.c. Si (Warren, C. N. 60, 158).

Preparation.—Si, prepared by reducing sand

by Mg (v. Silicon, Preparation, No. 1, p. 456) and removing MgO as described under Sunco-BROMOFORM (supra), is carefully heated in a stream of dry HCl gas in a tube connected with a receiver surrounded by a very good freezing mixture. The directions given for preparing SiCl₄ must be carefully followed (v. Silzoon Tetrachloride, Preparation, No. 2). [Gattermann recommends to make the HCl in a Kipp's generator, using conc. H₂SO₄ and fused NH₂Cl.] The liquid in the receiver is fractionated from a flask with a side tube 75 cms. long; if this is done, no condenser is required; the flask must be warmed by water at 90°, if a naked flame is used explosions of part of the SiHCl₅ may result (Gattermann, B. 22, 190).

Properties and Reactions.—A colourless, mobile liquid, with a very disagreeable smell; boils at 35°-39° (F. a. L., l.c.). Non-conductor

of electricity.

The vapour takes fire very easily in air, e.g. by bringing a hot glass rod to the mouth of a vessel in which a little SiHCl₂ has been shaken. Mixture of the vapour with air is explosive. Decomposed to Si and HCl by passing through a narrow tube heated to redness. Water at 0° forms Si₂O₃H₂ (v. Silicoformic annutration, p. 455) and HCl; at the ordinary temperature, SiO₂xH₂O, HClAq, and H are produced. Chloring reacts at ordinary temperatures, forming SiCl₄ and HCl; bromine reacts similarly at 100°. Alcohol produces SiH(OEt)₂ (v. Silicon tetra-hydride, Preparation, No. 1, p. 460). Combines with phosphorus hydride, under pressure, to form a solid unstable body (Besson, C. R. 112, 580).

M. M. P. M.

SILICO-ETHANE, PENTA-BROMO- DERI-VATIVE OF, Si, HBr₅. (Distilicon hydrogen pentabromide. Pentabromo-silico-ethane.) In making SiBr, by passing SiH, into Br, Mahn (J. Z. 5, 163) obtained a solid in the form of white needles, melting at 89° and boiling (out of contact with air) at 230°, and giving numbers on analysis agreeing with the formula Si₂HBr₅. Takes fire when heated in air. M. M. P. M.

SILICO-FLUORHYDRIC ACID H. SiF. Aq. (Hydroftuosilicic acid.) Known only in aqueous

solution.

Preparation.—SiF₄, prepared by the interaction of CaF₂, SiO₂, and H₂SO₄ (v. Sillicon TETRAFLUORIDE, p. 459), is passed into water, the exit tube dipping beneath a little Hg in the bottom of the vessel which contains the water; the vessel is shaken frequently to break up the gelatinous silica which forms in the water; when the liquid becomes thickish from separated silica, it is filtered through linen, the silica is pressed, and more SiF₄ is passed into the filtrate, as before.

SiF₄+3H₂O+Aq = SiO_xH₂O+4HFAq;
 4HFAq+2SiF₄=2H₂SiF₆Aq. The solution of H₂SiF₆ may be evaporated in a Pt vessel till it contains c. 34 p.c. of the acid.

To prepare the acid solution on a large scale, Tessié du Mothay recommends to fuse a mixture of fluorspar, silica, and charcoal in an oven, and to lead the gas that is given off through condensers containing water; c. 68 p.c. of the F in the fluorspar is thus obtained as H₂SiF₆ (C. C. 1868, 432).

Properties and Reactions.—H.SiF.Aq is a strongly-acid liquid, smelling like HClAq, and burning the skin. If kept in glass vessels it gradually withdraws alkali, lime, and iron

oxide (H. Rose, P. 80, 403). Truchot (C. R. 98, 821) gives H.F. [SiF',2HFAq] = 17,000 (cf. Sabatier, A. Ch. [5] 22, 91). Thomsen gives [SiO'Aq,6HFAq] = 32,800; and the heat of neutralisation as [H'SiF'Aq,2NaOHAq] = 26,600 (Th. 1, 236). The following table is given by Stolba (J. pr. 90, 193):—

s.G.	P.c.	8.G.	P.c.
H,SiF,Aq	H.SiF.	H_SiF_Aq	H.SiF.
1.0040	0.5	1.1559	18
1.0080	1	1.1653	19
1.0161	2	1.1748	20
1.0242	8	1.1844	21
1.0324	4	1.1941	22
1.0407	5	1.2038	23
1.0491	6	1.2136	24
1.0576	7	1.2235	25
1.0661	8	1.2335	26
1.0747	9	1.2436	27
1.0834	10	1.2537	28
1.0922	11	1.2639	29
1.1011	12	1.2742	30
1.1100	13	1.2846	31
1·1190	14	1.2951	32
1.1281	15	1.3056	33
1.1373	16	1.3162	34
1.1466	17		

The S.G. for any percentage of H_2SiF_6 , when n = number of half-per-cents, is given by the formula; S.G. = $1 + n \cdot 004 + \frac{n(n-3) + 2}{100000}$.

By passing SiF₄ into fairly conc. HFAq until the liquid is saturated, hard, colourless, very hygroscopic crystals of the *hydrate* H₂SiF₄.2H₂O separate; the crystals melt at 19°, and decompose at a higher temperature (Kessler, C. R. 90, 1285).

H₂SiF_eAq acts as a dibasic acid (the heat of neutralisation confirms this), giving salts M₂SiF_e and M¹¹SiF_s; with excess of a strongly basic metallic oxide it forms metallic fluoride and SiO₂·H₂O—thus

 $H_2SiF_6Aq + 3K_2O = SiO_2.H_2O + 6KFAq.$

H₂SiF_eAq is decomposed by conc. \hat{H}_2 SO₄, also by HCl gas, with formation of SiF₄ and withdrawal of water. Boric acid separates SiO₂ $_{\infty}$ H₂O and forms fluoboric acid (q. v., vol. i. p. 530). M. M. P. M.

SILICO-FLUORIDES. (Hydrofluosilicates.) Salts of H2SiF6 (v. preceding article). These salts are formed by dissolving various metalse.g. Fe or Zn-in H2SiF6Aq, or by neutralising the acid solution by the proper quantities of basic oxides or hydroxides; if excess of a strongly basic oxide is added to H2SiF6Aq, SiO2.H2O is separated and a fluoride is formed. Most of the silicofluorides are sol. water; the salts of Ba, K, Na, Ce, and a few elements allied thereto, are only slightly sol. water and insol. alcohol. Aqueous solutions of silicofluorides have an acid reaction to litmus. Silicofluorides are generally decomposed by heat, giving off SiF, and leaving fluorides; heated with AmCl, metallic chlorides remain (Stolba, J. pr. 101, 157). Some silicofluorides are de composed by heating strongly with Fe or K, forming fluoride and Si. Conc. H2SO4 evolves H₂SiF₆, when heated HF is given off; conc. HClAq and HNO, Aq react similarly, but more slowly and less completely; but H₂SiF_{*}Aq SILICON. 455

partially decomposes most chlorides and nitrates in solution. Truchot (C. R. 98, 821, 1330) gives the heats of formation of alkali silicofluorides.

M. M. P. M. SILICO-FORMIC ACID. By the action of water on (?) SiL2, Friedel a. Ladenburg (A. 203, 247) obtained a white solid that gave off H when treated with KOHAq; the amount of H obtained agreed fairly well with the formula SiH₂O₂. If this formula is correct, the compound is probably the Si analogue of formic M. M. P. M.

SILICO-FORMIC ANHYDRIDE H,Si,O, HSi:0

This body represents among Si com-HŚi:O.

pounds the hypothetical formic anhydride HÇ:0

Prepared by slowly distilling SiHCl, HĊ:O.

into water kept at 0°, filtering quickly the solid that separates, washing with ice-cold water, drying in vacuo over H₂SO₄ and then at 150°; the SiHCl, is distilled from a small flask and the distillation-tube is fused on to an inverted funnel which dips beneath the water; $2SiHCl_8 + 3H_2O = 6HCl + Si_2H_2O_3$ white, amorphous, light powder; floats in water, sinks in ether; sl. sol. water. Does not decompose at 300°, but at higher temperatures it glows and gives off H which takes fire explosively; burns when heated in O, emitting a brilliant light; burns when heated in a covered crucible, with formation of some amorphous Si. Heated in tube, SiH, is given off and a thin layer of amorphous Si remains. Not acted on by acids, except HFAq which dissolves it with rapid evolution of H. Alkali and alkali carbonate solutions, including NH₃Aq, dissolve this compound, giving off H, and forming alkali silicates. A freshly-prepared aqueous solution of H₂Si₂O₃ acts as an energetic reducer; AuClaAq is reduced to Au; Pd salts probably to Pd mixed with Pd silicate; SeO₂Aq, SO₂Aq, TeO₂Aq, and HgCl₂Aq are reduced to Se, S, Te, and HgCl. CrOsAq, indigo solution, and salts of Pt and Ir are not reduced.

The substance obtained by Wöhler, by the action of light and water on the body he named silicone (v. Silicon, compounds of, with hydrogen and oxygen, p. 459), and called by him leukon (A. 127, 268), and also the substance called by Geuther silicium oxide (J. pr. 95, 424), are probably identical with Si₂H₂O₃. M. M. P. M. SILICO-IODOFORM SiHI₃. (Silicon hy-

drogen iodide. Tri-iodo-silico-methane.) mula probably molecular, from analogy of SiHCla. Obtained in small quantities by the interaction of HI and crystalline Si at red heat (Buff a. Wöhler, A. 104, 99); better prepared by diluting the HI vapour with H; the product, which is a mixture of SiI, and SiHI, is repeatedly fractionated; SiI, boils at 290°, SiHI, at c. 200°. A mixture of Sil, and SiHI, is also obtained by heating SiH₄ with I (Mahn, J. 1869. 248). colourless, refractive liquid; S.G. 3.362 at 0°, 3.314 at 20°; boils at c. 200°. Water produces Si₂H₂O₃ (Friedel, A. 149, 96) (v. Silico-formio M. M. P. M. ANHYDRIDE, supra).

SILICO-MOLYBDATES. Compounds containing SiO₂, MoO₂, and basic radicles; v. MoLYB-DATES, vol. iii. p. 427.

SILICON. Si. (Silicium.) At.w. 28.3; mol.w. known. H.F. $[Si, O^2] = 219,240$ amorphous unknown. Si; 211,120 crystalline Si (Troost a. Hautefeuille, C. R. 70, 252). C.E. (linear at 40°) ·00000763 (Fizeau, C. R. 68, 1125). For other

properties v. Properties, p. 456.

Historical.—The existence of a special earth in rocks that could be melted to glass-like substances was indicated by Pott in 1746. Scheele (Opuscula, 2, 67) and Bergmann (Opuscula, 2, 26) showed that this earth could not be changed into lime or alumina; Smithson in 1811 found that the earth acted like an acid; Berzelius, in 1823, isolated the element of this earth, and in 1854 Deville obtained the same element in crystalline form. The new element was shown to resemble carbon by the researches of Buff and Wöhler and of Friedel and Ladenburg. The name silicon was given from acidum silicum or silex.

Occurrence .-- Si is not found uncombined; compounds of Si are very widely distributed in vast quantities; next to oxygen, Si (in combination) is the most widely distributed element. Silicates occur in very many rocks and soils and in plant-ashes; SiO2 is found in many mineral springs and in sea-water (Forchhammer, Pr. E. 2, 303; Bunsen, A. 62, 7, 25), and in small quantities in various animal organisms (von Gorup-Besanez, A. 61, 46; Henneberg, A. 61, 261). Si is found in pig-iron; it is generally supposed to exist therein as crystallised Si, but the experiments of Jordan a. Turner (C. J. 49, 215) make it probable that the Si is combined with Fe as a silicide.

Formation.—Amorphous silicon.—1. By passing the vapour of SiCl, over heated K, removing excess of SiCl, by a current of dry air, and washing out KCl with water (Berzelius. Lehrb. (1st ed.) 3, 327).-2. By passing vapour of H₂SiF, over neared A, washing the residue in a covered crucible (to heating the residue in a covered crucible (to weeking with HFAq (to of H2SiF, over heated K, washing with water, remove H and C), washing with HFAq (to remove SiO₂) and then with water (B., l.c.).—3. By heating a mixture of K₂SiF_q and K, and washing with water (B., l.c.); Wöhler (A. 104, 107) heats a mixture of Na2SiF, with NaCl and Na.-4. By heating Mg in SiF, vapour (Warren, C. N. 58, 215).-5. By reducing SiO₂ by heating with Mg (v. Preparation, No. 1).-6. By elec trolysing a fused silicate in a carbon crucible, with Pt as the positive, and gas-carbon as the negative, electrode (Hampe, Chem. Zeitung, 12, 841). Crystalline silicon.—7. By heating a mixture of K₂SiF₆, Na, and Zn (v. Preparation, No. 2).—8. By reducing K₂SiF₅ or a silicate by Al (v. Preparation, No. 3) .- 9. By passing H and SiCl, vapour over Zn heated to redness in a porcelain tube (Beketoff, Bl. 1, 22).—10. By heating a mixture of SiO₂, K₂CO₃, Iceland spar, and Na (Deville, A. Ch. [8] 49, 62).—11. By passing vapour of SiF, or SiCl, over Si heated to redness (Troost a. Hautefeuille, C. R. 73, 443).-12. By passing vapour of SiCl, over heated Na, or Al, and heating the product (Si with excess of Na or Al) in a carbon crucible (Deville, A. Ch. [8] 49, 62).—13. A piece of siliceous pig-iron is con nected with the positive pole of a battery and 456 SILICON.

immersed in dilute H₂SO₄Aq, in which a Pt plate connected with the negative pole of the battery is also placed; after some hours the iron has dissolved; the mixture of graphite, amorphous Si and SiO₂ which remains is heated to full redness for some time with Zn and the fused mass is treated with dilute HClAq (Warren, C. N. 57, 54).

Preparation. - Amorphous silicon. 1. An intimate mixture of 10 g. Mg powder and 40 g. thoroughly dry sand is placed in a testtube, of fairly thick glass, c. 2-3 cms. diameter and c. 15 cms. long; the tube is heated throughout by a large flame, and then the lower part is very strongly heated, when reduction quickly occurs. If the tube is gradually moved downwards so that one part is strongly heated after another, the whole of the SiO_2 is reduced in a few minutes. The contents of the tube are shaken out, pulverised, and treated with HClAq (1:2), the solid being added to the acid in small successive quantities (to prevent explosion from evolution of SiH,) (Gattermann, B. 22, 186). Crystalline silicon.—2. A mixture of 15 parts dry K₂SiF₆, 20 parts dry granulated Zn, and 4 parts dry Na în small pieces is placed in a clay crucible, which is covered and heated to redness; when the reduction of the K2SiF6 is effected (there is a visible, but not violent, reaction) the temperature is raised till the mass melts, care being taken not to heat to the b.p. of zinc, else considerable loss will occur. The crucible is broken when cold, the regulus is heated to the m.p. of zinc, the molten zinc is poured off, and SiO2, Zn, &c., are removed by treatment, in succession, with conc. HClAq, conc. boiling HNO, Aq, and The Si, mixed with a little K.S.F. thus obtained is placed in a crucible and covered with a layer of powdered glass; the trucible is set inside another, and heated to the m.p. of pig-iron (c. 1100°). When all is melted the crucible is allowed to cool to dull redness and is then plunged into water; the regulus is separated from the slag, and is treated with HFAq and then washed with water (Caron, A. Ch. [3] 63, 26; Deville a. C., A. Ch. [3] 67, 435).—3. A mixture of 1 part Al with 20-40 parts thoroughly dry K₂SiF₆ or Na₂SiF₆ is heated to c. 950° in a Hessian crucible, and the contents are kept molten for c. a quarter of an hour; the crucible is broken when cold, the regulus is separated, crushed, heated with conc. HClAq as long as H is given off, then with HFAq, and washed with water and dried (Wöhler, A. 97, 261). Or, a mixture of 10 parts powdered cryolite and 5 parts K2SiO2 or powdered glass is divided into two equal parts; one part is placed in a Hessian crucible, 1 part Al is added, the rest of the mixture is placed over this, and the whole is heated to redness for half an hour. The regulus is treated with conc. HClAq, HNO, Aq, &c., as directed above.

Properties.—Si exists as a brown amorphous powder, and also as greyish-black, very lustrous, octahedral crystals.

Amorphous silicon is a lustrous, brown powder, which adheres to the fingers or to glass tenaciously. The S.G. has not been determined, but the powder is heavier than oil of vitriol; it is a non-conductor of electricity; burns when

heated in air or O, but the SiO, formed soon stops the process; dissolves in cold HFAq, forming SiF, and H; also dissolves in warm alkali solutions; is oxidised with incandescence when thrown into molten nitre; combines with S when warmed therewith. After being strongly heated in a covered Pt crucible, and the SiO, formed has been removed by HFAq, amorphous Si is a dark chocolate-brown powder, which is not burnt by heating in air or O, is insol. HFAq and boiling alkali solutions, does not combine with S, and is not acted on by molten KNO, or KClO,

Crystalline silicon is obtained in leaflets or needles. The leaflets are opaque, very lustrous, metal-like, greyish black, resembling graphite; they consist of regular octahedra. The needles are also octahedral. Crystalline Si has S.G. 2.49 at 10° (Wohler, J. 9, 437); 2.194 to 2.197 (Winkler, J. 17, 208). Scratches glass, but not topaz. According to Deville (C. R.39, 321) crystalline Si 'conducts electricity like graphite.' M.P. between those of pig-iron and steel, i.e. between c.1100° and c.1300°. Crystalline Si is not changed when heated, even to whiteness, in O; it is unacted on by HFAq; dissolves in warm alkali solutions; combines with S vapour. When heated in the electric arc crystalline Si melts and then boils, and the ends of the electrodes become covered with crystals of SiC (Moissan, C. R. 117, 423).

For the lines in the emission spectrum of Si v. B. A. 1884. 441.

Deville (A. Ch. [3] 49, 70) thought that a graphite-like form of Si existed different from the ordinary crystalline (or diamond-like) form; but Miller (P. M. [3] 31, 397) showed that both the leaflets and the needles consisted of octahedra. Kopp (A. Suppl. 5, 72) found differences between the specific heats of crystalline Si and the graphite-like form prepared by the interaction of K,SiF, and Al, and these differences were confirmed by Winkler (J. pr. 91, 198).

Warren (C. N. 63, 46) thought he had obtained a crystalline form of Si (oblique octahedra) different from the ordinary by the action of impure Al on K₂SiF₆ at a very high temperature; he described the crystals as very perfect, and sometimes half-an-inch across the faces.

Specific heat of silicon. Kopp, Regnault, and others obtained values for the S.H. of Si varying from '138 to '173 at c. 30°-100°. In 1875 Weber (P. M. [4] 49, 161, 276) showed that S.H. of crystallised Si increased rapidly from -40° to c. 200°, and attained an almost constant value at the latter temperature. Weber's results gave the following values for S.H. of crystallised Si:-

Temp.	8.H.
- 40°	•136
+ 57	·183 3
128	·196
184	•2011
232	.2029

The atomic weight of Si has been determined (1) by finding the ratio of Si to SiO₂ formed therefrom (Berzelius, P. 1, 226 [1824]); (2) by converting BaSiF₄ into BaSO₄ (B., P. 8, 20

[1828]); (3) by ppg. Cl in SiCl, by Ag (Pelouze, C. R. 20, 1047 [1845]; Dumas, A. Ch. [8] 55, 183 [1859]); (4) by converting SiCl, into AgOl (Schiel, A. 120, 94 [1861]); (5) by decomposing SiBr, by water and determining SiO, produced (Thorpe a. Young, C. J. 51, 576 [1887]); (6) by finding V.D. of SiH, SiCl, SiBr, SiF, &c.; (7) by determining S.H. of Si. The atom of Si is tetravalent in the gaseous molecules SiH, SiCl, SiBr, SiF, SiF, &c.; Acc.; (7) by determining S.H. of Si. The atom of Si is tetravalent in the gaseous molecules SiH, weight of Si is unknown.

Si is closely related to C, and less closely to Ti, Ge, Zr, Sn, Ce, Pb, and Th (v. Carbon Group

OF ELEMENTS, vol. i. p. 682).

Reactions and Combinations.—I. Amorphous silicon which has not been heated.—1. Heated in air or oxygen burns to SiO₂, which coats the Si, so that the process soon stops.—2. Heated with sulphur forms SiS, (Berzelius) .- 3. In molten nitre forms potassium silicate, with incandescence.—4. Dissolves in cold solution of fluorhydric acid to form SiF, and H.-5. Dissolves in conc. alkali solutions on warming, forming H₂SiO₃Aq (or Na₂SiO₃Aq) and H. For differences between amorphous silicon before and after heating, v. Properties of amorphous silicon, p. 456. -II. Crystalline silicon.—6. Is not acted on by oxygen, nor by fluorhydric acid. -7. Heated in dry chlorine forms SiCl,; in bromine vapour forms SiBr,; in a mixture of iodine vapour and CO₂ forms SiI₄; takes fire in fluorine, forming SiF₄ (Moissan, C. R. 103, 256).—8. Heated with sulphur vapour forms SiS₂.—9. Strongly heated in nitrogen forms Si, N, .- 10. Heated to c. 800° in a mixture of oxygen and chlorine, Si₂OCl_a is formed (Troost a. Hautefeuille, Bl. [2] 35, 360). 11. Dissolves slowly in warm alkali solutions of medium concentration, giving off H.—12. Burns when heated with alkali carbonates, separating C, and giving off CO.—13. SiS₂ is produced (according to Sabatier, Bl. [2] 38, 153) by heating to bright redness in hydrogen sulphide .-14. Schutzenberger (C. R. 114, 1089) obtained a mixture of Si₃N₄ and a carbide (to which he gives the formula SiC) by heating to bright redness, in a carbon crucible placed inside another crucible packed with lamp-black, a mixture of 1 part Si and 2 parts SiO2.—The following reactions apply generally to silicon.—15. Heated in hydrogen chloride SiHCl, is formed; in hydrogen iodide mixed with H, SiHI, is produced; and SiHBr, is obtained by heating in hydrogen bromide.—16. Heated with silicon tetrachloride, Si₂Cl₂ (and ?SiCl₂) is formed.— 17. Oxidised by hydrogen iodide solution (v. Ditte, Bl. [2] 13, 322).—18. With molten caustic soda or potash, H is given off and a silicate formed; similar but slower reactions occur with baryta and lime (v. Berzelius, A. 49, 247).-19. Heated with several metallic oxides, such as PbO or Ag₂O, reduction occurs, generally with formation of a silicide. If a mixture of powdered Si, Al, and PbO is heated, a violent explosion occurs.—20. According to Colson (Bl. [2] 38, 56), Si₂C₂O is formed by heating Si in carbon dioxide; SiS, SiSO, and Si,C,S are formed by heating Si to white heat in carbon disulphide SiCO, is formed by heating with bensene; and Si₂C₂O₂ is produced when Si is heated to whiteness in a carbon orucible.

Silicon, amidonitride of. By the interaction of NH, and SiCl, or SiF, Harris (C. C. 1889 [ii.] 283) obtained a snow-white powder, to which he gave the formula Si.NH₂.N.

Silicon, carbide of. According to Schutzen-berger (C. R. 114, 1089), a compound of Si and C, having the composition of SiC, is formed along with a little Si₃N₄, by placing a mixture of 1 part crystallised Si and 2 parts SiO₂ in a covered gas-carbon crucible, imbedding this in lamp-black in a larger crucible, and this again in more lamp-black in another larger crucible, and heating to bright redness for some hours; boiling with moderately conc. HFAq dissolves SiO₂ and Si₃N₄, and leaves the SiC. Heated to low redness in chlorine, SiC is said to give SiCl, and C. Moissan (C. R. 117, 425) obtained colourless crystals of SiC by fusing C and Si in an electric furnace. The crystals are very hard, act strongly on polarised light, S.G. 3.12; they are not acted on by O or S vapour at 1000°; Cl begins to react at c. 600° and the action is complete at 1200°; the crystals are not acted on by boiling HClAq, HNO₃, H₂SO₄, aqua regia, or a mixture of HNO, and HFAq, nor by fused KNO₃ or KClO₃; molten KOH gradually forms K2CO3 and K silicate.

Silicon, bromides of. Two bromides of Si

are known, Si₂Br_s and SiBr₄

DISILICON HEABROMIDE Si₂Br_e. (Silicon tribromide SiBr_s.) Formula probably molecular from analogy of Si₂Cl_e. A colourless liquid, boiling at c. 240°. Formed by adding the proper quantity of Br to Si₂I₆ in CS₂, pouring off from I, purifying by shaking with Hg, filtering in dry air, and fractionating (Friedel a. Ladenburg, A. 203, 254).

SILICON TETRABROMIDE SiBr. Mol. w. pro-

bably 347.3.

Preparation.—Amorphous Si, prepared as described under Silicon, Preparation, No. 1 (p. 456), after being partially purified from SiO₂ by HClAq, washing, and drying, is heated in a glass tube, while Br vapour is passed over it; the tube passes into a small flask, which is surrounded by cold water. A mixture of SiBr₄ and Br collects in the flask; this is fractionated, and the last traces of Br are removed by shaking with Hg and distilling (Gattermann, B. 22, 186). SiBr₄ may also be prepared by heating a mixture of SiO₂ and O (cf. Silicon tetrachloride, p. 458) in Br, shaking with Hg, and fractionating (Serullas, P. 24, 341; modified by Reynolds, C. J. 51, 590).

Properties and Reactions.— A colourless liquid, having a disagreeable odour, and fuming much in air. S.G. 2.8128 at 0°. B.p. 153.4° at 762.5 mm. (Pierre, A. Ch. [3] 20, 26; Freyer a. Meyer, Zeit. für anorgan. Chemie, 2, 1); 148° to 150° (Serullas, l.c.). Solidifies at c. -12°. (F. a. L., l.c.). Decomposed by water to HBrAq and SiO₂; shaken with conc. sulphuric acid, slowly changes to Br and SiO₂. Heated to 250° with lead oxide gives PbBr₂ and Pb silicate (Friedel a. Ladenburg, A. 147, 362). Ammonia probably forms SiBr₁.NH₄ (Persoz, A. Ch. [2] 44, 315). Phosphorus hydride forms a white amorphous compound, by repeated compression with SiBr₁, the pressure being maintained for some hours (Besson, C. R. 110, 240).

For the compounds SiBraH and SizBraH v.

SILICOBROMOFORM, and SILICO-ETHANE, PENTABBOMO- DERIVATIVE OF, pp. 453, 454.

Silicon, bromochlorides of. Three compounds of Si with Br and Cl have been isolated. The V.D. of SiBr,Cl, and also that of SiBrCl₂, has been determined; the formulæ of all are probably molecular.

Šilicon uromo-trichloride SiBrCl₂. Mol. w. 214·16. A colourless liquid, boiling at c. 80°, fuming in air, and decomposed by water. Prepared by heating Br and SiHCl₂ at 100°; SiHCl₂+Br₂=SiBrCl₃+HBr (Friedel a. Ladenburg, A. 145, 187). Also by the interaction of Br and SiCl₃SH (? SiCl₂SH+3Br=SiBrCl₃+SBr+HBr) (F. a. L., l.c. p. 179). V.D. 104·7 at c. 130° (F. a. L., l.c.). Reacts with NH₃ to form an amorphous compound, 2SiBrCl₂-11NH₃, easily decomposed by water (Besson, C. R. 112, 788).

SILICON DIBROMO-DICHLORIDE SiBr₂Cl₂. A colourless liquid, boiling at 103°-105°, and not solidifying at -60°. Prepared by heating SiHCl₂ and Br above 100° (F. a. L., *l.c.*). Also by passing the vapour of HBr and SiCl₄ through a red-hot porcelain tube (Besson, *l.c.*). Combines with NH₃ to form SiBr₂Cl₂.5NH₃, decom-

posed by water.

SILICON TRIBROMO-CHLORIDE SiBr₃Cl. Mol. w. 302-92. A colourless liquid, fuming in air, boiling at 140°-141°, not solidifying at -40°; S.G. 2·432. V.D. 150·5 at c. 185° (Reynolds, C. J. 51, 590). Obtained, along with SBr₄, by passing Br vapour over a mixture of SiO₂ and C heated in a wind-furnace, passing dry H through the warm product, shaking with Hg, and fractionating (R., l.c.). Besson (C. R. 112, 788) seems to have obtained the same compound, along with SiBr₂Cl₂, by passing the vapours of HBr and SiCl₄ through a redhot porcelain tube. B. gives the b.p. 126°-128°, and m.p. -39°; he says a compound SiBr₃Cl.11NH₃ is formed by interaction with NH₃.

Silicon, bromo-iodides of, v. SILICON 10DO-

BROMIDES, p. 461.

Silicon, chlorides of. Two chlorides of Si have been isolated with certainty, Si₂Cl₆ and SiCl₄; these formulæ are molecular. There are indications of the existence of another chloride, SiCl₆.

SILICON HEXACHLORIDE Si₂Cl₈. (Silicon tri-

chloride SiCl₃.) Mol. w. 268-22.

Preparation.—1. By heating HgCl₂ with Si_I_s, fractionating, and distilling the portion boiling at 146°-148° from HgCl₂ (Friedel a. Ladenburg, A. 203, 253).—2. Vapour of SiCl₄ is passed over Si kept molten in a porcelain tube (3SiCl₄+Si=2Si₂Cl₆); the product is rapidly cooled, and fractionated from SiCl₄, Si oxychlorides (and? SiCl₂) (Troost a. Hautefeuille, A. Ch. [5] 7, 459).

Properties. — A colourless, mobile liquid, boiling at 146°-148°; S.G. 158 at 0°; solidifies at -14° to large leaflets, which melt at 1° (F. a. L., l.c.). V.D. 140 at c. 240° (T. a. H., l.c.). Fumes in air, decomposed by water; vapour takes fire when Si_oCl_a is heated in air.

takes fire when Si_cCl_s is heated in air.

Reactions and Combinations.—1. Decomposed to SiCl_s and Si by heat in a closed tube; decomposition is extremely slow at 350°, nearly complete at 800°; heated rapidly above 1000°

the dissociation-pressure falls; the compound is stable below 350° and above 1000° (T. a. H., l.c.).—2. Decomposed by water. At ordinary temperatures products remain in solution, and are ppd. by NH₂Aq; at 0° Si₂H₂O₄ is formed (v. Silico-oxalio acid), p. 463) (F. a. L., l.c.).—3. Caustic potash forms SiO₂ and gives off H (F. a. L., A. 203, 254).—4. Phosphorus hydride is changed to the solid hydride at -10° (Besson, C. R. 110, 516).—5. Combines with ammonia to form Si₂Cl₆.5NH₃ (B., l.c.).

SILICON TETRACHLORIDE SICI. Mol. w. 169.78. Boils at 57.57° at 760 mm. pressure (Thorpe, C. J. 37, 327). S.G. % 1.52408 (T., l.c.). V.D. 85.5 at 100° (Dumas, A. Ch. [2] 33, 368). S.H. of SiCI, vapour, at 90° to 234°, = 1322, at constant pressure, referred to equal weight of water; 12, at constant volume, referred to equal weight of water (Regnault, Acad. 26, 1). Regnault (l.c.) gives the vapour-pressure.

sures of SiCl, as follows:-

Temp			Vap	our-pr	essure.
- 20°	•	•	. 26.49	mm.	mercury
 1 0		•	. 46.46	,,	**
0			. 78.02	,,	••
+10		•	. 125.90	,,	**
20		•	. 195.86	,,	17
80	•	•	. 294.49	,,	,,
40		•	. 429.08	,,	**
50			. 607:46	,,	**
60			. 837.23	,,	,,

H.F. [Si,Cl4] = 157,640 (Troost a. Hautefeuille,

A. Ch. [5] 9, 70).

Formation.—1. By passing dry Cl, or HCl gas, over a mixture of SiO₂ and C at full red heat (Oerstedt, B. J. 6, 119; Deville, A. Ch. [3] 43, 23). According to Weber (P. 112, 649) C is not necessary if the temperature is high enough.

2. By heating crystalline Si, or Si containing H, in Cl (Berzelius, C. J. 4, 91).—3. Favre (C. R. 107, 339) heats impure Si, and passes HCl and vapour of naphthalene over it.—4. By heating iron containing c. 15 p.c. Si in Cl (Warren, C. N. 60, 158).—5. By passing vapour of BCl₃ over SiO₂ heated to dull redness (4BCl₃ + 3SiO₂ = 3SiCl₄ + 2B₂O₃; Troosta. Hautefeuille, A. Ch. [5] 7, 476).

Preparation.-1. Amorphous SiO2 is mixed with an equal weight of lampblack, and the mixture is made into a paste with oil. Small pellets are made of this paste; these are imbedded in charcoal powder, and strongly heated. The dry pellets are placed in a porcelain tube, which is heated to very bright redness in a furnace while a stream of Cl (dried by H.SO, and then by CaCl.) is passed through. The exit then by CaCl2) is passed through. end of the porcelain tube is connected with a U-tube, from the bottom of the bend of which a short tube passes down into a small flask; the U-tube and small flask are surrounded by a freezing mixture. The SiCl, which condenses in the flask is shaken in a dry flask with Hg and a few pieces of K, and then distilled with sodium.

L. Meyer (A. 270, 238) recommends the use of charcoal-powder in place of lampblack, and starch-paste rather than oil.—2. Gattermann (B. 22, 186) places impure amorphous Si (containing some Mg, &c.), prepared as described under Siloon, Preparation, No. 1, in a tube c.

2 cms. wide, leaving a space of c. 1 cm. above the Si along the tube; he bends this tube at right angles, and connects it with a U-tube placed in a freezing mixture; he then passes dry Cl through the apparatus, while he heats the tube, laid in an iron trough in a gas-furnace so that the tips of the flames just touch the iron trough. The temperature must not get too high, else the Si glows, and MgCl₂ and O are formed, and the O oxidises the Si to SiO₂.

Properties.—A colourless, very clear, and very mobile liquid, with a disagreeable odour resembling that of cyanides; reddens litmus; fumes in moist air. Does not become viscid

at -75° (Haase, B. 26, 1052).

Reactions and Combinations.—1. Heated to redness with hydrogen a little SiHCl, is formed (Friedel a. Ladenburg, Bl. [2] 12, 92).—2. Oxychlorides are produced by the action of oxygen under the influence of induction sparks (Troost a. Hautefeuille, A. Ch. [5] 7, 465).—3. SiCl, is decomposed by passing the vapour with dry air through a white-hot tube, giving Si₂OCl₆ and Cl (F. a. L., l.c.). Berthelot (A. Ch. [5] 15, 185) says that SiO₂ is also produced.—4. Water produces H₂SiO₃ and HĈlAq.—5. Hydrogen sulphide forms SiCl₃.SH, when SiCl₁ and H₂S are passed through a red-hot tube (F. a. L., A. 145, 179; Pierre, A. Ch. [3] 24, 300).—6. Many metals—e.g. K, Na, Zn, Ag—withdraw all Cl from SiCl, when heated to redness in vapour of SiCl₄ (F. a. L.; cf. Rauter, A. 270, 235). 7. Most metallic oxides form SiO2 and metallic chlorides when heated with SiCl, (R., l.c.); CaO, MgO, Al₂O₄, and BeO form silicates and crystalline SiO₂ (Daubrée, C. R. 39, 135); TiO₂ is not changed (T. a. H., A. Ch. [5] 7, 476).—8. Decomposed by heating with alkali carbonates, potassium chlorate or nitrate, and various oxidising salts (R., l.c.).—9. SiO₂ and HCl are produced by reaction with sulphuric acid.— 10. Ammonia forms SiCl₄.6NH₃ (Persoz, A. Ch. [2] 44, 315); also, according to Gattermann (B. 22, 194), Si.NH₂.N.—11. Phosphoretted hydrogen does not react at ordinary temperatures; but at -40° c. 40 vols. of PH, are absorbed by SiCl, forming a solution which does not solidify at -60° (Besson, C. R. 110, 240). A colourless crystalline compound is produced by compressing PH, and SiCl, and then decreasing pressure to 20 atmospheres at 10° (B., l.c.).—12. Heated with sodium silicate, NaCl is formed. If SiCl, vapour is passed through a tube containing fragments of felspar, at a white heat, KCl, SiO2, and Si₂OCl₆ are formed (F. a. L., l.c.).

For the compound SiHCl, v. Silico-chloro-

говм, р. 453.

? SILICON DICHLORIDE SiCl₂. In the preparation of Si₂Cl₃ by the reaction of SiCl₄ with Si, Troost a. Hautefeuille (A. Ch. [5] 7, 463) obtained a liquid which took fire when heated in air, and reacted with ice-cold water to form a hydroxide of Si having reducing properties.

Silicon, chlorobromides of, v. Silicon Bromo-

CHLORIDES, p. 458.

Silicon, chlorohydrosulphide of, SiCl., SH. (Trichloro-silico-mercaptan.) Mol. w. 167:39. This compound may be looked on as a derivative of hypothetical ortho-thio-silicic acid Si(SH), obtained by replacing 3SH by 3Cl.

Prepared by passing dry H2S into SiCl, in a

retort, and then passing the mixed vapours of H₂S and SiCl, through a red-hot porcelain tube connected with a receiver surrounded by a freezing mixture, and fractionating the liquid that condenses (Friedel a. Ladenburg, A. 145, 179). Colourless liquid, with sharp, disgusting smell; boils at 96°; S.G. 145. V.D. 83·5 at c. 167°. Decomposed by moist air, more rapidly by water, to H₂S, S, HClAq, and SiO₂xH₂O; bromine produces SiBrCl₃ and HBr. Alcohol in excess forms Si(OEt)₄, H₂S, and HCl; by reacting in the ratio 3EtOH: SiCl₂:SH, F. a. L. (l.c.) obtained a liquid boiling at 164°-167°, probably Si(OEt)₄, SH; this liquid could not be obtained pure, as it changed to Si(OEt)₄, HCl, and H₄S.

Silicon, chloro-iodides of, v. Silicon iodo-

CHLORIDES, p. 461.

Silicon, compounds of, with carbon and oxygen. According to Colson (Bl. [2] 38, 56), the compound SiCO₂ is formed by heating Si with benzene vapour, the compound Si₂C₂O₂ by heating Si to whiteness in a carbon crucible, and the compound Si₂C₂O by heating Si in CO₂.

Silicon, compound of, with carbon and

Silicon, compound of, with carbon and sulphur. Colson (Bl. [2] 38, 56) says that a compound, to which he gives the formula Si_4C_4S , is produced by heating Si to a white heat

in CS2 vapour.

Silicon, compounds of, with hydrogen and oxygen. The compounds of the form SiO2.xH2O are described under Silica, HYDRATES OF (p. 447); the compounds Si₂H₂O₃ and Si₂H₂O₄ are described as silico-formic anhydride and silicooxalic acid respectively (pp. 455, 463). Wöhler (A. 127, 257) obtained a substance, by the decomposition of impure Ca silicide by HClAq, to which he gave the name silicone, and either the composition Si₄H₄O₃ or Si₃H₃O₂. On account of the yellow colour of this body, Miller proposed to call it chryscone. Silicone is described as orange-yellow leaflets, insol. water, alcohol, CS, SiCl, or PCl, when heated below redness it takes fire, burning to SiO, and amorphous Si; heated out of air, H is given off and SiO₂ and amorphous Si remain; not acted on by Cl, fuming HNO,, or conc. H.SO,; caustic alkali solutions, even dilute NH,Aq, produce SiO, with rapid evolution of H; acts as an energetic reducer towards solutions of many metallic salts.

Silicon, fluorides of. The existence of any compound of Si and F except SiF, is doubtful. Troost a. Hautefeuille (A. Ch. [5] 7, 464) obtained a fine dust by passing SiF, over molten Si and suddenly cooling, which was probably a lower fluoride than SiF,

SILICON TETRAFLUORIDE SiF₄. Mol. w. 104'3. Formation.—1. By passing BF₃ vapour through a red-hot porcelain tube (Troost a. Hautefeuille (A. Ch. [5] 7, 464).—2. By the interaction of HFAq and SiO₂ or silicates.

Preparation.—A fair-sized flask is one-third filled with a mixture of equal parts of powdered CaF₂ and quartz or white sand; sufficient conc. H.₂SO₄ is added to thoroughly moisten the mixture; the flask is gently warmed, and the gas that comes off is collected over Hg (2CaF₂ + 2H₂SO₄ + SiO₂ = 2CaSO₄ + 2H₂O + SiF₄).

Properties.—A colourless gas, with a disagreeable odour and sour taste; reddens dry litmus paper; fumes strongly in air. V.D. 51.0

(Dumas, A. Ch. [2] 33, 368). Condensed to a colourless mobile liquid at -106.5° and 9 atmos. pressure (Faraday, T. 1845. 155). According to Olszewski (M. 5, 127) SiF₄ solidifies at -102° . Acts as a powerful poison (v. Cameron, Dublin J. of Med. Sci. Jan. 1887).

Reactions.-1. Decomposed by electric sparks with separation of Si (T. a. H., C. R. 43, 443).-2. Potassium and iron, when hot, burn in SiF, forming fluorides and Si .- 3. Moist air forms Si₂O₃F.OH (Landolt, A. Suppl. 4, 27, and v. Dimetasilicio fluorhydrin, infra).—4. Water absorbs SiF₄, forming SiO₂xH₂O and H₂SiF₂Aq (v. SILICOFLUORHYDRIC ACID, p. 454).-5. Ammonia forms SiF. 2NH₃ (J. Davy, T. 1812. 352); also, according to Harris (C. C. 1889 (ii.) 283), SiNH, N (v. SILICON AMIDONITRIDE, p. 457).

Combinations.-1. Combines with dry phosphoretted hydrogen, when the gases are strongly compressed at -22° and 50 atmospheres in the ratio 2PH₃:3SiF₄, to form lustrous crystals (Besson, C R. 110, 80).-2. Alcohol absorbs SiF, freely, forming an acid liquid which gives off SiF, in the air (Knop, J. 1858. 146).—3. SiF, is absorbed by many metallic oxides.

Dimetasilicic fluorhydrin Si2O4.OH.F. Landolt (A. Suppl. 4, 27) obtained this compound as a white crystalline solid (v. also Schiff,

J. 1865. 196).

haloid compounds of. The com-Silicon, pounds of Si with halogens belong to the forms Si₂X₆ and SiX₄; the compounds SiX₄, where X is a single halogen, have been gasified, and the compound Si₂Cl₆ has also been gasified. The formulæ are probably all molecular. There are also indications of the existence of some compounds of the type SiX2. Besides the compounds where X is a single halogen, there exist the compounds SiXX', SiX₂X'₂, and SiX'X₃, where X and X' are Br and Cl, I and Cl, and I and Br respectively; the only one of these compounds that has been gasified is SiBr, Cl.

Silicon, hydrides of. The compound SiH4 The existhas been isolated and examined. ence of another hydride, probably SigHz, is

likely.

SILICON TETRAHYDRIDE SiH. (Siliciuretted hydrogen.) Mol. w. 32.3. Si and H do not combine directly, even in the electric arc (Friedel, C. R. 73, 497).

Formation.—1. Al containing Si is made the positive pole in NaClAq (Buff a. Wöhler, 4.103, 218).—2. Crude Mg silicide is decomposed by dilute HClAq (v. Preparation). These processes yield mixtures of SiH, and H .- 3. By the reaction of Na with SiH(OEt), (v. Prepara-

Preparation.-1. Small pieces of Na are dropped into SiH(OEt), (v. infra), and the liquid is gently warmed; the escaping gas is allowed to drive out the air (if the Na or SiH(OEt), was moist H comes off, and the gas takes fire), and is then collected over Hg (Friedel a. Ladenburg, 143, 123). The SiH(OEt), is prepared by adding dry absolute alcohol to pure SiHCl, in a longnecked flask, in the ratio 3EtOH : SiHCl., distilling after some time, and collecting the liquid that boils from 134° to 137° .- 2. Impure SiH, mixed with considerable quantities of H, is prepared by placing coarsely-pulverised crude Mg silicide (for preparation, v. infra) in a small two-necked flask, fitted with a funnel tube and a short wide delivery tube; the flask and delivery tube are quite filled with water from which all air has been driven out by boiling, the delivery tube is made to dip under boiled water, and the bell-jar to receive the gas is filled with boiled water; conc. HClAq is added, little by little, by the funnel tube. The gas, which comes off very rapidly, is collected in a bell-jar fitted with a stop-cock; this jar is then connected with a U-tube containing CaCl2 and carrying a short, narrow piece of glass tubing, which is made to dip under a dry vessel full of By depressing the bell-jar, the gas is forced out; it burns in the tubes till the air is exhausted, when it passes into the vessel full of Hg, where it is collected.

Preparation of thecrude silicide. A mixture of 1 part finely-powdered, dry, white sand and $1\frac{1}{2}$ parts Mg powder is heated in a stout glass tube; reduction takes place with production of much light, and a bluegrey, semi-molten mass is obtained (Gattermann, B. 22, 186). For other methods of preparing the Mg silicide required, v. Wöhler (A. 137, 369), Warren (C. N. 58, 215), and Mermet (Bl.

[2] 47, 306).

Properties.—A colourless gas; insol. water; liquefied at -11° and 50 atmos., -7 and 70 atmos., or -1° and 100 atmos. pressure (Ogier, A. Ch. [5] 20, 5). V.D. 15.9 (F. a. L., A. 143, 123). H.F. [Si,H'] = 32,900 (O., Lc.). Does not

react with N, NO, NH_s, H₂SO, Aq, or HClAq. Reactions.—1. Decomposed completely to Si and H by heating to above 400° (O., l.c.).—2. Induction sparks cause increase of volume from 100 to 121-129; H is produced, and yellow solid separates, probably Si₂H₃ (v. Disilicon Tra-HYDRIDE, infra). - 3. SiH, takes fire in air when slightly heated, or when the pressure is decreased; if the pure gas is passed into a tube filled with Hg, the pressure is lowered to 100-150 mm., and air is admitted, the gas takes fire (F. a. L., l.c.). The gas as prepared from Mg silicide is mixed with H, and takes fire in air at ordinary temperature and pressure.-4. Burns in chlorine with violent explosion. 5. Conc. potash solution forms K2SiO3 and H; the volume of H is four times that of the SiH, $(SiH_4 + 2KOHAq + H_2O = K_2SiO_8 + 8H)$ (F. a. L., i.c.).-6. Reduces solutions of many metallic salts; ppts. Ag and Si from AgNO, Aq, Pd from Pd salts, Cu silicide from CuSO, Aq, &c. Does not react with PtCl, Aq or Pb(C, H, O,), Aq.— 7. The action of electric discharges on SiH. mixed with nitrogen produces NH, and (?) Si2H, which combines with some of the N (Ogier, A. Ch. [5] 20, 31).

DISILICON TRIHYDRIDE SigHs. This formula is given by Ogier (A. Ch. [5] 20, 31) to a yellow solid obtained by the action of induction sparks on SiH4. The composition of the substance is somewhat doubtful. It takes fire when rubbed or heated in air; heated in H or N, inflammable SiH, is produced, or at higher temperatures Si

and H are formed.

Silicon hydrogen bromides v. Silico-Bromo-FORM, p. 453, and SILICO-ETHANE, PENTABROMO-DERIVATIVE OF (Si2HBrs), p. 454.

Silicon hydrogen chloride v. Silico-chloro**говм**, р. 453.

Silicon hydrogen iodide v. Silico-iodoform,

Silicon, hydroxides of, v. Silica, Hydrates OF, p. 447; SILICO-FORMIC ACID, p. 455; SILICO-FORMIC ANHYDRIDE, p. 455; SILICO-OXALIC ACID, p. 463; and Silicone, under Silicon, compounds of, with hydrogen and oxygen, p. 459.

Silicon, iodides of. Two compounds have been isolated, SiI, and Si₂I₆, and a third, which

is probably SiI2, seems also to exist.

SILICON TETRA-IODIDE SII, Mol. w. 534·42. Melts at 120·5°; boils at 290°. V.D. at 360°

= 268.5. H.F. [Si,F'] = 58,000 (Berthelot).

Preparation.—1. The crude product obtained by reducing a mixture of 4 parts fine white sand by heating with 1 part Mg powder (v. Silicon, Preparation of, No. 1, p. 456) is strongly heated in a mixture of dry CO, and I vapour, in a tube which projects c. 20 cm. rom the furnace; SiI, collects in the cold part of the tube, it is dissolved in CS₂ (1 part CS₂ dissolves c. 2.2 parts SiI4), shaken with Hg till colourless, and the CS₂ is evaporated by passing dry CO₂ through it at the lowest possible temperature (Gattermann, B. 22, 190).-2. A mixture of CO2 and I vapour is passed over Si strongly heated in a porcelain tube; the SiI, that collects in the cold part of the tube is purified as in 1 (Friedel, A. 149, 96).

Properties and Reactions.—Colourless, transparent, regular octahedra; isomorphous with (For M.P., &c., v. supra.) The vapour burns when heated in air, with separation of I. Water produces SiO₂xH₂O and HIAq. Alcohol forms EtI, HI, and SiO₂. Ether, at 100°, produces Si(OEt), and EtI (F., l.c.).

Silicon HEXA-IODIDE Si₂I₈. Formula pro-

bably molecular, from analogy of Si₂Cl₆. pared by heating SiI, with finely-divided Ag formed by reducing AgCl) to 290°—300° for some hours, removing SiI, by washing with a little dry CS₂ (1 part CS₂ dissolves c. 26 parts SiI, and c. 22 parts SiI,), dissolving in much hot CS2, and crystallising (Friedel a. Ladenburg, 4. 203, 254). Colourless, six-sided, double refractive plates. Melts at 250° in vacuo, with partial decomposition. When heated, decomposes to Sil,, and a substance that is probably Sil, Fumes in air; with water gives H,Si,O, and HIAq.

SILICON DI-IODIDE. The yellow solid that is formed by heating SiI, is probably SiI, it is insol. in the ordinary solvents; with water becomes grey, perhaps forming H₂SiO₂ (v. Silico-formic ACID, p. 455) (Friedel a. Ladenburg, A. 203, 247).

For the compound SiHI, v. SILICO-IODOFORM,

p. 455. Silicon, iodobromides of. Three compounds. corresponding with the three bromochlorides and the three iodochlorides, are formed by passing IBr (alone or mixed with H) over crystalline Si heated to dull redness; also by passing the vapour of a solution of I in SiBr, over crystalline Si at a low red heat (Besson, C. R. 112, 1447). The compounds are solids which decompose in air with separation of I; they all combine with NH₂ to form white compounds that are decomposed by water.

SILICON IODOTRIBROMIDE SilBr. Melts at 14° and boils at 192°. Prepared as described above, also (probably) by the interaction of I and

SiHBr, at 200°-250°, also by passing HI over SiBr, heated to low redness (B., l.c.).

SILICON DI-IODODIBROMIDE Sil Br. Melts at c. 38° and boils at 230°-231°.

SILICON TRI-IODOBROMIDE Sil, Br. Melts at c. 55° and boils at c. 255°.

Silicon, iodochlorides of. Three compounds have been isolated, corresponding with the three bromochlorides, and the three iodobromides. The V.D. of none has been determined, but from the analogy with the bromochlorides the sim-

plest formulæ are probably molecular.

SILICON IODOTRICHLORIDE SIICl. Obtained by passing HI mixed with SiCl, vapour through a red-hot tube, also by the interaction of HI and SiHCl, at 200°-250°, also by distilling ICl over crystallised Si heated to redness (Besson, C. R. 112, 60, 1314). A colourless liquid, boiling at 113°-114°; fumes in air; decomposed by water; I separates on standing, especially in sunlight. With NH_s forms white amorphous 2SiICl,.11NH,.

SILICON DI-IODODICHLORIDE Sil, Cl., Formed from HI and SiCl., also from ICl and Si; also by heating HI with SiI, Cl at 250°. Colourless liquid, boiling at 172°. With NH, forms amor-

phous Sil, Cl, 5NH, (B., l.c.).

SILICON TRI-IODOCHLORIDE Sil, Cl. A solid obtained in preparation of the two former compounds; melts at 2°. Fumes in air, with separa-

tion of I.

Silicon, nitrides of. Schutzenberger (C. R. 114, 1089) obtained small quantities of a compound to which he gave the formula Si, N, along with a carbide of Si, by heating 1 part Si and 2 parts SiO₂ in a covered carbon crucible, imbedded in lampblack in an outer crucible, to bright redness for some hours. By passing NH, through a porcelain tube kept at a white heat, Colson (C. R. 94, 1710) found a black layer in the hottest part of the tube, which was probably a mixture of Si and nitride of Si. A compound of Si, N, and H, probably Si.NH2.N, was obtained by Harris (C. C. 1889 (ii.) 283) by the reaction of NH, with SiCl, or SiF₄.
Silicon, oxide of, SiO₂; v. Silica, p. 446.
Silicon, oxychlorides of. The compound

Si₂OCl₆ is formed by heating SiCl₄ vapour in air or O. According to Troost a. Hautefeuille (Bl. [2] 35, 360) several oxychlorides are formed by passing a mixture of SiCl, vapour and O through a red-hot tube, or, better, by passing a mixture of 1 vol. Cl and \(\frac{1}{2} \) to \(\frac{1}{2} \) vol. O over crystalline Si heated to not above 800°. T. a. H. isolated the following oxychlorides:-

Si₄O₃Cl₁₀; liquid, b.p. 152°-154°. Si₂O₃Cl₂; liquid, b.p. 198°-202°. Si₂O₃Cl₂; oily liquid, b.p. above 400°. Si₄O₇Cl₂; solid, m.p. above 400°.

SILICON OXYCHLORIDE Si₂OCl₆. (Perchlorosilico-methyl ether (SiCl₃)₂O.) Mol. w. 284-78. Prepared by passing vapour of SiCl, through a porcelain tube heated in a wind furnace burning coke, condensing the product, repeating the operation with the portion which boils above 70°, fractionating, and separating the liquid boiling at 137°-138° (Friedel a. Ladenburg, A. 147, 855). A colourless liquid, boiling at 187°-138°. V.D. 144 at c. 200°. Fumes in air, decomposed by water to HClAq and SiO, xH,O. Miscible in all proportions with CHClas CClas CS.

SiCl, and Et,O. Alcohol produces Si₂O₇(Et)₆; zinc-ethide at 180° forms SiEt, and Si2OEt. (F. a. L., l.o.)

The compound SiP2O6Cl2 is described as SILICOPHOSPHORIC OXYCHLORIDE (p. 461).

Silicon, oxysulphide of. According to Colson (Bl. [2] 38, 56), a compound SiOS is formed, along with SiS and Si, SC, by heating Si to white heat in CS2.

Silicon, sulphides of. The only compound of Si and S certainly isolated is SiS2. Colson (Bl. [2] 38, 56) says that silicon monosulphide, SiS, is produced, along with SiOS and Si, SC, by heating Si to white heat in CS2; he describes SiS as a yellow solid, decomposed by water giving off H₂S, sol. in very dilute KOHAq with evolution of H. SiS is also said to be formed, along with SiS₂, by heating Si to reduces in H₂S (Sabatier, Bl. [2] 38, 153).

SILICON DISULPHIDE SiS. Mol. w. not determined. Sabatier (Bl. [2] 38, 153) gives H.F. [Si,S] = 19,900. Amorphous Si and S combine when strongly heated (Berzelius). Prepared by heating Si to redness in a stream of dry H2S; the other products are a yellow solid, probably SiS, and a brown substance that is likely a mixture of SiS₂ and SiS, or of SiS₂ and Si (S., *l.c.*; v. also Fremy, A. Ch. [3] 38, 324). Also prepared by strongly heating dried pellets of oil and SiO₂ (separated from SiF₄) in a slow stream of dry CS₂, quickly separating the white needles that form on the cooler part of the tube, and keeping in a closed tube (Fremy, l.c.). The residue that remains on distilling the products of the interaction of SiCl, and H₂S (v. Silicon chlorohydrosulphide, p. 459) contains SiS, and S; S may be removed by carefully heating in a stream of N (Gay-Lussac a. Thénard, A. Ch. [2] 69, 204). Long, lustrous, white needles; volatilised at very high tempera-tures (Fremy, *l.c.*). Unchanged in dry air at ordinary temperatures; decomposed by moist air to H₂S, and crystalline SiO₂ pseudomorphous with SiS₂; burns to SO₂ and SiO₂ when heated in air. Decomposed rapidly by water to SiO₂xH₂O and H₂S; also decomposed by alcohol and ether (Fremy, I.c.). HNO, oxidises rapidly, producing H2SO. M. M. P. M.

Silicon, sulphocyanide of; v. p. 463. SILICON, ORGANIC COMPOUNDS OF.

Silicic ethers are described elsewhere -v. AMYL, AMYLPHENYL, ETHYL, METHYL, PHENYL, THYMYL, TOLYL, and XYLYL SILICATES.

Tetra-methyl-silicane SiMe, Mol. w. 88. (31°). V.D. 3.08 (calc. 3.04). Formed by heating SiCl₄ with ZnMe₂ at 120° (Friedel a. Crafts, A. 136, 203). Light oil, burning with bright flame emitting SiO₂. Not attacked by potash or

HNO, (S.G. 1.4).

Tetra-ethyl-silicane SiEt. Mol. w. 144. (153°). V.D. 5·13 (calc. 4·99). S.G. 2·834. Prepared by heating SiCl, with ZnEt, at 160°. Formed also by the action of ZnEt, and Na on ethyl silicate (Friedel a. Crafts, Bl. 1863, 468; 1865, 858; A. 127, 31; 138, 19). Oil, insol. H₂SO, Chlorine forms liquid Et,SiC,H,Cl

Formed by distilling Si₂I₆ with ZnEt₂ (Friedel a. Ladenburg, A. 203, 251). Oil, burning with bright flame.

'Silicopropionic acid' EtSiO.OH. Got by heating its ortho- ether with HIAq (Ladenburg, A. 159, 271; 164, 305). Amorphous powder, insol. water and Na CO, Aq, sol. conc. KOHAq.

Methyl ortho- ether EtSi(OMe), (126°). S.G. 9747. Formed from Si(OMe), sodium,

and ZnEt₂ (Ladenburg, B. 5, 1081). Oil.

Ethyl ortho-ether EtSi(OEt), (159°).
S.G. 2 927. Formed by the action of Na and ZnEt, on ClSi(OEt), or Si(OEt), Oil. Converted by BzCl into EtOBz and EtSiCl, (100°), which fumes in the air and is converted by water into silicopropionic ether.

'Silicodiethyl ether' Et2Si(OEt)2. (156°). S.G. 2 875. V.D. (H=1) 87 (calc. 88). Formed from Na, ZnEt₂, and silicin ether. Oil, sol. alcohol and ether. AcCl (1 mol.) at 200° forms EtOAc and Et₂SiCl(OEt) (147°). A larger quantity (2 mols.) of AcCl at 250° forms Et,SiCl, (129°), which is decomposed by water, forming syrupy SiEt₂O (above 360°), which may also be got by boiling SiEt (OEt), with HIAq.

'Silicoheptyl ether' SiEt3.OEt. Mol. w. 160. (153°). S.G. 4.840. V.D. (H = 1) 80.8(calc. 80). Formed from SiEt, (OEt)2, Na, and ZnEt₂. Oil, sol. alcohol and ether. Sol. conc. H₂SO₄. AcCl at 180° forms SiEt₃Cl (144°), S.G. 2.925, a fuming liquid converted by NH, Aq into SiEt,OH.

Tri-ethyl-silicol SiEt, OH. Mol. w. 132. (154°). S.G. 2 871. Formed by dropping SiEt, OEt into NH₃Aq, or by heating it with Ac₂O at 250° and decomposing the resulting SiEt₃.OAc with Na₂CO₂Aq. Thick liquid, smelling like camphor. Insol. water, miscible with alcohol and ether. Burns with bright flame. Sodium forms SiEt, ONa, which, in ethereal solution, combines with CO₂, forming SiEt₃.CO₂Na, an amorphous deliquescent solid, decomposed at a red heat into Si₂Et₈O and Na₂CO₃. AcCl forms HOAc and SiEt₃Cl. HIAq at 200° forms ethane and SiEt₂O. Oxidising agents have no action. H2SO4 forms EtSiO2H, ethane, H, and SO4.

Acetyl derivative SiEt₃OAc. (168°). S.G. ² '903. Formed by heating SiEt₃.OEt with Ac₂O at 250°. Liquid with ethereal odour. 'Silicoheptyl oxide' (SiEt₃)₂O. Mol. w. 246.

(231°). S.G. 2.859. Formed as above, and also by the action of P2O5 on tri-ethyl-silicol. Got also from SiEt, Cl and KOHAq. Colourless syrup, sol. H2SO, and separated unchanged on dilution, if heating be avoided.

Tri-ethyl-silicane SiEt,H. ² ·751. $\hat{V}.D.$ (H = 1) 59 (calc. 58). Formed, together with SiEt, by the action of excess of ZnEt, and Na on Et, SiO, Liquid, insol. water and H, SO, sol. alcohol and ether. Fuming H2SO, converts it into (SiEts)2O. HNOs attacks it vigorously. Bromine added slowly to the cooled liquid forms SiEt, Br (161°), which is converted by NaOHAq into (SiEt₃)2O, and by NH, Aq into SiEt, OH.

H₂SO₄. Chloring forms inquid Et₃SIC₂H₄OI (185°), converted by alcoholic KOAc at 180° into Et₃SiC₂H₄OAc (208°-214°), whence alcoholic potash at 120° forms oily Et₃SiC₂H₄OH (190°).

Hexa-ethyl di-silicane Si₂Et₆. (252°).

S.G. § *851; \$\frac{2}{38}\$* 840. V.D. 8*6 (calc. 7.96). Hygroscopic liquid, decomposed by alkalis with

evolution of H. Sodium decomposes it into SiH₄ and Et₄SiO₄.

Chloro-silicoformic ether v. ETHYL SILICATE. 'Silicoacetic acid.' Ethyl ortho-ether CH₃.Si(OEt)₃. (145°-151°). S.G. 2°928. Formed by heating silicic ether with ZnMe₂. Oil, sol. alcohol. Converted by HIAq into amorphous silico-acetic acid CH₃.SiO₂H, which is insol. water and ether (Ladenburg, B. 6, 1029).

The term silicoacetic acid is also sometimes used to denote $H_2Si_2O_4$, which is got by decomposing SiI_4 by water at 0° (Friedel a. Ladenburg, A. 203, 249; v. Silico-oxalic acid, infra). A compound $Si_2C_3O_2$ is formed, as a bottle-green powder, by heating silicon to whiteness in a crucible lined with lampblack. It is insol. KOHAg and HFAg (Colson. C. R. 94, 1316).

KOHAq and HFAq (Colson, O. R. 94, 1316).

Tetra-propyl-silicane Si(C₃H₁)₄. (213°).
S.G. 2 '7979; ½ '7883. V.D. 99·7 (obs.).
Formed, together with SiHPr₃, by heating ZnPr₂ (2 pts.) with SiHCl₃ (1 pt.) at 150° (Pape, B. 14, 1872; A. 222, 370). Colourless oil, sol. alcohol and ether, insol. H₂SO₄. Br has no action in the cold, but on warming it forms oily SiC₁₂H₂, Br, converted by alcoholic potash into oily SiC₁₂H_{2s} (206°-210°).

Tri-propyl-silicane SiHPr_s. (170°). S.G. 2 '7723; ¹⁵ '7621. V.D. 82 (obs.). Formed as above. Oil, v. sol. alcohol and ether, insol. conc. H₂SO₄. Burns with bright flame, forming SiO₂. Br acts violently, forming SiPr_sBr (213°), a fuming liquid, which is slowly converted by

water into SiPr₃.OH.

Tri-propyl-silicol SiPr, OH. (205°-208°). Formed by the action of NH, Aq on SiPr, Br, or of Na₂CO, Aq on SiPr, OAc (Pape). Oil.

Acetyl derivative SiPr. OAc. (212°-216°). Formed from SiPr. Br and AgOAc. Oil.

Hexa-propyl-di-silicyl oxide (SiPr₃)₂O. (280°-290°). Formed by warming SiHPr₃ with H₂SO₄ and, together with SiPr₃·OH, by boiling SiHPr₃ with Na₂CO₃Aq (Pape, A. 222, 369). Liquid, sol. alcohol, ether, and H₂SO₄.

Silicon sulphocyanide Si(CNS). [142°]. (c. 300°). Formed by distilling lead sulphocyanide with SiOl, (Miguel, A. Ch. [5] 11, 343). Colourless prisms, insol. ether and CS₂, soluble in a solution of HCNS in benzene. Burns with violet flame. Dyes the skin red. Decomposed

by water into HCNS and SiO₂.

Tetra-phenyl-silicane SiPh₄. [233°]. (above 860°). S.G. $\frac{25}{20}$ 1·0780. Formed from chlorobenzene, SiCl₄, and Na in presence of a little EtOAc (Polis, B. 18, 1540; 19, 1012; 20, 3331). Dimetric crystals; a:o=1: '440, sol. hot benzene, sl. sol. alcohol and ether. May be sublimed. Sulphonated by fuming H_2SO_4 . HNO₃ forms Si(C₅H₄,NO₂)₄ [105°], a yellow powder, v. sol. benzene. Converted by PCl, into liquid SiPh₂Cl₂ (234° at 90 mm.) and SiPh₃Cl [89°], from which water produces tri-phenyl-silicol SiPh₃.OH [141°], which separates from ether in colourless crystals.

Pheuyl-tri-chloro-silicane SiPhCl_s. (197°). Formed by heating SiCl₄ with HgPh₂ at 300° (Ladenburg, B. 6, 379). Oil. Decomposed by hot water, or by NH₂Aq, yielding 'silicobenzoic acid' SiPh(OH)_s [92°]. Absolute alcohol forms 'silicobenzoic' ethyl ortho- ether SiPh(OEt)_s (237°). S.G. 2 1·013; ½ 1·006. Silicobenzoic acid is converted at 100° into amorphous 'silicobenzoic anhydride' (SiPhO)₂O.

Phenyltriethyl-silicane SiPhEt₃. (230°). S.G. ² 904. Formed, together with SiEt, and SiPh₂Et₂ (c.310°), by heating SiPhCl₃ with ZnEt₄ at 150°. Oil, smelling like cloves. Yields a bromo-derivative (270°-280°). Cl forms SiCl₁₂H₁₅Cl (260°-265°), S.G. ² 10185.

Tetra-m-tolyl-silicane Si(C,H.). [151°]. (above 550°). S.G. 2º 1·119. Formed from m-bromo-toluene, SiCl., and Na (Polis, B. 19, 1021). Pale-yellow needles (from ether), v. sol. benzene and chloroform, insol. alcohol.

Tetra-p-tolyl-silicane Si(C,H,). [228°]. (above 360°). S.G. 22 1·079. Formed from p-bromo-toluene, SiCl., and Na (Polis, B. 18, 1542). Colourless crystals, sol. benzene.

p-Tolyl-tri-chloro-silicane C,H,SiCl_s. (219°). Formed from Hg(C,H₁)₂ and SiCl_s at 310° (Ladenburg, A. 173, 165). Fuming liquid, decomposed by water. Converted by NH₃Aq into viscid C,H,SiO₂H, which is converted at 200° into the solid anhydride (C,H,SiO)₂O, which is not melted at 200°.

Tetra-benzyl-silicane v. vol. i. p. 502.

Silicon tetra-phenyl-tetra-amide Si(NHPh),. [138°]. Formed from SiBr, and excess of aniline diluted with benzene, the product being distilled in a current of H at 105° (Reynolds, C. J. 55, 477). Colourless monoclinic crystals; a:b:o=985:1:1.043; $\beta=110^{\circ}20'$. V. sol. benzene, sol. CS₂, insol. ligroïn. Decomposed by water and alcohol. Not decomposed by heating at 210°. HCl forms SiCl, and aniline.

Silicon di-chloro-di-phenyl-di-amide SiCl₂(NHPh)₂. Formed from SiCl₄ and aniline (Harden, C. J. 51, 40). Amorphous solid. Decomposed by water into aniline hydrochloride and silica.

Silicon tetra-o-tolyl-tetra-amide Si(NHC,H₁)₄. Formed from SiCl₄, o-toluidine, and benzene (Reynolds). Prisms, v. sol. benzene and CS₂. The compound SiCl₂(NHC,H₁)₂, which is also formed from SiCl₄ and o-toluidine, is a white granular powder (Harden).

Silicon tetra-p-tolyl-tetra-amide Si(NHC,H,),. [132°]. Formed from SiCl₄ and p-toluidine in benzene (Reynolds). Botryoidal aggregate of needles, v. sol. benzene and ether, decomposed by water and alcohol.

Silicon tetra-naphthyl-tetra-amide $Si(NHC_{10}H,)_4$. Formed from (β)-naphthylamine and $SiCl_4$ (R.). Nodules, decomposed by water. Sl. sol. ligroïn, sol. benzene.

SILICO-NITRATES. SiO₂ seems to form some compounds with N₂O₃ and basic oxides the salt 3SiO₂.N₂O₃.7Ag₂O (=3Ag₄SiO₄.2AgNO₂) was obtained, in ruby-coloured prisms, by Rousseau a. Tite (C. R. 114, 294), by heating AgNO₃ with a little water and fragments of marble in a sealed tube at 180°-300° for several hours; heated to dull redness, it gave off Noxides, and Ag and Ag₂SiO₃ remained.

M. M. P. M.

SILICO-OXALIC ACID $H_2Si_2O_4 = {0:Si.OH \atop O:Si.OH}$

This compound is obtained, as a white amorphous powder, by the interaction of water and Si₂Cl₅ or Si₂I₆ at 0°; the gelatinous pp. is washed with ice-cold water, dried in vacuo and then at 100° (Friedel a. Ladenburg, A. 203, 118). It is also produced by the interaction of Si₂I₆ with

absolute alcohol. The compound is decomposed even by weak bases, with evolution of H. M. M. P. M.

SILICO PHOSPHORIC OXIDES. (Silicophosphoric acids.) SiO₂.P₂O₅ and SiO₂.2P₂O₅.4H₂O. The compound SiO₂.P₂O₁ = SiP₂O₂ is formed by dropping SiO₂, prepared by decomposing SiF, by water and drying the pp., into molten HPO₃, and washing with water (Hautefeuille a. Margottet, C. R. 96, 1052; v. also Skey, C. N. 16, 187). Large, hard, transparent crystals; S.G. 3·1 at 14°; melts when strongly heated to a glass-like mass. crystals are polymorphous; they are hexagonal under 800°; at c. 800° they form leaflets resembling tridymite, between 700° and 800° they are regular octahedra, and between 800° and 1000° they assume the form of clinorhombic prisms. Water attacks the hexagonal, but not the octahedral or prismatic, orystals. Molten AgNO₃ forms Ag₃PO₄ and SiO₂ with all the forms (H. a. M., C. R. 99, 789).

The compound SiO₂.2P₂O₃.4H₂O was obtained (H. a. M., C. R. 104, 56) by one-fourth saturating H₂PO₄Aq with SiO₂xH₂O, heating the solution in a Pt dish to 125° for 7-8 hours, and drying the solid which separated. The compound is a crystalline powder; decomposed by moist air; sol. water at 0°, but decomposed by water at the ordinary temperature to $H_{2}PO_{4}Aq$ and gelatinous $SiO_{...}xH_{2}O$. M. M. P. M.

SILICOPHOSPHORIC OXYCHLORIDE SiP₂O₅Cl₂. (Silicopyrophosphoryl chloride.) Prepared by heating SiCl₃.OEt, or Si(OEt)₄, with excess of POCl₃ to 180° for 2 hours, distilling off EtCl, and SiCl, if prepared from SiCl, OEt, and heating the solid that remains to 150°-200° in a stream of dry air to remove adhering POCl_s. A white, loose, very hygroscopic, amorphous powder. Very sol. water, but with separation of SiO₂.xH₂O; easily sol. alcohol, insol. ether. or SiO₂:xH₂O; easily sol. alcohol, insol. ether. Decomposed very slowly below 200°, more rapidly above 200°, giving off POCl₃; at a red heat P₂O₅ comes off, and a glassy mass remains, probably consisting of SiO₂ and SiO₂:P₂O₅. Addition of excess AgNO₃Aq to solution in water (with a little alcohol to prevent separation of silicol followed by HNO Ag and AgCl tion of silica), followed by HNO, Aq, ppts. AgCl, and on filtering this off and adding NH, Aq drop by drop Ag, P₂O, is ppd. mixed with some Ag, P₂O. Heated to 100° with PCl, the equation $SiP_2O_aCl_2 + 4PCl_5 = SiCl_4 + 6POCl_5$ is realised. The constitution of the compound is probably

O.PO.Cl either $Si \leqslant_{O_2:PO.Cl}^{O_2:PO.Cl}$ or O:Si(Stokes. 0.PO.Cl

Bulletin U.S. Geolog. Survey, No. 90 [1892] 47; also in B. 24, 933). M. M. P. M.

SILICOTITANATES. Compounds of silicates and titanates are found in certain minerals; sphene, for instance (CaO.3SiO_2CaTiO_s), may be called a silicotitanate. M. M. P. M.

SILICOTUNGSTATES: Compounds containing SiO2, WO2, and basic radicles; v. TUNGSTO-SILICATES.

SILICOVANADATES v. VANADO-SILICATES.

SILK v. PROTEÏDS.
SILVER. Ag. At. w. 107.66. Mol. w. not known (v. infra, Properties). Melts at c. 950°;

Person (A. Ch. [8] 27, 250) gave 1000°, Daniell (T. 1830. 237) 1024°, Violle (C. R. 85, 543) 954°, Becquerel (J. 1863.) 916°, Deville (B. 12, 791) 916°. B.p. not known; Meyer (B. 12, 1428) says that Ag does not appreciably evaporate at c. 1570°. S.G. c. 10.5; Playfair a. Joule (C. S. Mem. 3, 66) gave 10.53, and 9.13 to 9.28 for molten Ag, Dumas (C. N. 37, 82) gave 10.512 for Ag heated in vacuo, Roberts-Austen gave 10.57, and 9.46 to 9.5 for liquid Ag (Pr. 23, 495); for further data v. Clarke's Table of Specific Gravities [new ed.] 14. S.H. 0°-100° -.0557 (Dulong a. Petit, A. Ch. [2] 7, 113); - '0557 (Dulong a. Feut, A. Ch. [2] 1, 110, '0559 (Bunsen, P. 141, 1); '05722 (Louguinine, A. Ch. [5] 27, 398). C.E. '00002 between 0° and 100° (v. Matthiessen, P. 130, 50; Fizeau, C. R. 68, 1125); '00003721 from 0° to the m.p. (Roberts-Austen, Pr. 23, 495). T.C. 100 (that of copper = 73.6, and of Au = 53.2, Wiedermann a. Franz, A. 88, 191); heat sufficient to raise 1096 mgm. water from 0° to 1° passes in 1 second through each sq. mm. of a plate of Ag 1 mm. thick, the two sides of which differ in temperature by 1° (Weber, B. B. 1880. 467). E.C. (Hg at 0°=1) 57.226 for soft Ag, 63.845 C. R. 76, 382). H.C. [Ag²,O] = 5.900 (Th. 3, 381). Heat of fusion (for 108 g.) 2·275 (Person, A. Ch. [8] 24, 275); 2·67 (Pionchon, A. Ch. [6] 11, 100). S.V.S. c. 10·3. Emission spectrum gives a line 5464 in the yellow, 5209 9 in the green, and many lines in the orange, green, blue and violet (v. B. A. 1884. 442); for absorptionspectrum of Ag vapour v. Lockyer a. Roberts-Austen (Pr. 23, 344). Refraction equiv.

 $=\frac{\mu-1}{d}$. At. w. = 12.62 (Kanonnikoff, J. R. 1884.

[1] 119); 13·2 (Gladstone, Pr. 18, 49).

Occurrence. - Found native, generally alloyed with Au, Cu, &c. The chief compounds found in ores are Ag.S (frequently in combination with Cu.S, Fe₂S₃, Sb₂S₃, As₂S₃, Bi₂S₃, PbS, &c.), AgCl, AgBr, AgI, compounds of Ag with Se. Te, As, Sb, Bi, &c. Ag is also found in small quantities in most lead ores. According to Malaguti (J. pr. 42, 422; cf. Field, D. P. J. 143, 397), traces of salts of Ag (c. 1 mgm. Ag in 100 litres) are found in sea-water. Small quantities of Ag, probably as AgCl, have been found in volcanic dust (Mallet, Pr. 47, 277). Silver has been known and used from the earliest times (for a full historical account of silver, including the derivation of the name, v. Silber in Ladenburg's Handwörterbuch der Chemie, 10, 635).

Formation.—1. By heating Ag₂S with PbO or PbSO4, SO2 and a compound or alloy of Ag and Pb are formed; the Pb may be separated from this alloy by cupellation.—2. By shaking AgCl with Hg and water, HgCl and an amalgam of Ag and Hg are formed; Ag remains when the Hg is removed from this amalgam by heating.-3. Ag is ppd. from solutions of its salts by Zn, Cu, Fe, &c.-4. AgCl is reduced to Ag by treatment with many reducing agents, such as Zn and H₂SO₄Aq, also by fusion with Na₂CO₃ and C.—5. Most easily-oxidised metals ppt. Ag from solutions of its salts. For an account of the extraction of Ag from its ores v. Dictionary of Applied Chemistry, vol. iii. p. 386.

Preparation.—Silver coins are dissolved in-

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pure dilute boiling HNO, Aq, the solution is evaporated to dryness, the residue is heated till it melts, after cooling it is dissolved in NH, Aq, after standing forty-eight hours the liquid is passed through a filter made of specially good paper, and the filtrate is diluted till it contains not more than 2 p.c. of Ag. This ammoniacal solution contains AgNO₂ and Cu(NO₂); the whole of the Ag is ppd. by ammoniacal solution of (NH₄)₂SO₃ at 60°. A solution of (NH₁)₂SO₃ is prepared by saturating pure NH₂Aq with SO₂ (made by heating pure Cu with pure H2SO. diluted with 1/2 to 2/8 its volume of water); a measured quantity of this solution is mixed with excess of pure NH,Aq, heated to boiling, and the ammoniacal solution containing AgNO, and Cu(NO₃)₂ is run from a burette into the boiling liquid until there is the faintest trace of blue colour in the liquid above the ppd. Ag. Cuprous sulphite is formed, and this reduces the AgNO, in the ammoniacal liquid; when all the AgNO, is reduced, the blue colour of the ammoniacal copper nitrate appears. The whole of the ammoniacal solution of AgNO, and Cu(NO_s)₂ is now mixed with the proper quantity of ammoniacal (NH₄)₂SO₃Aq required for complete reduction of the AgNO₃, the mixed liquids are set aside in a closed vessel for fortyeight hours, when c. \frac{1}{8} of the Ag is ppd.; the blue liquid is decanted off and heated on a water-bath to 60°-70°, whereby the whole of the Ag in solution is ppd. The two quantities of ppd. Ag are washed with ammoniacal water, by decantation, until the washings show no trace of blue colour on standing (or until BaCl2Aq gives no trace of pp.), the Ag is kept in contact with conc. NH, Aq for some days, and is then washed with water until every trace of NH_s is removed. If it is desired to have the Ag in bars, the finely-divided metal obtained by ppn. is mixed with 5 p.c. of its weight of pure borax which has been strongly heated, and 5 p.c. pure NaNO, fused in a porcelain crucible and poured into kaolin moulds lined with a paste made of a mixture of kaolin that has been strongly heated and unignited kaolin; when cold the bars are cleaned with fine sand, then strongly heated with pure KOHAq (to remove traces of kaolin) and washed with water. The bars may be cut with a chisel of very hard steel; if this is done the pieces must be warmed for some time with pure conc. HClAq, washed with NH, Aq, and then with water (Stas. Rech. 38 et seq.). The Ag may be freed from traces of absorbed gases by distilling it; this is done by placing the Ag in a hole in a block of strongly-heated marble; this hole communicates, by a slit in the marble, with another hole from which an opening passes upwards to the surface of the marble; another opening admits the nozzle of an O-H blowpipe, so that the flame plays on the surface of the Ag. (For details v. Stas, l.c.) Dumas (A. Ch. [5] 15, 289) found that 1 kilo. of Ag considered pure gave from 59 to 174 c.c. O when heated in a Sprengel vacuum; it was supposed by some chemists (v. Mallet, T. 1880. 1003; Clarke, Recalculations of the Atomic Weights, 262) that this result proved that Ag made by Stas' process was not quite pure; but Brauner has proved experimentally (C. J. 55, 899) that Ag prepared Voz. IV.

by Stas's method described above, and distilled by the O-H flame in a block of burnt marble, gives off no O when heated in vacuo (v. also Stas, Bull. Acad. roy. Belg. [3] 18, 23; abstract in C. J. 58, 561).

Properties.-A white, very lustrous metal. Crystallises in forms in the regular system, chiefly cubes, octahedra, and rhombododecahedra; these crystals are obtained by slowly cooling molten Ag, or by the electrolytic deposition of the metal; they are also sometimes found native. Finely divided Ag—as obtained, for instance, by reduction of AgCl—is a grey powder, which becomes white and lustrous when compressed with a flat iron. A thin deposit of

Ag on glass transmits bluish light.

Ag is harder than Au and softer than Cu. Ag reflects light and heat strongly; its absorptive power is very small. It is very malleable; Ag foil has been obtained 003 mm. thick. Ag is very ductile; 06 gram may be drawn to a wire 130 metres long; it is so tenacious that a cold-drawn wire of 2 mm. diameter does not break till it is loaded with 80-85 kilos (Baudrimont, A. Ch. [8] 30, 304). Ag is slightly volatile at a low-red heat in vacuo, but not at the ordinary pressure at this temperature, nor in vacuo at 440°-450° (Stas, Bull. Acad. roy. Belg. [3] 18, 23). It may be distilled, by heating, by the O-H flame, Ag placed in a block of burnt marble. Stas distilled 50 g. Ag in c. 15 mins.; part of the Ag appeared as a blue vapour during the distillation. When an alloy of Ag with a more volatile metal, such as Hg, Pb, or Sn, is heated, part of the Ag volatilises with the other metal. Molten pure Ag absorbs c. 20 times its volume of 0 from the air (v. Lucas, A. Ch. [2] 12, 402; Gay-Lussac, A. Ch. [2] 45, 221; Graham, P. M. [4] 32, 503; Neumann, M. 13, 40; Levol, J. pr. 57, 192); as the Ag cools it 'spits' and gives off the O it has absorbed; Dumas' experiments (A. Ch. [5] 15, 289) show that a little O is retained, but Brauner (C. J. 55, 399) has proved that no O remains if the Ag is distilled in a (For the chemical block of burnt marble. action of O on Ag v. Reactions, No. 1.) Pure Ag suffers no change when heated in air, H, or CO (van der Plaats, Mandblad voor Natuur-wetenschappen, 1886. No. 2). Molten Ag ab-sorbs P, but the whole of the P, except c. 002 p.c., separates on cooling (Warren, C. N. 56,

The at. wt. of Ag has been determined (1) by finding the ratio of Ag to Cl in AgCl (Berzelius, P. 8, 17 [1826]; Turner, T. 1829. 297 [1829]; Marignac, A. 44, 23 [1842]; ibid. Bibl. univ. Genève, 46, 354 [1843]; Maumené, A. Ch. [8] 18, 41 [1846]; Dumas, A. Ch. [3] 55, 184 [1859]; Stas, Stas. Rech. 88 et seq. [1860]); (2) by finding the ratio of Ag to Br in AgBr, and of Ag to I in AgI (Marignac, l.c. 46, 860 [1843]; Stas, Stas. Nouv. R. 158 et seq. [1865]); (3) by determining the ratio of Ag to Ag.S formed therefrom (Dumas, A. Ch. [8] 55, 147 [1859], Stas, Stas. Rech. 53 [1860]); (4) by reducing Ag₂SO₄ to Ag (Struve, A. 80, 203 [1851]; Stas, Stas. Rech. 125 [1860]); (5) by converting Ag₂SeO₅ into Ag (Pettersson a. Ekman, Bl. [2] 27, 205 [1876]); (6) by converting Ag into AgNO₅ (Marignac, A. 59, 289 [1848]; Stas, Stas. Stas. Stas. Converting Ag in the Agent So. et as (1960)). (7) by Agrandician Rech. 50 et seq. [1860]); (7) by determining

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S.H. of Ag; (8) by finding V.D. of AgCl at white heat (Biltz a. Meyer, B. 22, 725). The classical work of Stas has placed the value for the at. wt. on a very sure foundation.

Ramsay (C. J. 55, 521) measured the depression of the vapour-pressure of Hg by dissolving Ag therein; assuming that the mol. w. of liquid Hg is 200, and that equal volumes of dilute solutions of metals in Hg contain equal numbers of molecules, the results led to the conclusion that the mol. w. of Ag is the same as the

AgCl is the only compound of Ag the vapourdensity of which has been determined; the value found shows that the atom of Ag is monovalent in the gaseous molecule AgCl.

Ag is fairly closely related chemically to Cu and Au (v. Copper group of Elements, vol. ii.

p. 250).

Allotropic forms of silver. According to Schneider (B. 25, 1281, 1440), an aqueous solution of Ag, containing c. 20 g. Ag per litre, is obtained by mixing 150 g. FeSO. 7aq in 500 c.c. water and 280 g. Na citrate in 700 c.c. water, pouring the mixture into 500 c.c. of 10 p.c. AgNO2Aq (it is best to divide the mixture into five parts, as small quantities are more easily dealt with), removing liquid from ppd. Ag after half an hour (by a pipette), filtering through a good filter under decreased pressure, and dis-solving the ppd. Ag in as little water as possible; by adding absolute alcohol, and filtering, after some days, with help of a good waterpump, colloidal Ag containing from .8 to .5 p.c. of Fe salts is obtained. A very little HClAq ppts. Ag with a little AgCl and Ag₂Cl (?) from solution of colloidal Ag. Carey Lea has carried out a series of researches on the properties of Ag ppd. from solutions of its salts by ferrous citrate, ferrous tartrate, and dextrin in presence of alkali (Am. S. [8] 37, 476; 38, 47, 129, 237; 41, 179, 259; P. M. [5] 31, 238, 320, 497; 32, 337). The pps. of Ag show almost every shade of colour, such as blue, red, green, purple, golden; some of the pps. are soluble in water, and some are not; most of the pps. are very sensitive to light, but they vary much in this respect; some of the pps. yield ordinary Ag when heated; ordinary Ag is also ppd. from solutions of soluble colloidal Ag by a little very dilute HClAq. In no case does any so called allotropic Ag seem to have been obtained quite free from organic compounds; the percentage of Ag in the pps. varied from c. 90 to c. 98. For accounts of the properties of the various coloured pps. the memoirs of Carey Lea must be consulted; v. also Schneider, B. 24, 8370; Barus a. Schneider, Z. P. C. 8, 278; Prange, R. T. C. 9, 121; Muthmann, P. 20, 983. B. a. S. determined many of the physical properties of a solution of colloidal Ag, and concluded that the colloidal variety consists of extremely minute particles of ordinary Ag that remain suspended in the liquid for a considerable time.

Reactions and Combinations.—1. According to Debray (C. R. 66, 735), Ag is partially oxidised by very strongly heating in the O-H flame, using excess of oxygen. According to Le Chatelier (Bl. [2] 48, 342) pure ppd. Ag is oxidised by heating in oxygen at 300° under 15 atmos. pressure; after 50 p.c. of the Ag was

oxidised the change stopped.—2. Osonised oxygen acts on moist, finely-divided Ag at the ordinary temperature, forming a peroxide (v. SILVER PEROXIDE, p. 471).--3. Ag combines easily with chlorine, bromine, iodine, and sulphur (v. SILVER CHLORIDE &c.); also, by heating, with selenion, arsenic, and phosphorus (v. SILVER SELENIDE &c.).—4. Finely-divided Ag is said to dissolve, with oxidation, in ammonia solution in presence of air (Carey Lea, Zeit. für anorg. Chem. 8, 180).—5. Ag reacts with steam at a white heat, absorbing O while H is given off.— 6. Finely-divided Ag is oxidised by hydrogen peroxide (Berthelot, A. Ch. [5] 21, 164).— 7. Finely-divided Ag is said to be oxidised by heating with oxides and salts that readily give up oxygen—e.g. MnO₂, Pb₃O₄, Pb(NO₃)₂, &c.— 8. Hydrogen sulphide forms Ag.S. Alkali sulphides also produce Ag₂S.—9. Molten sodium chloride forms AgCl, some Na vapour being given off; solution of sodium chloride, as also of potassium or ammonium chloride, dissolves some Ag as AgCl .- 10. Potassium iodide solution, in air, forms soluble AgI.KI.—11. Hot solution of potassium cyanide dissolves Ag, forming AgK(CN), (v. Christomanos, Fr. 7, 301) .- 12. Fusion with silicates, such as glass, forms Ag₂O, which dissolves in the molten silicate.—13. Ag dissolves in solution of ferric sulphate on warming; as the solution cools the Ag₂SO₄ is again decomposed, with formation of Fe₂(SO₄)₈ and ppn. of finely-divided Ag.—14. A compound of Ag and Si is said to be formed by strongly heating Ag with potassium silico-fluoride and potassium (Warren, C. N. 60, 5).— 15. Ag dissolves in moderately dilute nitric acid, forming AgNO₂, and giving off N oxides; the gaseous products are N₂O and NO, according to Montemartini (G. 22, 384, 397, 426).—16. Conc. hydrobromic and hydriodic acids dissolve Ag on warming, with evolution of H; the dissolved AgBr or AgI is ppd. on dilution.—17. Hot conc. sulphuric acid dissolves Ag, giving off SO₂; if a little Fe₂(SO₄), solution is added the Ag goes Very finelyinto solution without heating. divided Ag, obtained by reduction of Ag salts by milk-sugar and alkali, is dissolved by very dilute H₂SO₄Aq (Carey Lea, Zeit. für anorg. Chem. 8, 180). Friedheim (B. 19, 2554; 20, 307) says that Ag dissolves in dilute H₂SO₄Aq containing KMnO,Aq.—18. Chromic acid (i.e. solution of CrO, in water) produces red Ag₂CrO₄; HNO,Aq, S.G. 1.2, in which K₂CrO₄ is dissolved, gives a red deposit on the surface of Ag; this test is used to distinguish Ag from other white metals, such as Sn.—19. Ag in pieces is scarcely acted on by hydrochloric acid, even when hot and conc.; very finely-divided Ag is said to form a little AgCl, with evolution of H. Hydrofluoric acid is without reaction.-20. Ag is not acted on by molten alkalis or alkaline carbonates; hence KOH, &c., may be fused in dishes of Ag.—21. Ag

alloys with many metals (v. Silver, Alloys of).
Silver, alloys of. Ag forms alloys with many metals. They are generally formed by heating the metals together; in a few cases—s.g. alloy with Ba or Mn—they are formed by heating Ag with metallic oxides and C.

heating Ag with metallic oxides and C.

With aluminium. An alloy of 100 parts
Al with 5 parts Ag is harder and more easily

polished than Al, but otherwise has most of the properties of Al; it is used for making the pans, &c., of balances. Hirzel (J. 1858. 137) obtained alloys corresponding in composition with the formulæ Ag,Al, Ag₂Al, and AgAl (v. also Tissier, C. R. 43, 885; 52, 931; Debray, C. R. 43, 925).

With antimony, and with arsenic; v. SILVER

ANTIMONIDE and ARSENIDE, infra.

With copper. Ag becomes harder, more elastic, more capable of being polished, and more slowly worn away when alloyed with Cu without losing much in malleability and duc-Alloys with less than 50 p.c. Cu are tility. white, with more than 50 p.c. Cu the colour becomes reddish. The S.G. of a Ag-Cu alloy is rather smaller than the mean of those of the constituent elements (v. Karmarsch, D. P. J. 108, 278). The composition of these alloys is rarely identical throughout a mass (v. Levol, A. Ch. [5] 36, 193). Alloys of Ag and Cu are used for making silver vessels and ornaments, and for coinage; the standard silver coinage alloy of the United Kingdom contains 925 parts Ag and 75 parts Cu per 1,000.

With mercury. The amalgams of Ag are formed (1) by the action of Hg on AgNO₂Aq, (2) by the action of finely-divided Ag on HgCl₂Aq, (3) by dissolving Ag in Hg. The compositions of many of these amalgams can be represented by formulæ, but it is very doubtful whether any of them are definite compounds. For accounts of these amalgams v. Campani, J. 1870. 373; Joule, C. J. [2] 1, 378; Crookewitt, J. 1847-48. 393; Malaguti a. Durocher, Ann. M. 17, 336;

Gay-Lussac, A. Ch. 58, 218.

Alloys of Ag with the following metals have been described:—Ba (Clarke, G. A. 62, 373; Lampadius, S. 15, 146); Cd (Wood, C. N. 6, 135); Au (G. Rose, P. 23, 181; Levol, A. Ch. [3] 15, 55; 27, 310); Ir; Fe; Pb (Levol, A. Ch. [3] 39, 173); Mg; Mo; Ni; Pd (Graham, J. 1868. 144); Pt; K; Rh; Na; Sn; Zn (v. Wright a. Thompson, Pr. 48, 25).

Silver, antimonide of. The mineral dys-

Silver, antimonide of. The mineral dyskrasite has approximately the composition Ag_sSb; when heated to dull redness in H, the compound Ag_sSb is said to remain (von Bonsdorff, S. 34, 225). The compound Ag_sSb is also obtained by ppg. AgNO_sAq by Sb hydride; using conc. AgNO_sAq the compound Ag_sSb.3AgNO_s was formed (Poleck a. Thümmel, B. 16,

2435).

Silver, arsenide of. No definite compound of Ag and As has been obtained. By heating Ag powder with As about 16 p.c. of As is absorbed by the Ag to form a grey, brittle soil (Gehlen). By passing As hydride into conc. AgNO, Aq the compound Ag, As, 3AgNO, was obtained (P. a. T., Lc.).

Silver, bromide of, AgBr. (Argentic bromide.) Formula probably molecular, from analogy of AgCl. Occurs native in Chili and Mexico.

Preparation. — HBrAq or alkali bromide solution is added to AgNO₂Aq till ppn. is complete; the pp. is washed with water in the dark and dried.

Properties.—A pale-yellow solid; melts at Ag pyrotartrate, filtering from the brown pp. c. 434° (Carnelley, C. J. 29, 489). S.G. 6·245 at 0°, 5·595 at m.p. (Rodwell, Pr. 31, 291). S.H. a dark-brown, metal-like pp., probably AgC (P. 415° to 98°) ·07891 (Regnault, A. Oh. [8] 1, 129). 86, 28; cf. Regnault, A. 19, 153).

For effects of heat on AgBr v. Rodwell (Pr. 25, 280). Obtained in octahedra by crystallising from HBrAq. Solubility of AgBr dried at 100° in NH, Aq S.G. 986, = 051 (Pohl, J. pr. 82, 52). Granular AgBr is slightly sol. water above 50° flocculent AgBr is slightly sol. water above 33° (Stas, A. Ch. [5] 3, 289). Sol. hot Hg(NO₂),Aq; AgBr crystallises out on cooling (Debray, C. R. 70, 995). Sol. solutions of alkali bromides. H.F. [Ag,Br] = 22,700 (Th. 3, 381). According to Berthelot (Bl. [2] 39, 19), the reaction [Ag,Br] has the value 23,700 when crystalline AgBr is formed, and the value 20,000 when the AgBr is amorphous (in both cases using liquid Br). B. also says that in the formation of AgBr by adding KBrAq to AgNO, Aq 20,700 cals. are produced, and that when AgNO, Aq is added to KBrAq only 17,600 cals. are produced at first, but that temperature then rises until 20.600 cals. are evolved; B. thinks that two forms of AgBr are formed, one more stable than the other. Vogel (B. 16, 1160) distinguishes two modifications of AgBr, one ppd. from aqueous solutions and very sensitive to blue rays, the other ppd. from solutions in absolute alcohol and sensitive to the violet rays. Stas (A. Ch. [5] 3, 289) distinguishes six forms of AgBr:—(1) flocculent, white, by adding dilute alkali bromide solution to excess of cold AgNO₂Aq; (2) flocculent, yellow, by using excess of bromide as pptant.; (3) powdery, white, by shaking (1) or (2) with water; (4) intense yellow, by heating (4); (5) granular, white-yellow, by pouring the flocculent or powdery variety, shaken with water, into boiling water, also by adding very dilute NH, BrAq to boiling AgNO, Aq (1:1000); (6) pure yellow, crystalline variety.

Reactions and Combinations. — 1. AgBr darkens rapidly when exposed to sunlight. Carey Lea says that the dark product of the action of light on AgBr reacts with HNO₂Aq to re-form AgBr, and Ag which dissolves in the acid (Am. S. [3] 15, 189). The action of light on the Ag haloids is treated more fully under Silver chloride (q. v.).—2. Heated in chlorine, AgCl and Br are produced. — 3. Dissolves slowly in hot silver nitrate solution; the salt AgBr.AgNO₃ separates on cooling (Risse, A. 111, 43; Riche, A. 111, 39).—4. Combines with silver chloride and iodide to form compounds Ag_aBr_bI_cCl_d (v. Silver iodobromochiorides, p. 470).

AgBr resembles AgCl in most of its reactions; NH, is not absorbed by dry AgBr (Rammelsberg, P. 55, 248).

Silver, bromo-iodochlorides of, v. SILVER IODOBROMOCHLORIDES (p. 470).

Silver, carbides of. Gay-Lussac noticed that Ag absorbed c. 3 p.c. C when heated with lampblack (A. Ch. [2] 58, 222); Ag₄C requires 2.7 p.c. C. A yellow powder, approximately agreeing with the composition Ag₂C, was obtained by Gerhardt and Cahours (A. Ch. [3] 1, 76) by heating Ag cuminate; the same compound was probably produced by Liebig by heating AgCN (A. 38, 21). By heating an aqueous solution of Ag pyrotartrate, filtering from the brown pp. produced, and boiling the pp., Berzelius obtained a dark-brown, metal-like pp., probably AgC (P. 36, 28; cf. Regnault, A. 19, 153).

m n 2

Silver, chlorides of. The isolation of a chloride of Ag containing less Cl than AgCl is not yet certain. Cl is given off when AgCl darkens in sunlight, but whether the darkened product is a chloride or an oxychloride is not finally determined (v. Action of light on silver chloride, infra).

SILVER SUBCHLORIDE ? Ag₂Cl. (Argentous chloride.) According to Güntz (C. R. 112, 861) Ag₂Cl is formed by passing dry HCl over Ag₂F (v. Silver subfluoride, p. 469) as long as darkening occurs and the solid continues to gain in weight. The compound is also said to be formed by passing vapour of CCl4, SiCl4, or PCl3 over dry Ag.F. A subchloride is formed, according to von Bibra (J. pr. [2] 12, 55), by digesting with conc. HClAq the product of the reduction of Ag citrate by H at 100°; after drying, the black solid is said to have the composition Ag. Cl., and with NH₃Aq to give Ag and AgCl.3NH₃. Newbury (Am. 8, 196) failed to obtain any subchloride by this method; he also found that the reduction of Ag citrate by H at 100° does not yield any definite compound, but probably a mixture of unchanged citrate and Ag.

Silver Chloride AgCl. (Argentic chloride.) Mol. w. 143.03. V.D. 80 at c. 1700° (Biltz a. Meyer, B. 22, 725). Melts at c. 457° (Carnelley, C. J. 29, 489). S.G. at $0^{\circ} = 5.505$; at m.p. =4.919 (Rodwell, Pr. 31, 291). S.H. (13° to 98°) .09109 (Regnault, A. Ch. [3] 1, 129). For expansion of AgCl by heat v. Rodwell (Pr. 25, 280). H.F. [Ag,Cl] = 22,700 (Th. 3, 381); 29,200 (Berthelot, Bl. [2] 39, 19). For solubilities in various liquids v. Properties.

Occurrence. — Native, as hornsilver, in Siberia, Mexico, Peru, the Harz, Alsace, &c.

Preparation.—1. By adding dilute HClAq, or solution of a chloride, to solution of any salt of Ag except Ag₂S₂O₃, washing the white curdy pp., drying, and heating to incipient fusion. Obtained in octahedral crystals by dissolving in large quantity of conc. HClAq and evaporating.—2. By heating Ag to redness in a stream of Cl or HCl gas.—3. By fusing NaCl with finely-divided Ag, washing, and drying.—4. Very slowly by the action of HClAq on Ag in the presence of air.

Properties.—A white solid. According to Cooke (Am. S. [3] 21, 220) AgCl is very slightly sol. cold water, and very distinctly sol. boiling water; he found that 1 4561g. AgCl washed with 66 litres boiling water lost '2241 g. = 15.39 p.c.; the solvent action was entirely prevented by the addition of '05 g. AgNO₃ per litre of water. Stas (C. R. 73, 998) distinguished four forms of AgCl: (1) gelatinous; (2) curdy, flocculent; (3) powdery; (4) granular, crystalline, or fused. The curdy, flocculent form is the most sol. in water; it is formed by ppg. cold dilute AgNO₃Aq by a chloride solution; on shaking it changes to the less sol. powdery variety. The granular, crystalline, or fused form is insol. cold water (10 million parts cold water dissolve 1 part) but it begins to dissolve at 30° (cf. Berthelot, A. Ch. [5] 29, 241, for the heats of solution in KONAq of different preparations of AgCl). AgCl is sol. conc. HClAq and in saturated solutions of various metallic chlorides; Vogel (C. C. 5, 578) gives the following table:—

100	C.C. 00	old HOLA	q 8.G.	1.165	dissolve	-298	g	AgO
,		ling "		99	99	.260	10	39
,	, cono.	HOLAq	diluted	with				
		t. H ₂ O			,,,	*056	99	,
		HÖLAq	diluted	with				
		ts, H ₂ O			79	-018	m	90-
•	, conc.	HOLAQ	diluted	i with		-0000		
	8 p	ts. H.O			19	-0089	**	90-
1	, cone.	HCIAq	dilute	1 With	l	.000#		
	5 p	ts. H ₂ O			99	•0035	**	91
,	, satur	ated Ba(يال.AQ		99	.0148	91	10"
1	, ,	, SrC	l.Aq		99	0884	97	r
•	91		l,Aq		29	-0930	"	75
1	, ,	, Mg	Ol, Aq		99	•171	"	97
1	9		DLAq		79	•095	**	99
,	, ,	KO			29	.0472	"	"
		, NH	OlAa			·1575	**	**

One part AgCl dissolves in 50,000 parts boiling cone. HNO, Aq (Thorpe, C. N. 25, 198). NH, Aq dissolves AgCl; 1 litre NH, Aq S.G. 924 dissolves 69.5 g. freshly ppd. AgCl, according to Millon and Comaille (C. R. 56, 309). AgCl crystallises from solutions in NH,Aq, on evaporation; explosive silver is obtained from boiling solutions (v. Silver nitrides, p. 470). According to Terreil (Bl. [2] 51, 598), crystals of AgCl.2NH, separate from a hot solution of AgCl in NH.Aq; these crystals lose NH, in the air. AgOl dissolves easily in Na₂S₂O₃Aq, forming Ag₂Na₄(S₂O₃)₃Aq; AgCl also dissolves readily in KCNAq forming AgK(CN)2, and in NH4HSO3Aq forming double Ag-NH, sulphites (v. Svensson, B. 4, 714). hot solution of AgNO, Aq dissolves AgCl; Risse says that a double compound of AgNO, and AgCl separates on cooling (A. 111, 39), but Debray asserts that simple solution occurs (C. R. 70, 995). Hg(NO₃)₂Aq also dissolves AgCl; Wackenroder (A. 91, 317) and Field (J. 1857. 255) say that AgCl separates on cooling a hot saturated solution; Stas (A. Ch. [5] 3,

180) found that a little AgNO, is formed.

Reactions.—1. AgCl is reduced to Ag by many metals, e.g. Zn or Fe, in presence of dilute acid; Cu reduces AgCl in presence of NH₂Aq, with formation of Ag and CuO.H₂O.—2. Heated in hydrogen, or with carbon, or with carbon, or with carbon of Ag.—3. Solution of hydriodic acid forms AgI and HClAq.—4. Caustic alkali solutions pro-

duce Ag₂O.

Action of light on silver chloride. AgCl darkens when exposed to sunlight in ordinary During the darkening Cl is given off air. (Scheele; von Bibra, J. pr. 122, 39; Hitchcock, Am. 11, 474; Béchamp, Bl. [3] 6, 836; Richardson, C. J. 59, 536; Baker, C. J. 61, 728). Only a small portion of the AgCl is changed to the dark compound (v. R., l.c.; B., l.c., got 09 g. of the dark body from 51 g. AgCl). AgCl does not darken in thoroughly dried oxygen (B., l.c.; nor in dried air according to Carey Lea, Zeit. für anorg. Chem. 3, 184). AgCl does not darken in vacuo, when thoroughly dry (Abney; B., l.c.). Carey Lea (l.c.) found that AgCl darkened under dry petroleum; Baker noticed darkening under benzene in absence of air, and he showed that the dark substance was Ag. B. (l.c.) noticed darkening, with formation of Ag, under CCl., but he proved that if the OCl, contained no trace of free Cl darkening did not occur. For the influence of HCl and chlorides in water upon the rate of darkening of AgCl in the water v. Richardson (l.c.). The dark substance was supposed by some to be a subchloride, but Baker has shown that it contains O (Hodgkinson had formerly arrived at this conclusion; v. Meldola's Chemistry of Photography [1889], p. 856). B. kept pure AgCl, dried at 150°, in sunshine, in a current of air freed from CO, and dried by H₂SO₄, until it darkened; he measured the O obtained from the darkened product, by decomposing by Cl, and also determined the Ag and Cl; the results agreed approximately with the formula Ag₂OCl; the atomic ratio of Ag to Cl found in various experiments was almost exactly 2:1, and of Ag to O was nearly 5:2; the darkened product dissolved in conc. KClAq, on dilution AgCl was ppd. and the solution had a slightly alkaline reaction, as would be expected if an oxychloride were present in the dark-coloured substance. B. also found that a little H₂O was obtained by heating the dark substance in dry H; Richardson (C. J. 59, 536) failed to obtain any water by treating darkened AgCl in this way; R. also failed to get O from the darkened AgCl by heating it to the m.p. in connection with a Sprengel pump. B. (l.c.) found that the darkcoloured product became white when left in the dark for some days, and that a small quantity of O was absorbed during this change; he, therefore, concludes that two oxychlorides of Ag are formed, one of which is dark-coloured and absorbs a little O to form another, white, oxychloride. Carey Lea (Am. S. [3] 38, 356) thinks that the darkening action of light on AgCl is preceded by the formation of what he calls a photochloride. The photochloride is supposed to contain less Cl than the normal chloride, and probably to be a compound of AgCl with a subchloride or oxychloride; it is extremely sensitive to the action of light, and undergoes many changes of colour when exposed to light (Am. S. [3] 33, 349, 480, 489; 34, 38; and full abstract in C. J. 54, 1). For the application of the effects of light on AgCl to photography v. Photographic CHEMISTRY, this vol. p. 154; and Photography, in Dictionary of Applied Chemistry, vol. iii.

Combinations.—1. With several metallic chlorides, especially the alkaline chlorides; generally forming compounds of the type AgCl.MCl. Wells and Wheeler (Am. S. [8] 44, 155) obtained the compound AgCl.2CsCl.— 2. With ammonia gas. According to H. Rose (P. 20, 157), 2AgCl.8NH₃ is formed. Isambert (C. R. 64, 1259; v. also Horstmann, B. 9, 749) studied the vapour-pressures of the NH, given off, and concluded that two compounds are formed when NH, is absorbed by AgCl-viz. 2AgCl.3NH, and AgCl.3NH, (v. curves given in Dissociation, vol. ii. p. 397). Bodländer (Z. P.C. 9, 730) measured the reduction of the freezingpoint of water produced by the compound 2AgCl.3NH_a; as this reduction was much larger than the normal, he concluded that the compound is dissociated in aqueous solution; a similar conclusion was come to by measuring the electrical conductivities of aqueous solutions. The dissociation cannot have been into NH, and AgCl, says B., else AgCl would have separated out; hence the dissociation must be electrolytic, and the solution must contain Ag and Cl as free ions. This conclusion was confirmed by finding that PbAc,Aq ppd. PbCl₂ from 2AgCl.8NH₂ in NH,Aq, KIAq ppd. AgI, and (NH₄)₂SAq ppd. Ag₂S; confirmatory evidence

was also found in the fact that addition of NH_Cl or AgNO, reduced the solubility in water of 2AgCl.8NH, in keeping with the law regarding the effect on the solubility of an electrolyte of the presence of one of its ions. B. concludes that 2AgCl.3NH, is a definite compound, which is electrolytically dissociated in aqueous solution. -8. With silver iodide and bromide; v. SILVER IODOBROMOCHLORIDES, p. 470.

Silver, chloro-iodobromides of; v. SILVER

IODOBROMOCHLORIDES, p. 470.

Silver, cyanide of; v. vol. ii. p. 346.

Silver, ferricyanide of; v. vol. ii. p. 340. Silver, ferrocyanide of; v. vol. ii. p. 837.

Silver, fluorides of. Two fluorides of Ag probably exist, Ag₂F and AgF, although there is still some doubt as to the isolation of the first

of these. SILVER SUBFLUORIDE (? Ag₂F). (Argentous

fluoride.) By electrolysing saturated AgFAq, using Ag electrodes, and a current so strong that the liquid became hot, Güntz (C. R. 110, 1887) noticed the separation of brass-coloured crystalline plates at the negative pole (if a weaker current was used, Ag was deposited). Treatment of these crystals with water gave AgFAq and Ag, the ratio of Ag separating to Ag going into solution being practically unity; hence G. gives the formula Ag₂F to the yellow crystals. G. says that the same compound is obtained by heating finely-divided Ag with AgFAq in a sealed tube to nearly 90°. Ag₂F does not change in dry air; it decomposes slowly in moist air, and rapidly in water. Heated in dry HCl, CCl,, PCl₃, &c., Ag₂Cl is said to be formed (G., C. R. 112, 861).

SILVER FLUORIDE AgF. (Argentic fluoride.) Formula probably molecular, from analogy of AgCl. Prepared by dissolving Ag₂CO₂ that has been freshly ppd. and well washed, in HFAq, evaporating to dryness in a Pt dish, with constant stirring, dissolving the black residue in water, filtering, and evaporating in vacuo over H_2SO_4 , in the dark (Moissan, Bl. [3] 5, 456; cf. Güntz, A. Ch. [6] 3, 42). A yellow, transparent, elastic solid; sol. water; melts at c. 435° (M., l.c.). S.G. 5.852 at 15.5° (Gore, Pr. 18, 157). Fremy (A. Ch. [3] 47, 39) obtained large colourless prisms, to which he gave the formula AgF. 2aq, by evaporating a solution of Ag₂CO₂ or Ag₂O in HFAq; these crystals lost 2aq in vacuo, but some exyfluoride was always formed (cf. Marignac, Ann. M. [5] 15, 270). Heated in air, AgF is decomposed, by the moisture present, to Ag, O, and HF, according to Gore (l.c.). AgF interacts violently with POCl₂, PCl₃, SiCl₄, and BCl₃, giving AgCl and fluoride of P, Si, or B (Moissan, l.c.). Molten AgF attacks vessels of Ag or Pt. The dry salt is said to absorb c. 800 times its volume of NH, (Gore, l.c.). Heating with I produces AgI and IF, (Gore, Pr. 30, 70; where a few other reactions of compounds with AgF are given).

Silver, fulminating; v. SILVER NITRIDES, p. 470; Silver fulminate (q. v., vol. ii. p. 317) is sometimes called fulminating silver.

Silver, hydroxides of; v. SILVER OXIDES AND

HYDRATED OXIDES, p. 470.
Silver, iodides of. The isolation of an iodide other than AgI is very doubtful.

SILVER SUSIDDIDE (? Ag₂I). Güntz (C. R. 112, 861) states that this compound is formed by the interaction of HI and Ag₂F; but no analyses are given.

SILVER IDDIDE AgI. (Argentic iodide.) Formula probably molecular, from analogy of AgCl. Melts at c. 530° (Carnelley, C. J. 29, 489); at 527° (Rodwell, Pr. 25, 280). S.G. at 0° 5.675, at 527° = 5.522; maximum density at 142° (R., l.c.). V.D. said to be 214 (Dewar a. Scott, B. A. 1881. 597). S.H. (15° to 98°) '06159 (Regnault, A. Ch. [3] 1, 129; v. also Bellati a. Romanese, Pr. 34, 104). H.F. [Ag,I] = 18,800 (Th. 3, 381); 14,300 for crystalline AgI (Berthelot, Bl. [2] 39, 18).

Occurrence.—As iodite in Peru, Mexico, the Harz, Spain, &c.; forms hexagonal tablets; also in combination with AgBr as iodobromite

in small quantities in Nassau.

Preparation.—1. By adding HIAq, or solution of an iodide, to AgNO₃Aq, washing, and drying.—2. By the interaction of finely-divided Ag with HIAg, H is evolved at ordinary temperatures and AgI formed; a solution saturated by heating HIAq with Ag deposits crystals of AgI.xHI on cooling (v. infra, Combinations, No.2), and the liquid decanted from these deposits hexagonal crystals of AgI on standing in air.—3. Crystals of AgI are also obtained by boiling saturated AgNO₃Aq with HgI., and allowing to cool (Field, J. 1857. 255).—4. The interaction of a mixture of HClAq and HIAq with Ag produces only AgI, according to Deville (A. 101, 197).

Properties.—As obtained by adding alkali lodide to excess of AgNO, Aq, AgI is a curdy pp., yellow, with a slight orange tint; by adding AgNO, Aq to excess of alkali iodide, a pure yellow, powdery pp. of AgI is formed; the former pp. is sensitive to light, the latter, after washing in the dark, is not discoloured by light. AgI melts to a yellow liquid, which becomes red and then dark-red as temperature rises; the liquid solidifies to a yellow, horny, crystalline mass. Rodwell (Pr. 25, 280) found that AgI contracted considerably at the moment of solidification; regular contraction then took place to 142°, whereat there was sudden and considerable expansion accompanied by change from the amorphous to the crystalline form; this was followed, as temperature fell, by slight expansion (for measurements v. R., l.c.). AgI exists in two forms: above 142° up to the m.p. it is yellow, transparent, and flexible; below 142° it is pale green, crystalline, opaque, and brittle (R., l.c.). AgI is nearly insol. NH, Aq; according to Martini (S. 56, 153) 1 part AgI requires 2560 parts NH₂Aq, S.G. 96, for solution. AgI dissolves in Na₂S₂O₃Aq, in KCNAq, and in NaClAq; it dissolves readily in hot Hg(NO₃)₂Aq. and crystallises therefrom on cooling (v. also Combinations).

Reactions.—1. Heated in chlorine, AgCl is formed.—2. Reduced by contact with metals, such as Zn or Fe, in presence of acids. Most of the reactions of AgI closely resemble those of AgCl.

Combinations.—1. Dry AgI absorbs ammonia; Rammelsberg (P. 48, 170) says that 2AgI.NH, is formed (for dissociation-pressures of this compound v. Isambert, C. R. 64, 1259). By heating

AgI with excess of NH, Aq in a sealed tube at 100°, Terreil obtained crystals of AgI.2NH, (Bl. [2] 41, 598).—2. AgI dissolves in hot conc. hydriodic acid solution; the liquid deposits large, colourless, unstable crystals of AgI.HI (?) (Deville, C. R. 42, 895); Berthelot obtained transparent crystals of 3AgI.HI.7H₂O (A. Ch. [5] 28, 89).—3. According to Preuss (A. 29, 328) a solution of AgI in a hot solution of mercuric nitrate deposits crystals of 4AgI.2Hg(NO₂)2. aq on cooling.—4. AgI dissolves in silver nitrate solution containing more than 3 p.c. AgNO. (Vogel); water re-ppts. AgI. For accounts of different compounds obtained v. Kremers (J. pr. 71, 54), Stürenberg (Ar. Ph. [2] 143, 112), Weltzien (A. 95, 127), Riche (A. 111, 39), Risse (A. 111, 43). AgI also combines with silver nitrate and lead nitrate, and with antimonide, arsenide, phosphide, and sulphide of silver (v. Poleck a. Thümmel, Ar. Ph. [3] 22, 1).—5. With alkali iodides and alkaline earth iodides (v. Boullay, A. Ch. [2] 34, 377; Berthelot, Bl. [2] 39, 24; Johnson, C. N. 37, 110; Maxwell Simpson, Pr. 27, 120).

Silver, iodobromochlorides of. Rodwell (Pr 25, 292) prepared various compounds of AgI AgBr, and AgCl by melting the constituents together. R. studied especially the effects of heat on the S.G., change of volume, and m.p. of these bodies. The formula xAgI.yAgBr.zAgCl expresses the composition of the compounds examined; x varied from 1 to 4, y was always 1,

and z varied from 1 to 2.

Silver, nitrides of. Raschig (A. 233, 93) has shown that the explosive substance first prepared by Berthollet, by the action of conc. cold NH₃Aq on Ag₂O, and supposed by B. to be a compound of Ag₂O and NH₃, is a nitride of Ag, Ag₃N, mixed with finely-divided Ag. Ag₃N is best prepared by allowing a conc. solution of Ag₂O in NH₃Aq to stand in air, or by adding alcohol. Ag₃N dissolves in KCNAq, forming AgCN, KHOAq, and NH₃.

The Ag salt of HN, may be called silver nitride; v. Hydrazoic Acid, vol. iii. p. 559. Silver, nitroprusside of, v. vol. ii. p. 341.

Silver, oxides and hydrated oxides of. Only one oxide of Ag has been certainly isolated; the existence of any hydrated oxide is doubtful.

SILVER OXIDE Ag₂O. (Argentic oxide.) Mol. w. not known.

Formation.—1. By heating Ag much above the temperature of volatilisation in the O-H flame, using excess of O (Debray, C. R. 66, 735).—2. By passing an electric current through KNO₄Aq, using a plate of Ag as positive electrode (Wöhler, A. 146, 264).—3. By boiling freshlypd. AgCl with KOHAq, S.G. 1.25 to 1.3 (Mohr, A. 66, 65).

Preparation.—KOHAq is added in slight excess to AgNO₃Aq, the pp. is thoroughly washed with hot water, best in air freed from CO₂, and

dried at c. 80°.

Properties.—A black powder (brownish when ppd.), with an unpleasant, metallic taste. S.G. 7.52 (Schröder, B. 9, 1888); 7.15 (Playfair a. Joule, C. S. Mem. 3, 84): 7.25 (Boullay, A. Ch. [2] 43, 266). H.F. [Ag²,O] = 5,900 (Th. 3, 381). Slightly sol. water; 1 part dissolves in 3,000 parts water according to Bineau (C. R. 41, 509), but according to Abl in 96 parts water at 18.50

(Oester. Zeit. f. Pharm. 8, 201 [1884]). Sol. NH₂Aq (for so-called explosive silver, formed by dissolving in conc. NH₂Aq, v. SILVER NITHIDES, p. 470). Also sol. aqueous solutions of alkali thiosulphates, chlorides, and cyanides. Sol. aqueous MeNH₂ and EtNH₂ (Wurtz, A. Ch. [3] 30, 453). Moist Ag₂O absorbs CO₂ from the air, forming Ag₂CO₃, and reacts with many salts like the alkalis (v. Reactions). Ag₂O is decomposed to Ag and O by heat (v. Reactions).

to Ag and O by heat (v. Reactions).

Reactions and Combinations. -1. Decomposed to Ag and O by heat; Carnelley and Walker (C. J. 53, 79) say that O begins to come off at c. 100°, rapid decomposition occurs at c. 270°, and reduction is complete at 300°-340° (v. also H. Rose, P. 85, 317; Joulin, Bl. [2] 19, 349; cf. Hydrated silver oxide, infra). - 2. Antimony sulphide, arsenic sulphide, finely-powdered sulphur and selenion, amorphous phosphorus, and some organic compounds, are ignited by rubbing with Ag₂O (Böttger, J. 1863. 284).— 8. Reduced to Ag by heating to 100° in hydrogen (Wöhler, A. 30, 4).—4. Finely-divided Ag₂O under water is reduced to Ag by cadmium, copper, tin, and zinc (not reduced by Fe or Hg) (Fischer, P. 10, 605).—5. AgI and HIO, are formed by the reaction of iodine with Ag₂O suspended in water (Naquet, Bl. 1860. 123); with chlorine the products are AgCl, HClOAq, AgClO, and AgClO₃ (Stas).—6. Moist Ag₂O decomposes many metallic salts in solution, ppg. metallic hydroxides, e.g. salts of Bi, Cr, Co, Cu, Fe, Hg (v. H. Rose, B. B. 1857. 245).—7. The moist oxide removes halogens from many halogencontaining organic compounds, replacing the halogen by OH (e.g. 2NMe,I+Ag₂O.H₂O halogen by OH (e.g. $2NMe_4I + Ag_2O.H_2O$ = $2NMe_4OH + 2AgI).$ —8. Ag_2O reacts with acids as a decidedly basic oxide, forming salts Ag₂X, where $X = SO_4$, $2NO_3$, &c.—9. A little AgNO₂ is said to be formed when nitric oxide is passed into water with Ag₂O suspended therein (Sabatier a. Senderens, C. R. 114, 1476).—10. Ag₂O combines with lead monoxide; by allowing a mixture of PbO.xH₂O and freshly ppd. Ag₂O to stand in contact with NaOHAq, a compound was obtained, probably 2Ag₂O.PbO (E. Aston, C. J. 59, 1093).

HYDRATED SILVER OXIDE (? AgOH). According to H. Rose (P. 85, 904), no compound of Ag₂O and H₂O exists. By ppg. an alcoholic solution of AgNO, by KOHAq at -40°, Bruce obtained a white flocculent pp. which he supposed to be a hydroxide of Ag (no analyses are given); the pp. darkened on heating, forming Ag₂O and H₂O according to B. (C. N. 50, 208). Carey Lea (Am. S. 1892. 249) found that ppd. Ag₂O retained some water after drying at 100° for 40 hours, and that all water was removed at 160°-165°, but some O went off also. and Walker (C. J. 53, 79) say that the product of drying ppd. silver oxide in air for 6 months had the composition AgOH, and that this was scarcely decomposed at 100°, but that above this temperature it gave off water and a little O, and was gradually changed to Ag₂O mixed with a little Ag. Some reactions of moist Ag₂O point to the existence of such a compound as AgOH (v. Silver oxide, Reactions, Nos. 6 and 7).

Oxnde of silver with less oxigen than Ag.O. (Silver suboxide.) Many experiments are recorded on the interactions of reducing

agents and Ag salts whereby salts of a lower oxide than Ag₂O were supposed to be formed, and the corresponding oxide was supposed to be ppd. from these salts by alkali. Wöhler (A. 30, 1) reduced Ag citrate by heating in H; W. a. Rautenberg reduced ammoniacal solutions of chromate, molybdate, and tungstate of Ag (A. 114, 119); W. reduced arsenate and phosphate of Ag by FeSO₄ (A. 146, 263); H. Rose reduced AgNO, in NH, Aq by ferrous and manganous salts (P. 101, 264, 321, 497; v. also Weltzien, A. 142, 105). More recent experiments have shown that the products of such reductions in presence of alkali are mixtures containing Ag (v. Pillitz, Fr. 27, 496; Newbury, Am. 8, 196; Muthmann, B. 20, 983; Bailey a. Fowler, C. J. 51, 416).

Faraday (Q. J. S. 4, 268) supposed that a suboxide of Ag was formed by the action of air on an ammoniacal solution of Ag₂O; but Bailey and Fowler (l.c.) have shown that the substance is a mixture of Ag₂O with a little explosive silver.

The result of a long series of experiments by O. v. d. Pfordten (B. 18, 1407; 20, 1458; 21 2288, 3375) on reducing AgNO, Aq by H, PO, Aq, alkali bisulphites, and tartaric acid, is that the black pp. thus obtained is not a suboxide of Ag (for some time it was asserted to be Ag₄O); the accuracy of this result has been strengthened by the critical experimental work of Bailey (C. N. 55, 263; C. J. 51, 466), and especially of Friedheim (B. 20, 2554; 21, 307). F. regards the supposed Ag₄O as a mixture of finely-divided Ag with more or less Ag 0, or with some organic compounds. V. d. P. (B. 21, 2288, 3375) thinks his latest work points to the black substance being a compound of Ag, H, and O in the ratio 4Ag:2H:O; he gives it the formula Ag. H.O, or Ag. OH.H, and calls it silver hydrate. According to Guntz (C. R. 112, 861) Ag,O is produced by the action of steam on Ag₂F at 160°

The question of the existence and isolation of a compound with less O than Ag₂O cannot be regarded as settled.

Oxides of silver with more oxygen than Ag₂O. By electrolysing conc. AgNO₂Aq, using Pt as positive and Ag as negative electrode, Ritter obtained a black deposit on the Pt; to this black substance R. gave the formula Ag₂O₂, and described it as crystallising in octahedra, S.G. 5.474, giving up half its O when carefully heated, with NH, Aqevolving N, and with H,O,Aq producing Ag, H₂O, and O. A similar production of what seemed to be a peroxide of Ag was noticed by Fischer (J. pr. 83, 237) and by Wöhler (A. 146, 264). Bottger (B. 6, 1398) noticed that the peroxide produced by electrolysis formed explosive Ag and rapidly evolved N, with NH,Aq. Ag peroxide is formed by the action of ozone on Ag, according to Schönbein (J. pr. 41, 321); and of ozone on Ag₂O according to Schiel (J. 1864. 118). Ag peroxide is an energetic oxidiser; it sets fire to H₂S, and Au sulphide when rubbed

therewith (v. Böttger, l.c.).

The analyses made by Fischer (J. pr. 33, 237),
Mahla (A. 82, 289), and Gmelin (J. 6, 105) of the
peroxide showed the presence of H.O and AgNO,;
their preparations were probably impure. The
analyses of Wallquist (J. pr. 31, 179) agreed well

with the formula AgO (or Ag₂O₂).

The supposed peroxide of Ag, prepared by electrolysing AgNO₂Aq, is regarded by Berthelot (Bl. [2] 84, 138) as a compound of AgNO₂ with an oxide to which he gives the formula Ag₄O₂. By the interaction of H₂O₂Aq with Ag₂O, Berthelot (l.c.) obtained black flocks, which were separated from Ag by washing; to this substance B. gives the formula Ag₄O₂ as the result of a not very complete analysis; he looks on it as probably identical with the product of the interaction of moist ozone and Ag or Ag₂O. This substance loses O when dried over H₂SO₄; with acids it gives salts of Ag₂O and evolves O.

Silver, oxychlorides of. The action of light on AgCl probably produces two oxychlorides of Ag; v. Silver chloride, Action of light on,

p. 468.

Silver, exyfluoride of. To the yellow crystals obtained by evaporating an aqueous solution of AgF, Pfaundler gives the composition AgF.AgOH

(W. A. B. 46 [2] 259).

Silver, phosphides of. Molten Ag absorbs P; according to Pelletier (A. Ch. 13, 101), some of the P remains in combination on cooling, but Warren (C. N. 56, 113) says that all except c. 002 p.c. separates as the Ag cools. According to Emmerling (B. 12, 152), a compound AgP is formed by heating Ag and P in a sealed tube to a dark-red heat; on heating the compound P is given off and Ag remains (v. also Landgrebe, S. 60, 128). By heating powdered Ag in vapour of P, Schrötter obtained a grey solid, S.G. 4·63, to which he gave the composition Ag₂P₃ (J. 1869. 247).

Fresenius and Neubauer (Fr. 1, 340) obtained a pp. said to contain Ag phosphide and Ag, by leading PH₃, or CO₂ laden with vapour of P, into AgNO₂Aq; using PH₃ diluted with CO₂, Poleck and Thümmel (B. 16, 2435) obtained a

compound, probably Ag,P.8AgNO,

Silver, photosalts of. Carey Lea (Am. S. [8] 83, 349, 480, 489; 34, 33; abstract in C. J. 54, 1) gives the name of photosalts to variouslycoloured compounds of Ag with the halogens which are very sensitive to light. These bodies are regarded by C. L. as identical with the substances that form the latent images on exposed photographic plates. The photosalts are obtained by the action of reducers, such as FeSO, Aq or FeCl, Aq, on the normal Ag haloids, by reducing Ag salts by such bodies as dextrose or tannin in presence of alkali and treating the products with HClAq, or by treating finelydivided Ag with the haloid salts of Cu or Fe and certain other halogenating reagents. The photosalts are regarded by C. L. as compounds of the normal haloids with sub-haloids; from the methods of preparation it seems fairly certain that the coloured bodies called photosalts contained various substances besides Ag and halogen.

Silver, platinocyanide of, v. vol. ii. p. 344. Silver, platinosulphocyanide of, v. vol. ii.

p. 351.

Silver, salts of. Compounds derived from soids by replacing H by Ag. These salts belong to the type AgX, where X = NO₂, ½SO₄, ½CO₅, ½PO₄, &c. Subsalts of the type Ag₂X possibly exist, where X is a halogen. The principal silver salts of oxyacids are: arsenate and arsenite, borate, bromate, carbonates, chlorate,

chlorite, perchlorate and hypochlorite, chromate and dichromate, iodate and periodates, molybdates, nitrate, nitrite and hyponitrite, phosphates, selenate and selenite, sulphates, sulphite and thionates, tellurates and tellurite. Several salts derived from sulphur-containing acids also exist, the chief of which are: thio-antimonates and antimonites, thio-arsenates and arsenites, thio-carbonate, thio-molybdates, thio-phosphates and phosphites, and thio-tellurite.

Silver, selenide of, Ag₂Se. Occurs native, also in combination with selenides of Cu and Pb. Obtained by heating Ag with Se, SeO₂, or H₂Se; also by passing H₂Se into solution of a Ag salt; and by heating AgNO₄Aq with Se (Senderens, C. R. 104, 175). A grey solid, melting at red heat to silver-white regulus. A little Se is lost by roasting in air; sol. conc. hot HNO₄Aq, crystals of Ag₂SeO₄ separating as liquid

cools.

Silver, selenocyanide of, v. vol. ii. p. 848.
Silver, silicides of. Berzelius found that
Ag after fusion with SiO₂ and C gave a residue
of SiO₂ when dissolved in HNO₂Aq. Percy
(Silver and Gold, 1, 131) failed to obtain any
compound of Ag and Si. Warren (C. N. 60, 5)
obtained compounds or alloys of Ag and Si by
heating Ag with K₂SiF₄ and Na.

Silver, silico-fluoride of, Ag,SiF_e.2H₂O. White, deliquescent crystals, melting below 100°; when strongly heated gives a residue of Ag with some SiO₂. Obtained by evaporating a solution of Ag₂O in H₂SiF_eAq (Marignao, Ann.

M. [5] 15, 270).

Silver, sulphides of. The isolation of any sulphide other than Ag₂S is very doubtful.

SILVER SULPHIDE Ag₂S. (Argentic sulphide.) Mol. w. not known. Occurs native as argentits or silver glance. Prepared by heating Ag in vapour of S, or in H2S, or by the action of H2S on red-hot AgCl (Durocher, C. R. 32, 823). H₂S quite free from admixed O is said not to react with Ag. Also prepared by passing H2S into a solution of a salt of Ag; by boiling Ag, SO, Aq or AgNO, Aq with S (Filhol a. Senderens, C. R. 104, 175); and by the contact of H2SAq with moist Ag.O. Spring formed Ag.S by subjecting a mixture of Ag powder and amorphous S to a pressure of several thousand atmospheres, powdering, and compressing again, and repeating this treatment seven or eight times (Bl. [2] 89, 641; 41, 488, 492; cf. Friedel, Bl. [2] 40, 526). Thomsen (Th. 3, 381) gives [Ag²,S] = 5,340. Winssinger (Bl. [2] 49, 452) obtained a dilute aqueous solution of colloidal Ag. S by ppg. from a very dilute solution of a Ag salt, by H2S, and dialysing; the solution was reddish brown, and did not change after keeping for more than two

Ag₂S obtained by ppn. is a black solid; when melted, out of air, it solidifies to a greyblack, lustrous, malleable mass, that can be cut with a knife; S.G. 6.85 (Karsten, S. 65, 394). S.G. of argentite 7.27 to 7.32 (Dauber, J. 13, 748). Heated in air slowly goes to Ag, giving off SO₂. With conc. boiling HClAq gives AgCl and H₂S; hot HNO₂Aq forms AgNO₂Aq and H₂S; conc. H₂SO₄ produces Ag₃SO₄ and SO₂. Heated in Cl. AgCl and S₂Cl₂ are formed. Reduced to Ag by action of metals and dilute acid; also reduced by heating with PbO, CuQ,

Fe powder, or Hg (cf. Rammelsberg, C. J. 89,

Many compounds of Ag2S with metallic sulphides occur in minerals, e.g. with Cu2S, Sb₂S₃, As₂S₃, Bi₂S₃, &c. Schneider (J. pr. 41, 414) obtained a compound of Ag2S and Bi2S. by fusing the two sulphides together in covered crucible. Poleck and Thümmel (Ar. Ph. [3] 22, 1) obtained Ag₂S.AgNO₃ by passing H₂S into conc. AgNO, Aq so long as it was completely absorbed, washing the greenish pp. with HNO,Aq, and drying; a black amorphous powdecomposed by water to Ag2S and AgNO,Aq.

SILVER SUBSULPHIDE. According to Guntz (C. R. 112, 861), Ag,S is obtained by the action

of H₂S on Ag₂Cl.

Silver, sulphocyanide of, v. vol. ii. p. 352. Silver, telluride of, Ag2Te. Occurs native as hessite. Prepared by melting together Ag and Te, by passing vapour of Te diluted with much N over red-hot Ag (Margottet, C. R. 85, 1142), and by heating AgNO₄Aq with Te in a sealed tube (Senderens, C. R. 104, 175). Grey, lustrous solid; obtained in needles (octahedral) by the second method given above. M. M. P. M.

SINALBIN $C_{90}H_{44}N_2S_2O_{16}$. A glucoside occurring in the seed of white mustard (Will, Z. [2] 7, 89; A. 199, 150). Prepared by washing the seeds with CS2 and extracting the residue with alcohol. Concentric groups of pale-yellow needles (from alcohol), v. sol. water, insol. CS2 and ether. Reduces Fehling's solution, with formation of Cu2S. Split up by the ferment myrosin into glucose, sinapine sulphate $C_{18}H_{23}NO_3H_2SO_4$, and C_7H_7OCNS , a pungent oil. Aqueous AgNO₃ produces a similar decomposition, forming a pp. whence H2S removes Ag, leaving in solution sulphate of sinapine and C₈H,NO [69°]. Coloured yellow by alkalis, and yields Na₂SO₄ and sodium sulphocyanide on boiling with NaOHAq. HgCl₂ gives a white pp.

SINAMINE v. ALLYL CYANAMIDE.

SINAPIC ACID $C_{11}H_{12}O_8$. [192° cor.]. Formed, together with neurine, by boiling sinapine sulphocyanide with alkalis (Von Babo a. Hirschbrünn, A. 84, 19; Remsen a. Coale, Am. 6, 50). Small yellow prisms (from alcohol), sl. sol. cold water, insol. ether. The ammoniscal solution turns brown in air. Salt.—BaA" (dried at 110°). Pp.

Acetyl derivative C11H11AcOs. [281°]. Got by boiling the acid with Ac.O. White

crystals, sol. hot water.

SINAPINE C16H22NOs. Occurs as sulphocyanide in seeds of white mustard (O. Henry a. Garot, J. Ph. 17, 1; 20, 63; Pelouze, A. Ch. [2] 44, 214; J. Ph. 17, 271; Boutron a. Robiquet, J. Ph. 17, 279; O. Henry a. Plisson, A. Ch. [2] 46, 198; Winckler, Rep. Pharm. 41, 169; 67, 257; Simon, P. 43, 651; 44, 593; Boutron a. Fremy, J. Ph. 26, 50; Von Babo a. Hirschbrünn, A. 84, 10; Remsen a. Coale, Am. 6, 50). Formed also by the action of myrosin on sinalbine (Will a. Laubenheimer, A. 199, 162). The free base is known only in solution, and is decomposed on evaporation, leaving a resin. Boiling baryta-water splits it up into neurine C₂H₁₈NO, and sinapic acid. Salts.—B'H₂SO, 2aq: rectangular plates, v.

sol. water and hot alcohol, nearly insol. ether.-B'HHgCl₁: thin prisms.—B'HONS. [176°]. SINAPOLINE v. DI-ALLYL URFA.

SINCALINE v. NEURINE.

SINISTRIN v. INULIN.

SIPERINE. An amorphous, resinous, alkaloid occurring, together with bebirine, in the greenheart tree (Nectandra Rodiæi or Bebeeru Sipeeri). It is v. sl. sol. water, m. sol. alcohol, insol. ether (Maclagan, A. 48, 106).

SKATOLE v. METHYL-INDOLE.

SKELETINS v. Proteïds, Appendix C. SKIMMIN C₁₈H₁₈O_s. [210°]. A glucoside extracted by alcohol from Skimmia japonica (Eijkman, R. T. C. 3, 204). White needles, sl. sol. cold water. Its alkaline solutions show blue fluorescence. It is not poisonous, and does not

reduce Fehling's solution.

Skimmetin C.H.O. [223°]. Formed, together with glucose, by boiling skimmin with dilute mineral acids. Colourless crystals, sol. alcohol and ether. Its solutions exhibit blue fluorescence. Does not reduce Fehling's solution. FeCl, gives a blue colour. Gold chloride gives a rose colour, turning blue and violet.

SMILACIN C18H20O6 (?). Occurs in sarsaparilla root (Smilax Sarsaparilla) (Reinsch, Rep. Pharm. 82, 145; Thubeuf, A. 5, 204; 14, 76; Batka, A. 11, 805; Poggiale, A. 13, 84; Henry, A. 14, 77; Petersen, A. 15, 74; 17, 166; Wright a. Rennie, C. J. 89, 237). Nodular 166; Wright a. Rennie, C. J. 89, 237). groups of leaflets, m. sol. hot water and alcohol, sol. alkalis.

Reference.—Parillin.

SNAKE POISON v. PROTEÏDS.

SOAP v. DICTIONARY OF APPLIED CHEMISTRY. SOBREBOL $C_{10}H_{10}O_2$. [150°]. $[a]_D = \pm 150^\circ$. A product of atmospheric oxidation of turpentine in sunlight (Armstrong a. Pope, C. J. 59, Long tables (from alcohol) or monoclinic prisms (from water); a:b:c=2.411:1:.853; $\beta=83°.38'$. Tastes bitter. Converted by boiling dilute H_2SO_4 into an oily isomeride of cam-By crystallising a solution of equal quantities of dextro- and lævorotatory sobrerol there is obtained an inactive sobrerol [131°] crystallising from alcohol in trimetric tables; a:b:c=2.424:1:.827.

SOCALOÏN v. ALOÏN.

SODA; v. Sodium hydroxide, p. 479. SODIUM Na. At. w. 22.995. Mol. w. very probably same as at. w. (v. infra). Melts at 95.6° (Bunsen, A. 125, 367), 97.6° (Hagen, W. 19, 480); solidifies at 97.6° (Regnault, J. 1856. 48). Boils between 860° and 950°, according to Carnelley and Williams (C. J. 85, 565); at 742° (Perman, C. J. 55, 826). S.G. 985 0 (Schröder, P. 106, 226); 9748 at 10° (Baumhauer, B. 6, 655); c. ·74 at b.p. (Ramsay, C. J. 89, 49). 12.7 at 1200°-1500° (Scott, Pr. E. 14, 410; v. infra). S.H. 2784 from -28° to 6° (Regnault, infra). S.H. '2734 from -28° to 6° (Regnault, A. Ch. [8] 46, 257); '21 when liquid (Joannis, A. Ch. [6] 12, 358). C.E. (0° to 50°) '00007105 (Hagen, W. 19, 436). E.C. (Hg at 0°=1) 18·3 solid at 0°, 8·3 liquid at 120° (Matthiessen, P. M. [4] 12, 199; 13, 81). T.C. (Ag=100)=86·5 (Calvert a. Johnson, P. M. [4] 16, 381). H.G. [Na²,0]=99,760 (Th. [3] 232). Refraction-equid valent $\frac{\mu-1}{d}$ × at. w. = 4.4 (Gladstone, Pr. 18, 49);

474 SODIUM.

4.03 (Kanonnikoff, J. R. 1884 [i.] 119). Emission spectrum characterised by a double line D, the components of which have the wave-lengths 5895 and 5889 (for detailed measurements of lines v. B., A. 1884. 443). For absorption spectrum v. Roscoe a. Schuster (Pr. 22, 362). S.V.S. solid 23.3; at b.p. 31 (Ramsay, C. J. 39, 49)

Occurrence. Compounds of Na occur widely distributed in large quantities. NaCl is found abundantly as rock-salt, also in sea-water and many mineral springs. Large quantities of NaNO, or Chili saltpetre occur in S. America. Carbonate, sulphate, and borate of Na are found, in comparatively small quantities, in rocks, and in lake and spring waters. Cryolite is a double fluoride of Na and Al. Many silicates contain Na silicate-e.g. natrolite, albite, labradorite, &c. The ashes of many plants contain Na salts, especially NaCl, NaI, and Na, SO,; some plant ashes are free from Na salts, even when the plants grow near the sea; the ashes of algae and some other sea plants are richer in salts of K than in those of Na. NaCl, Na₂CO₂, Na₂HPO₄, and some other compounds of Na, are found in animal organisms.

Historical. - Caustic soda was decomposed by Davy in 1807 (T. 1808. 1) by electrolysis; Gay-Lussac a. Thénard heated NaOH with Fe filings and obtained Na (A. Ch. 65, 325 [1808]). Brunner (S. 71, 201) reduced Na₂CO₃ by heating with charcoal; this process was made applicable on the large scale by Deville (A. Ch. [8]

43, 5 [1855]).

The word neter is used by the older Hebrew writers to denote a substance employed for washing clothes; the same, or a similar, substance seems to have been known as virpov or nitrum, and by the fifteenth century this name was given to saltpetre; when the existence of two similar but distinct substances having detergent properties was recognised, the name nitrum was given to one, and the other was called natrum or natron. The metal obtained from natron was called natrium; this name is retained in German, but in English and French the metal is named from soda, a word applied in the Middle Ages to designate alkaline substances, and used as synonymous with nitrum and natron. The symbol Na is universally employed for the metal.

Formation.—1. By electrolysing NaOH (cf. Potassium, Formation, No. 1, p. 298).—2. By reducing Na₂CO₈ by heating it with charcoal, or with charcoal and very finely-divided iron (Castner, C. N. 54, 218).—3. By electrolysing fused NaCl, or conc. NaClAq, using Hg as the

negative electrode.

Preparation.—1. An intimate mixture of 30 parts calcined Na₂CO₂, 13 parts coal, and 5 parts chalk is heated in an iron bottle with a short exit tube, in a furnace, until vapours of Na appear at the mouth of the tube, when a small flat receiver, made of sheet iron, is adapted to the tube; when the receiver is full of Na it is removed, and the metal is taken out under mineral oil (cf. Potassium, Preparation, p. 298). As Na does not combine with CO, the difficulty which attends the preparation of K because of the formation of KCO, does not occur with Na.-2. A mixture of NaOH, C, and Fe is heated in egg-shaped iron retorts, connected with cast-iron condensers

about 3 ft. long and 5 in. diameter, and having small openings near the nozzles through which the molten Na runs into pots placed beneath. The mixture of C and Fe is made by heating a mixture of very finely divided iron and pitch; this is heated with NaOH in approximately the proportion 6NaOH:2C:4Fe. (For details v. Dictionary of Applied Chemistry, vol. iii. p. 422.)

The metal is purified by pressing through linen under rock oil kept at c. 100°; it must be kept under oil free from O. Rosenfeld (B. 24, 1658) recommends to soak the metal in a mixture of 1 part amyl alcohol and 3 parts petroleum. and to rub it with a rag soaked in the same mixture, till the surface is bright; then to let it remain under petroleum with 5 p.c. amyl alcohol; to wash it with pure petroleum, and to keep it under petroleum containing from } to 1 p.c. amyl alcohol. The film of sodium-amyl oxide which forms is easily rubbed off with filter

Properties.—A silver-white, lustrous metal; usually described as whiter than K, but according to Long (C. J. 13, 122) a clean surface has a pale rose colour. By melting Na in a glass tube, narrowed at one place, and filled with coal-gas or H, and allowing the semi-solidified metal to flow slowly through the narrowed part of the tube, Na is obtained in lustrous, white, quadratic octahedra, with a rosy sheen (Long, l.c.; cf. Böttger, B. 7, 1536). According to Linnemann (J. pr. 75, 128), the surface of a freshly-out piece of Na shows green phosphorescence that soon disappears, but returns again at 60°-70° (cf. Baumhauer, J. pr. 102, 123, 361). A sublimate, in a very thin film on glass, shows a greenish blue colour in transmitted light (Dudley, C. N. 66, 163; Newth, N. 47, 55). The vapour obtained by boiling Na in an atmosphere of H is colourless in thin layers, and violet to purple when seen in thick layers (Roscoe a. Schuster, Pr. 22, 362; Carnelley a. Williams, C. J. 35, 565). At the ordinary temperature Na is as soft as wax; it is ductile at 0° , and fairly hard at -20° ; at c. 50° it is pasty; it melts at c. 96° , and boils at c. 750° (v. beginning of this article). Na dissolves in liquid NH, forming a blue liquid (v. Sodium-ammonium, p. 475). A freshly-cut surface of Na exposed to air becomes rapidly covered with a film of oxide; Na is said to be unchanged at the ordinary temperature in perfectly dry air. It decomposes cold water rapidly, but the H evolved does not generally take fire. Na must be kept under a liquid free from O, such as mineral oil or naphtha (v. Preparation, supra).

The atomic weight of Na has been determined (1) by converting NaCl into AgCl (Berzelius, P. 8, 189 [1826]; Pelouze, C. R. 20, 1047 [1845]; Dumas, A. Ch. [3] 55, 182 [1859]; Stas, Stas. Rech. 78); (2) by reducing NaClO, to NaCl (Penny, T. 129 [1] 25 [1839]); (3) by converting NaCl into NaNO, (Stas, Nouv. R. 248 [1865]);

(4) by determining S.H. of Na.

The molecular weight of Na is not known with certainty. Dewar a. Dittmar (C. N. 27, 121) and D. a. Scott (Pr. 29, 206 [1879]) obtained numbers that indicated a mol. w. of 23; but the values varied considerably. D. a. S. used a vessel of wrought iron. Scott (Pr. E. 14, 410 [1888]) ch-

tained fairly concordant results, giving values for V.D. varying from 11.8 to 15, using a Pt vessel heated above the m.p. of cast iron; but V. Meyer's experiments (B. 13, 391 [1880]) seemed to indicate that Na vapour attacks vessels of Ag or Pt. Mitscherlich (A. 12, 173 [1834]) found that vessels of hard glass were also attacked; and Rieth (B. 4, 807 [1871]) got no constant results by using a vessel of Cu. The values obtained by Scott were so constant that there is a large probability in favour of the monatomicity of the molecule of gaseous Na. Ramsay (C. J. 55, 521) dissolved Na in Hg and measured the decrease of vapour-pressure of the Hg thereby produced; assuming the mol. w. of liquid Hg to be 200, and assuming that equal vols. of dilute solutions in Hg contain equal numbers of molecules, the results gave values for mol. w. of Na varying from 15 to 21.5.

The V.D. of no compound of Na has been determined; from the analogy of K, Rb, Cs, and Ag, the atom of Na is probably monovalent in

gaseous molecules.

Sodium is a strongly positive element; it comes after Rb, Cs, and K in order of decreasing electropositiveness (Bunsen). In all its compounds Na reacts as a metal; it does not enter into the composition of any negative radicles. Na forms few basic salts and no oxyhaloid compounds; its haloid compounds are very stable towards heat. Na is closely related to Li, K, Cs, and Rb (v. Alkalis, Metals of the, vol. i. p. 114), and less closely to Cu, Ag, and Au (v. Copper group of elements, vol. ii. p. 250).

Reactions and Combinations.—1. Heated in air or oxygen, Na₂O and Na₂O₂ are formed (v. Oxides, p. 482). According to von Bonsdorff (P. 41, 296), Na is unchanged in quite dry air at the ordinary temperature.—2. Na absorbs hydrogen at c. 300°, probably forming a compound (v. Hydride, p. 479).—3. Combines with chlorine, bromine, and iodine when heated with these elements (v. Chloride, bromide, and iodide). -4. Forms sulphides, by reacting with sulphur at the ordinary temperature; combines, when heated, with selenion and tellurium, also with arsenic and phosphorus (v. Sulphide, p. 483, Selenide, p. 482, Telluride, p. 483, Arsenide, p. 476, and *Phosphide*, p. 482).—5. Alloys with several metals (v. Alloys, infra).-6. Na decomposes water at the ordinary temperature, forming NaOHAq and H; much heat is produced, but the H does not generally take fire unless the reaction is slackened, and friction is increased, by thickening the water by gum &c., or placing the Na on moist filter paper (v. Ducatel, Am. S. 25, 90; Serullas, A. Ch. [3] 40, 329). If Na is allowed to react with water in a cylinder standing over water, the H collecting in the cylinder, a violent explosion sometimes occurs, probably due to formation of a peroxide that is then decomposed with evolution of O, which mixes with the H (Böttger, J. pr. 85, 397; Griffin, J. 1856. 320).—7. Na reduces many oxides at high temperatures, e.g. CO, N₂O, B₂O₄, P₂O₅, As₂O₂, HgO, CuO, Ag₂O, PbO, ZnO, Fe₂O₃, &c.—8. Many carbonates, sulphates, nitrates, phosphates, and silicates are reduced when strongly heated with Na.—9. Many metallic chlorides are reduced by heating with Na, e.g. MgOl₂, AlCl₂ &c.—10. By heating with solid caustic potash to c. 170°, an alloy of Na and K is formed (v. Williams, C. N. 3, 21; Wanklyn, C. N. 3, 66).—11. Heated in ammonia gas, NaNH₂ is formed (v. Amide, infra), and when this is more strongly heated Na₃N is formed (v. Nitride, p. 482).—12. Na reacts with aqueous solutions of acids to form salts NaX, where X is a monovalent acidic radicle.

Sodium, alloys of. Na forms alloys with many metals; these alloys are generally formed by directly melting the metals together, or by reducing the chlorides by Na. By the action of sodammonium (v. infra) on Sb, Bi, Pb, Joannis (C. R. 114, 585) says that alloys of Na with these metals are obtained agreeing in composition with the formulæ Na₃Sb, Na₃Bi, and NaPb. Schumann (W. 43, 101) obtained a definite alloy of Na and Hg corresponding with the formula NaHg₅ (v. also SODAMMONIUM, infra).

Sodium, aluminates of, v. vol. i. p. 141.

Sodium, amide of, NaNH₂. (Sodamide.) Mol. w. not determined. First prepared by Gay-Lussac and Thénard (Recherches physicochimique, 1, 354) by heating Na in NH₃; the Na absorbed 142 to 163 vols. NH₃ and 100 vols. H were produced. Prepared by Beilstein and Geuther (A. 108, 88) by driving out the air from several flasks, connected by tubes, by a stream of dry H, then placing a few grams Na in each flask, replacing the H by dry NH₃, free from CO₂ (Drechsel, J. pr. [2] 16, 203), and heating the flasks on sand-trays; the Na melts and swims on the blue-green liquid that forms. When the Na has all disappeared the liquid is allowed to cool; the solid that forms is at first brown, but when quite cold it forms a crystalline, olive-green mass. Also obtained by the gradual decomposition at the ordinary temperature (hastened by light) of sodammonium; as the liquid decomposes, crystals of NaNH₂, 1 mm. long, are formed; these crystals are said to be white (Joannis, C. R. 112, 392).

Heating in CO produces NaCN and H₂O, along with NaOH and NH₂. When heated in CO₂ much NH₃ is given off, and the final products are H₂O and CN.NNa₂ (disodium salt of cyanamide; v. vol. ii. pp. 313, 314); B. a. G. (l.c.); Drechsel (l.c.). CS₂ reacts to form NaSCN and H₂S, then NH₃ is given off, and NaSH remains. Dilute HCLAq reacts violently, producing NaCl and NH₄Cl. EtCl forms NaCl, NH₃, and C₂H₄, but no EtNH₂. Heated with C the products are NaCN and H (Drechsel, J. pr. [2] 21, 91). When strongly heated NaNH₂ gives off NH₃ and leaves NaN₃ (v. SODIUM MITRIDE,

o. 482).

Sodium-ammonium or Sodammonium. Weyl (P. 121, 611; 123, 350) obtained a blue liquid, by the action of liquid NH₃ on Na, which he regarded as a compound or alloy of Na and NH₄. As Na remained when the NH₃ was allowed to evaporate, Seeley (C. N. 23, 169) looked on the blue liquid as a solution of Na in liquid NH₃. Joannis (C. R. 109, 900, 965; 110, 238) found that the vapour-pressure of a solution of Na in liquid NH₄ decreased to a limit whereat it became constant if temperature remained unchanged; that on then removing more NH₃ a copper-red solid separated, and that NH₄ was then given off at constant pressure till only Na remained. When the last part of the change commenced, the copper-red substance contained

Na and NH, in the ratio Na:NH,; the evolution of NH, at constant pressure was regarded by J. as due to the dissociation of a compound NaNH,. The heat of formation of NH, Na from gaseous NH, and solid Na was determined by J. to be 5,200, and from liquid NH, and solid Na to be 800, cals. J. determined the lowering in the freezing-point of liquid NH, by solution in it of NaNH,, and so calculated the mol. formula of this body to be Na, N₂H₆ (C. R. 115, 820). The solution of Na₂N₂H₆ in liquid NH₂ is decomposed by Sb, Pb, and Hg, but not by Al, Cu, Ag, or Zn; the final product of the action of Pb is Pb, Na. 2NH,; by dropping the liquid on to Hg, and washing the product with liquid NH, the crystalline amalgam NaHg, was obtained (J., C. R. 113, 795). According to J. (C. R. 112, 392), Na₂N₂H₆ decomposes slowly at the ordinary temperature to NaNH, and H. With excess of NaCl the blue liquid becomes colourless and NaNH2 separates; the compound NH, Na, Cl is first formed, and is then decomposed by the liquid NH, to NaNH, and NaCl. By passing dry O into Na,N,H, in liquid NH, at -50°, J. (C. R. 116, 1370) obtained NH,Na,OH; by the prolonged action of O he obtained Na₂O₂.
Sodium, arsenates of, v. vol. i. p. 809.
Gay-Lussa

Gay-Lussac and Thénard found that Na combined directly with As. According to Landolt (A. 89, 210) NaAs is formed, as a silver-white solid, by heating the elements in the ratio 3Na:As. A compound of Na and As is also formed when Na is heated in AsH₂. Decomposed by water, giving off AsH₂.

Sodium, arsenites of, v. vol. i. p. 307. Sodium, aurobromate of, NaAuBr₄. Orange-red prisms, by dissolving NaBr in AuBr₄Aq (cf. vol. ii. p. 649).

Sodium, aurochlorate of, NaAuCl4. By dissolving NaCl in AuCl, Aq (cf. vol. ii. p. 650).

Sodium, borates of, v. vol. i. p. 529. Sodium, boride of. The product of the reduction of B₂O₂ by Na may contain a compound of B and Na.

Sodium borofluoride v. vol. i. p. 526. Sodium, bromide of, NaBr. Mol. Mol. w. not certainly known, but probably 102.745 (= NaBr). Melts at 708° (Carnelley, C. J. 83, 279); at 727° (v. Meyer a. Riddle, B. 26, 2443). S.G. 3.079 at 17.5° (Kremers, P. 99, 443); 3·198 at 17·3° (Favre a. Valson, C. R. 77, 579); 2·448 when molten (Quincke, P. 138, 141). Kremers (P. 99, 443) gives solubilities as follows:-

Temp.	s.	Temp.	8.
0°	77.5	60°	111-1
20	88.4	80	112.5
40	104.2	100	114.9
	C1. F43 OF	. FOR SET OF	4

Coppet (A. Ch. [4] 25, 506; [5] 80, 411) gives

the following:-

Temp.	S.	Temp.	8.
44·1°	115.6	86°	118.8
51.5	116.2	90.5	119.7
55.1	116.8	100.3	120.6
60.8	117	110-6	122.7
64.5	117.8	114.8	124
74.5	118-4		

C. represents S. of NaBr as 110.84 + 1075t when
 t varies from 44° to 114°.
 S. in alcohol 90 p.c.

of NaBrAq given by Kremers a. Gerlach (Fr. 6, 279) at 15° as follows :—

S.G. NaBrAq	P.O. NaBr	S.G. NaBrAq	P.C. NaBr
1.04	5	1.281	80
1.08	10	1.344	85
1.125	15	1.410	40
1.174	20	1.483	45
1.226	25	1.565	50

H.F. [Na,Br] = 85,770; [Na,Br,Aq] = 85,580 (Th. 8, 232).

Formation.—1. By passing Br vapour over Na. According to Merz and Weith (B. 6, 1518), Na remains unchanged for months in liquid Br at the ordinary temperature, and even at 100° or 200° the formation of NaBr proceeds very slowly.—2. By treating BrAq with iron filings, boiling the solution of FeBr₂Aq with NaOHAq, filtering, and evaporating (Henry, J. Ph. 15, 54).—3. By neutralising HBrAq by NaOHAq or Na₂CO₂Aq, and evaporating.—4. By decomposing CaBr₂Aq by Na₂SO₄, filtering after some time, adding Na CO, as long as turbidity is produced, filtering, and evaporating (Klein, A. 128, 237).-5. By decomposing pure NH, Br by NaOHAq or Na₂CO₄Aq, and evaporating (Castelhaz, C. R. 70, 1050).

Preparation.—Br is added, little by little, to moderately conc. pure NaOHAq (v. Sodium hydroxide, p. 479) till the liquid is slightly yellow; the liquid is evaporated to dryness; the residue of NaBrO, and NaBr is strongly heated, in a porcelain dish, till traces of Br are given off, the residue is dissolved in water, evaporated, and allowed to crystallise at a temperature not under 30° (NaBr. 2aq separates below 30°).

Properties.—Lustrous, white cubes; strong alkaline taste; reaction in water is neutral. Dissolves easily in water, with slight fall of temperature [NaBr,Aq] = -190 (Th. 3, 232); NaBr separates from this solution at c. 30° but at the ordinary temperature crystals of NaBr. 2aq are formed.

Reactions .- 1. Chlorine reacts with NaBrAq to give NaClAq and Br.-2. Heated with potassium chromate and sulphuric acid, Br is given off: NaCl under similar conditions gives CrO₂Cl₂.—8. Conc. sulphuric acid forms HBr. Br, and SO,; by using dilute H.SO.Aq (c. 30 p.c.) a mere trace of Br is obtained (v. Addyman. C. J. 61, 94).—4. Potassium permanganate does not react with NaBrAq even when boiled, but Br (Hempel, A. 107, 160).

Br (Hempel, A. 107, 160).

With water to form addition of a little H2SO4Aq causes evolution of

Combinations.—1. With water to form NaBr.2H₂O. This hydrate separates from solutions of NaBr saturated when hot and cooled to under 30°; it crystallises in monoclinic prisms, which melt at 50°, leaving NaBr.-2. With many bromides of less positive metals, e.g. with PbBr, to form PbBr, 2NaBr, and CdBr, to form CdBr, 5NaBr. 5aq (v. the various metallic bromides).—3. With arsenious oxide to form

NaBr.As,O, (Rüdorff, B. 21, 3051).

Sodium, chloride of, NaCl. (Common salt.) Mol. weight not known with certainty, but pro-74.5 118.4 bably 58.365 (= NaCl). Melts at 772° (Carnelley, C. represents S. of NaBr as 110.84 + .1075t when twaries from 44° to 114°. S. in alcohol 90 p.c. a. B.P. of saturated NaBrAq=121°. S.G. 188, 141); 2.157 at 4° (Schröder, P. 106, 226); (?) 1-612 at m.p. (Braun, B. 7, 958); 2·04 after fusion (Quincke, P. 185, 642); for other values v. Clarke's Table of Specific Gravities [2nd ed.] 20. S.H. 15° to 98° ·21401 (Regnault, A. Ch. [8] 1, 129; v. also Kopp, T. 155 [i.] 71). Vol. at 40° = vol. at 0° (1 + ·00012117 × 40) (Fizeau, C. R. 64, 314). H.F. [Na,Cl] = 97,690 (Th. 3, 232). Poggiale (A. Ch. [8] 8, 469) gives following data for solubility in water:—

Temp.	s.	Temp.	s.
-15°.	82.73	40°	36.64
-10	33.49	50	36.98
- 5	34.22	60	37.25
0	35.52	70	37.88
+ 5	85.63	80	38.22
9	35.74	90	38.87
14	35.87	100	39.61
25	86.13	109.7	40.35

Coppet (A. Ch. [5] 30, 411) gives S. above 20° to $109^{\circ} = 34 \cdot 359 + 0527t$. Gerlach (Fr. 8, 281) gives the following data:—

		S.G. NaO	lAq at 15°	
1	p.c.	1.00725	10 p.c.	1.14315
2	- ,,	1.01450	11 "	1.15107
3	"	1.02174	12 ,,	1.15931
4	11	1.02899	13 "	1.16755
5	,,	1.03624	14 ,,	1.17580
6	**	1.04366	15 ,,	1.18404
7	11	1.05108	16 ,,	1.19228
8	11	1.05851	17 ,,	1.20098
9	"	1.06593	18 "	1.20433

1000 c.c. of NaClAq saturated at 15° contain 318·479 g. NaCl and 888·669 g. water (Michel a. Kraft, J. 1854. 296). NaClAq containing 29·4 to 29·5 p.c. NaCl boils at 109·25° (at 760 mm. pressure) and freezes at -21·3° (Karsten, Salinenkunde [Berlin, 1847] 2,38). The following freezing-points are given by Karsten (l.c.) for NaClAq (cf. Rüdorff, P. 114, 63; and de Coppet, A. Ch. [4] 25, 509):—

P.C. NaCl	F.P.	P.C. NaCl	F.P.
2	-1·32°	16	-11.69
4	-3.024	20	-14.44
6	-4.52	22	-15.78
8	- 5.99	24	-17.11
10	-7.44	26	-18.42
12	-8.88		

For freezing-points of dilute NaClAq, containing from c. 006 to c. 2.6 p.c. NaCl, v. Jones (Z. P. C. 11, 210).

Solution of NaCl in water is accompanied by lowering of temperature and contraction; 36 parts NaCl in dissolving in 100 parts water at 12.6° lower the temperature to 10.1° (Rüdorff, B. 2, 68); Thomsen gives [NaCl,Aq] = -1180 (Th. 3, 232). By mixing 36 parts NaCl with 100 parts snow, temperature falls to -21.3° (Rüdorff, P. 114, 79; 122, 337).

NaCl is insol. absolute alcohol; Girardin (A. Ch. [4] 5, 146) gives the following data for S. in aqueous alcohol S.G. 9282; 10.9 at 4°, 11.1 at 10°, 11.43 at 13°, 11.9 at 23°, 12.3 at 32°, 13.1 at 44°, 13.8 at 51°, 14.1 at 60°. Schiff (A. 118, 365) gives S. in alcohol at 15° as follows; 28.38 in 10 p.c. alcohol, 13.25 in 40 p.c., 5.93 in 60 p.c., 1.22 in 80 p.c. NaCl crystallises in the regular system, generally in cubes, sometimes in octahedra.

Occurrence.—As rock salt, or habite, in very under Combinations, p. 478).

large quantities very widely distributed; in seawater, many salt lakes, brine springs, and most mineral springs.

Formation.—1. By burning Na in moist Ch. According to Wanklyn (C. N. 20, 271), dry Ch. does not combine with Na even at the M.P. of the metal; Donny a. Mareska (C. R. 20, 817) state that Na is unacted on by liquid Cl at -80°.—2. By neutralising NaOHAq or Na₂CO₄Aq by HClAq, and evaporating.—3. By the reaction of excess of conc. HClAq with several salts of Na, e.g. Na₂SO₄.—4. By strongly heating Na₂SO₄ with excess of NH₂Cl; also by decomposing Na₂SO₄Aq by several chlorides, e.g. MgCl₂ or CaCl₂.—5. As a by-product in making KNO₃ by the reaction of KCl with NaNO₃. 6. By the reduction of many chlorides by Na.

Preparation.—Pure NaHCO, is prepared by washing a large quantity of the ordinary 'pure salt, in fine powder, with cold water till the washings cease to be clouded on addition of a salt of Ag or Ba; after boiling with slight excess of HNO, Aq, the residue is dried and heated to dull redness in a Pt dish; a boiling saturated solution of the Na₂CO, thus obtained is prepared, filtered, and shaken while crystallising to insure the formation of small crystals; the crystals are dried by suction by a water-pump, they are then twice moistened with cold water to remove all mother-liquor, and crystallised three times from water. The pure Na₂CO₃ is dissolved, and NaCl is formed by passing in a stream of pure HCl gas (obtained by heating pure conc. HClAq); a little pure NH₂Cl is added to the solution (for preparation of pure NH Cl v. vol. i. p. 197, under Ammonia, Preparation, No. 3), which is then evaporated to dryness in a Pt retort; the residue is very strongly heated, and then dissolved in water; after standing for 24 hours the solution is poured off (from any Al₂O₈.xH₂O and SiO_2 that settle), a little pure \hat{NH}_4Cl is added, the solution is evaporated to dryness in a Pt retort, the residue is strongly heated for some time, and after partial cooling the still liquid NaCl is poured off (from Pt, SiO2, and Al₂O₃) into a Pt dish; when cold, the NaCl is dissolved in water, the solution is poured off after 2 days and evaporated to dryness in a Pt retort, and the residue is fused in a Pt vessel (Stas, Chem. Proport. 275)

For preparation of NaCl from rock salt, brine, and sea-water, v. Dictionary of Applied Chemistry, vol. iii. p. 429.

Properties .- A pure, white, lustrous solid; crystallises in the regular system, generally in cubes. Decrepitates when heated. NaCl is diathermanous. Solution in water is neutral to indicators, and has a bitter taste. Pure NaCl is very slightly hygroscopic, taking up c. \(\frac{1}{3} \) p.c. water from moist air (Stas, Karsten). Melts at c. 772°; vapourises markedly at white heat in a stream of N (v. Stas, l.c.). NaCl is almost wholly ppd. from an aqueous solution by leading in HCl (Margueritte, C. R. 43, 50); also ppd. from a conc. solution by cold conc. NaOHAq (Berthelot C. R. 76, 111); also by warming a conc. solu tion with excess of NaBr or NaI and cooling (von Hauer, J. pr. 98, 137). Saturated NaClAq deposits the hydrate NaCl. 2aq at -7° to c. -22° and at c. - 23° NaOl. 10aq separates (v. Hydrates

Reactions.--1. According to de Sanderval (C. R. 116, 641), when NaCl is vapourised around a porous tube Cl collects in the inside of the tube.—2. Electrolysis of NaClAq produces Cl and NaCHAq, but NaClOAq and NaClO, are soon formed by secondary reactions; if a dia-phragm of asbestos is used, the cathode being on one side and the anode on the other, and CO2 is passed into the NaClAq, Cl is given off and Na₂CO₃ is formed (v. Hempel, B. 22, 2475).-3. Fusion with potassium produces KCl and Na. 4. Fusion with sulphur is said to form Na₂S and S2Cl2, but this is denied by Karsten; NaCl is not decomposed by hydrogen, or oxygen, at a red heat.—5. Superheated steam has no reaction, according to Kunheim (J. 1861. 149); there is a slight reaction, according to Lunge (Soda-industrie, 2, 288).-6. Sulphuric anhydride vapour is absorbed, without heating, giving S2O6NaCl (Na salt of SO₂(Cl).O.SO₂.OH); according to Schultz-Sellack (B. 4, 112) more SO₂ is then absorbed, and a compound approximately NaCl.4SO₂ is formed; on heating Na₂SO₄, SO₂ and Cl are produced. By heating with *liquid* sulphuric anhydride S2O5Cl2 is formed (Rosenstiehl, C. R. 53, 658).—7. Decomposed by heating to c. 500° in a mixture of air, sulphur diocide, and water vapour, giving Na₂SO₄ and HCl (Hargreaves and Robinson's process for making Na₂SO₄; v. Dictionary of Applied Chemistry, vol. iii. p. 444).—8. NaCl is decomposed, with formation of oxychlorides or chlorides, by heating with phosphoric, chromic, or molybdic anhydride, or with antimonic oxide; decomposition occurs with silica, boric oxide, alumina, and tungstic oxide only in presence of air or water vapour, forming silicate, borate, aluminate, or tungstate of Na, and HCl; in presence of water vapour Cl is evolved in some of these reactions (v. Schulze, J. pr. [2] 21, 407; Delalande a. Prudhomme, Bl. 20, 74; Gorgeu, A. Ch. [5] 10, 145). Lead oxide decomposes NaClAq, forming PbCl2 and NaOHAq (Scheele); but molten NaCl is said not to interact with PbO. Moist NaCl mixed with caustic lime and exposed to air becomes covered with an efflorescence of Na₂CO₂ crystals. Magnesia interacts with NaClAq, in presence of CO₂, to form NaHCO₂ and MgCl₂ (Weldon; Wagner, J. 1873. 256).—9. Sulphuretted hydrogen partially decomposes hot NaCl to Na2S, more completely in presence of steam (Kingzett, C. J. [2] 11, 456).—10. Many metallic sulphides, e.g. Ou2S, produce Na₂SO, and metallic chlorides (sometimes Cl is set free) when roasted in air with NaCl.—11. Several metallic sulphates when fused with NaCl produce Na, SO, and metallic chlorides—e.g. PbSO, ZnSO, CuSO,; some of these sulphates react also with NaClAq. Some sulphates form Na, SO₄, and give off Cl when heated with NaCl, e.g. FeSO₄ (v. Barreswil, J. Ph. [3] 12, 456).—12. Fusion with potassium chlorate only decomposes a trace of NaCl (Schulze, J. pr. [2] 21, 407).-13. Ammonium oxalate heated with NaCl forms Na, CO, and distinct quantities of NaCN (Fresenius); addition of $(NH_4)_2C_2O_4$ to hot saturated NaClAq ppts. Na₂C₂O₄.—14. Potassium carbonate reacts with NaClAq to form Na, CO, Aq and KClAq; if CO, is passed in NaHCO, ppts.—15. Anmonium bicarbonats and NaClAq produce NaHCO, and

NH₄ClAq. Passage of CO₂ into NaClAq saturated with NH₃ ppts. NaHCO₃ (v. Ammonia-soda process, in Dictionary of Applied Chemistry, vol. iii. p. 494).—16. Sulphuric acid decomposes NaCl, forming HCl, and NaHSO₄ or Na₂SO₄ according to the temperature. The Leblanc cording to the temperature. process for making sodium carbonate takes advantage of this reaction (v. DICTIONARY OF APPLIED CHEMISTRY, vol. iii. p. 459).-17. Evaporation with excess of nitric acid produces NaNO, and HCl.-18. NaCl is readily decomposed by hydrofluoric acid, also by hydrofluo-silicic acid.—19. Carbonic acid is said to set free some HCl from NaClAq (v. H. Müller, B. 3, 40; Funk, Chem. Zeitung, 1879. 660).—20. Repeated evaporation with excess of oxalic acid completely decomposes NaCl to Na₂C₂O₄.

Combinations. - 1. With water to hydrates. The dihydrate NaCl. 2aq separates from saturated NaClAq cooled to -7° to -22° , in large clear monoclinic crystals (v. Hankel, P. 53, 623; Mitscherlich, J. pr. 83, 485). The crystals effloresce at -10°, and deliquesce at 0°. Ehrenberg (P. 36, 240) and Frankenheim (P. 37, 638) obtained this hydrate by evaporating dilute NaClAq at a moderate temperature. The crystals that separate from hot NaClAq containing HCl are hydrated, according to Bevan (C. N. 35, 17), and contain 5.48 p.c. water, but after a time they change to the ordinary crystals of NaCl. According to Naumann (Thermochemie, 106), crystals of the decahydrate, NaCl. 10aq, separate from NaClAq at -23°.-2. NaCl absorbs considerable quantities of ammonia, under pressure. Joannis dissolved NaCl in liquid N_{H_3} at -10° ; on cooling to -30° and letting NH, escape at the ordinary pressure, he obtained slender white needles, probably NaCl.5NH, (C. R. 112, 337).—3. Sulphuric anhydride is absorbed by NaCl, forming NaCl.2SO,, and then perhaps NaCl.4SO, (v. Reactions, No. 6).—4. Compounds with sodium iodate are formed, by adding excess of NaCl to NaIO, Aq, and by boiling NaIO, with HClAq; the compound NaCl.NaIO_s.4aq was obtained by Ditte, and 3NaCl.2NaIO_s.18aq by Rammelsberg (P. 44, 548; 115, 584).—5. With many chlorides of less positive metals, e.g. AlCl3, CdCl2, CrCl3, PbCl2, MgCl2 (v. Chlorides of various metals).

SODIUM SUBCHLORIDE. H. Rose (P. 120, 15) noticed that a grey-blue solid was produced by fusing NaCl with Na in H; he thought this might be a subchloride of Na. Bunsen and Kirchhoff (P. 113, 339) noticed the formation of a deep-blue substance during the electrolysis of molten RbCl; this substance dissolved in water, giving off H and producing a colourless, strongly alkaline solution; a similar reaction was obtained on electrolysing KCl; these blue substances were regarded by B. a. K. as probably subchlorides, but no analyses or proofs of compo-

sition were given.

Sodium, chromisulphocyanide of, v. vol. ii,

Sodium, cyanide of, v. vol. ii. p. 347. Sodium, ferrate of, v. vol. ii. p. 547. Sodium, ferricyanide of, v. vol. ii. p. 340. Sodium, ferrite of, v. vol. ii. p. 547. Sodium, ferrocyanide of, v. vol. ii. p. 837. Sodium, fluoboride of, v. vol. i. p. 526.

Sodium, fluoride of, NaF. Mol. w. not | with a sharp, acid taste. Sl. sol. cold water known with certainty, but the formula NaF is

probably molecular.

Formation.—1. By neutralising HFAq by NaOHAq and evaporating.-2. Formed along with NaHSO, by the reaction of HFAq with Na₂SO₄ (Weldon, D. P. J. 182, 228).—3. By fusing together CaF, Na, SO, and charcoal, and extracting with water (Jean, C. R. 66, 801, 918).—4. By boiling powdered cryolite with NaOHAq of at least 1.85 S.G. till a little taken out is entirely sol. boiling water, decanting the alkaline liquid from NaF, pressing the NaF, dissolving in boiling water, separating any Al₂O₃ in solution by Na₂SiO₃Aq, removing silica by CO, filtering, evaporating, and recrystallising the NaF that separates (Schuch, A. 126, 108).-5. Berzelius (Lehrbuch [5th ed.] 3, 216) prepared NaF by adding to 100 parts Na₂SiF, and 112 parts Na₂CO₂ water enough to make the whole pasty, boiling as long as CO, was given off, extracting the solid thus formed with boiling water, evaporating the solution to dryness, heating the residue gently, again extracting with hot water, filtering from SiO2, and crystal-

Preparation.—Pure Na₂CO₂ (for preparation v. Sodium chloride, Preparation, p. 477) is dissolved in pure HFAq, till the acid is just saturated, in a Pt dish, the solution is evaporated to dryness, the residue is heated strongly for some time,

and allowed to cool.

Properties.-Clear, lustrous cubes; presence of Na₂CO₂ crystallises in octahedra. Sl. sol. water; S. at $16^{\circ} = 4.78$ (Berzelius), at 15°=4 (Fremy, A. Ch. [3] 47, 32). Scarcely more sol. hot than cold water. Insol. alcohol. Güntz (A. Ch. [6] 3, 5) gives [NaF,Aq] = -600. Decrepitates on heating, and melts at a high temperature without decomposition (Berze-

Reactions. - 1. Partially decomposed by heating to redness in water vapour, with formation of NaOH and HF (Weldon). -2. Excess of caustic potash forms NaOHAq.—8. Boiling with magnesia partially decomposes NaF to NaOHAq and forms a double Na-Mg fluoride (MgF₂.2NaF) (Tissier, C. R. 56, 848).-4. Completely decomposed to NaCl by heating strongly in hydrochloric acid gas (Deville, C. R. 43, 970).-5. Partially decomposed to NaCl by heating with ammonium chloride (Rose, P. 74, 579). Fusion with magnesium chloride produces NaCl and MgF, (Geuther, J. Z. 2, 208).

Combinations .- 1. With several fluorides of less positive metals; 3NaF.AlF, occurs native as cryolite (v. Aluminum fluoride, vol. i. p. 145).—2. With silicon fluoride to form Na,SiF, (v. Sodium silicofluoride, p. 483).—3. With sodium borate to form 6NaF.Na,B₂O₄ (v. Fluoride). borates, vol. i. p. 530).—4. With sodium phosphate to form NaF.2Na,PO₄. 19aq and 22aq, and NaF.Na,PO₄. 24aq (v. Baumgarten, J. 1865. 219; Brieglieb, A. 117, 95).—5. With sodium sulphate to form NaF.Na,SO₄ (v. Marignac, Ann. M. [5]

12, 18). SODIUM-HYDROGEN FLUORIDE NaHF, NaF.HF. This compound is formed, according to Berzelius (Lehrbuch [5th ed.] 3, 217), by the spontaneous evaporation of a solution of NaF in HFAq. Forms small rhombohedral crystals, more sol. hot water. Decomposed by heat to NaF and HF (Marignac, J. 1857. 128). Güntz (A. Ch. [6] 3, 5) gives [NaF, HF] = 17,100 (to form NaF.HF).

Sodium, hydride of, Na.H. Gay-Lussac and Thénard (A. Ch. 74, 203) observed that Na absorbed H when heated therein. The process was more accurately studied by Troost and Hautefeuille (A. Ch. [5] 2, 273). They found that absorption of H by Na began at c. 300° and ceased at c. 421°, when the gas was at the atmospheric pressure. The product gave, in a Sprengel pump, 237 vols. H. for 1 vol. Na; the formula Na.H requires 238 vols. H. Na.H is a soft, silver-white solid; it can be melted without decomposition in H; S.G. 959. Traces of H are given off at 760 mm. pressure; heated in vacuo a regular dissociation takes place from 330° to 430° (for vapour-pressures of H given off v. Dissociation, vol. ii. p. 398). Moutier (C. R. 79, 1242) gives the thermal value [Na²,H] = 13,000 at c. 330°.

Sodium, hydrosulphide of, NaSH. (Sodium sulphydrate.) Sabatier (A. Ch. [5] 22, 5) obtained this compound by treating Na₂S. 9aq with H2S in absence of air, and then evaporating the liquid so formed in an atmosphere of H2S. A white, very hygroscopic solid. Passage of CO2 into NaSHAq drives out H₂S and converts all the Na into Na₂CO₂. NaSHAq contains H₂S and NaOH, according to Gernez (C. R. 64, 86). NaSHAq dissolves S with evolution of H₂S; it ppts. PbS, or MnS, from a neutral solution of a Pb or Mn salt, giving off H₂S; Na₂SAq does not give off H2S while dissolving S or ppg. PbS or

Sabatier (l.c.) obtained the hydrateNaSH.2H2O by passing H2S over Na2S. 9aq, and then evaporating in H2S till half the water was removed; very hygroscopic needles, becoming yellow in air.

Sodium, hydroxide of, NaOH. (Caustic soda. Sodium, or sodic, hydrate.) Mol. w. not determined. Melts at 1098° (v. Meyer a. Riddle, B. 26, 2443). S.G. 2·13 (Filhol, A. Ch. [3] 21, 415); 1·723 (Smith, Am. J. Pharm. 53, 145). S. (cold water) c. 212 (Bineau, C. R. 41, 509). H.F. [Na,O,H] = 101,870; [Na,O,H,Aq] = 111,810; $[Na^2O,H^2O] = 85,620$ (Th. 3, 232).

Formation.—1. By the interaction of Na, or Na₂O, with H₂O.—2. By decomposing Na₂CO₂Aq with CaO₂H₂ (v. Preparation, No. 1).—3. By the action of CaO₂H₂ with NaFAq (Tissier, J. pr. 90, 50).—4. By decomposing Na₂SO₄Aq by BaO or CaO under pressure.—5. From NaClAq by interaction with PbO (Knab, B. 11, 1458).—6. By heating NaNO, with charcoal, or with Cu (Wöhler, A. 87, 873), or Fe (Polacci, C. N. 26, 288).-7. By decomposing Na₂SAq by CuO, Fe₂O₃, PbO, ZnO, &c. (Kopp, D. P. J. 142, 841; Stromeyer, A. 107, 833).— 8. By the interaction of NaF and steam (Weldon).-9. By heating soda felspar with lime.

Preparation .- 1. By causticising soda crystals by lime. A solution of pure Na CO, crystals in 4-5 parts water is boiled, in a dish of silver or polished iron, with milk of lime till a little of the filtered liquid gives off no CO, when acidified; the weight of CaO required is c. } the wt. of soda crystals used, it should be

suspended in about as much water as the wt. of soda crystals used; as the boiling proceeds a little water should be added from time to time, otherwise the NaOHAq will decompose some of the CaCO, formed to CaO and Na₂CO₂Aq. The CaCO, formed is allowed to settle, the vessel being closed, the NaOHAq is drawn off by a siphon and evaporated to dryness in a silver dish, and the residue is heated till volatilisation begins, and allowed to cool in an exsiccator. By dissolving in absolute alcohol, allowing to settle, drawing off the liquid, evaporating it to dryness on a water-bath in a silver dish, removing resinous matter from the warm semisolid mass by a silver spatula, heating to bright redness, and pouring the molten mass on to a plate of polished iron-or, better, of silver-NaOH containing only traces of impurities, chiefly NaCl and Na₂CO₂, is obtained.—2. A large silver dish is surrounded by cold water; 2 or 8 drops of water are placed in the dish, and then a small piece of clean Na, 1 or 2 cms. long; the dish is constantly shaken, so that the Na flows over a large cold surface (explosions are thus avoided); when the reaction is finished 2 or 3 more drops of water are added, and then another piece of Na, while the dish is shaken, and so on. The semi-solid mass is heated till the water is all driven off and the NaOH melts, and the molten substance is poured on to a plate of polished iron or silver.

To obtain NaOH free from nitrites and nitrates Ilosva (Bl. [3] 2, 357) recommends to place Na in water with a layer of petroleum on the surface, the petroleum having been washed with water till free from nitrites and nitrates. He also says that if 2-4 p.c. ordinary NaOHAq is kept for some weeks in contact with granulated Zn all nitrites and nitrates are destroyed.

For preparation of ordinary caustic soda v. Dictionary of Applied Chemistry, vol. iii.

p. 480.

Properties.—NaOH is a white, hard, brittle solid. Melts below red heat; volatilises at very bright red heat, higher than temperature of volatilisation of KOH. Very deliquescent; dissolves in c. '47 parts water. During solution in water much heat is produced, [NaOH,Aq] = 9,940 (Th. 3, 232); the maximum amount of heat is produced when H₂O is added to NaOH in the ratio NaOH: 20H₂O, addition of water to this solution is attended with a slight disappearance of heat (Th. 3, 84). (For hydrates v. p. 480, Combinations.) Easily sol. alcohol. NaOHAq with 36:86 p.c. NaOH boils at 130°, with 70 p.c. boils 180°, with 77:5 p.c. boils 288°, with 78:7 p.c. boils 243°, and with 82:6 p.c. boils 260° (Lunge, Soda-industrie, 2, 548).

NaOHAq acts as a strong alkali; its affinity is c. equal to that of KOHAq and LiOHAq and c. 50 times greater than NH₂Aq (Ostwald). The properties of NaOHAq closely resemble those of potassium hydroxide solution (q. v. p. 302).

The table in the next column, given by Lunge, shows the composition of NaOHAq of different S.G. at 15°.

Regarding the freezing-points of NaOHAq of different concentrations v. Rüdorff (P. 116, 55); de Coppet (A. Ch. [4] 24, 551); and Pickering (who gives very complete data) C. J. 68, 890).

8.G.	Baumé	Twaddell	100 pts. by wt.		1 cub. metre contains kilos.	
PAHORN	Be	F	Na _s O	NaOH	Na _s O	NaOH
1.007	1	1.4	0.47	0.61	4	6
1.014	2	2.8	0.93	1.20	9	12
1.022	8	4.4	1.55	2.00	16	21
1.029	4	5:8	2.10	2.71	22	28
1.036	5	7.2	2.60	8.35	27	35
1.045	6	9.0	8.10	4.00	32	42
1.052	7	10.4	8.60	4.64	88	.49
1.060	8	12.0	4.10	5.29	48	56
1.067	9	13.4	4.55	5.87	49	68
1.075	10	15.0	5.08	6.55	55	70
1.083	11	16.6	5.67	7.31	61	79
1.091	12	18.2	6.20	8.00	68	87
1.100	13	20.0	6.78	8.68	74	95
1.108	14	21.6	7.30	9.42	81	104 112
1.116	15	23.2	7.80	10.06	87	123
1.125	16	25.0	8.50	10.97	96	184
1.134	17 18	26.8	9.18	11.84	104	144
1.142	19	28·4 30·4	9.80	12.64	112 121	156
1.152	20	32.4	10·50 11·14	13·55 14·37	121	167
1.162	20	84.5			137	177
1.171	22	36.0	11.78	15.13		188
1.180	23	88.0	12·33 13·00	15·91 16·77	146 155	200
1·190 1·200	24	40.0	13.70	17.67	164	212
1.210	25	42.0	14.40	18.58	174	225
1.210	26	44.0	15.18	19.58	185	239
1.231	27	46.2	15.96	20.59	196	253
1.241	28	48.2	16.76	21.42	208	266
1.252	29	50.4	17.55	22.64	220	283
1.263	80	52.6	18.35	23.67	232	299
1.274	31	54.8	19.23	24.81	245	816
1.285	82	57.0	20.00	25.80	257	882
1.297	88	59.4	20.80	26.83	270	848
1.308	34	61.6	21.55	27.80	282	364
1.320	85	64.0	22.35	28.83	295	881
1.332	86	66.4	23.20	29.93	309	899
1.345	87	69.0	24.20	81.22	326	420
1.357	38	71.4	25.17	32.47	342	441
1.370	39	74.0	26.12	83.69	859	462
1.383	40	76.6	27.10	34.96	875	483
1.397	41	79.4	28.10	86.25	892	506
1.410	42	82.0	29.05	37.47	410	528
1.424	43	84.8	80.08	88.80	428	553
1.438	44	87.6	81.00	89.99	446	575
1.453	45	90.6	32.10	41.41	466	602
1.468	46	93.6	33.20	42.83	487	629
1.483	47	96.6	84.40	44.38	510	658
1.498	48	99.6	35.79	46.15	535	691 721
1.514	49	102.8	86.90	47.60	559	750
1.530	50	106.0	88.00	49.02	581	700
Pegations 1 According to Deville (C. R.						

Reactions.—1. According to Deville (C. R. 45, 857), NaOH is decomposed to Na, O, and H by heating to white heat in an iron bottle.—2. Strongly heated with non-volatile acidic anhydrides, water and salts of Na are formed.—3. When molten NaOH is exposed for a considerable time to air or oxygen, some Na,O, is formed (Gay-Lussac; Thénard).—4. Moist or molten NaOH is decomposed by electrolysis, with production of Na (Davy, T. 1802. 1).—5. Heating with sulphur produces polysulphides, sulphite, and sulphate; S heated with NaOHAq forms polysulphides and sulphite. Similar reactions occur by heating with selenion or tellurium.—6. Chlorine interacts with NaOHAq to

form NaClOAq and NaClAq; on heating | NaClO, Aq is formed. Bromine reacts similarly. If Cl is led into NaOHAq containing iodine in suspension, NaIO, is produced.—7. When NaOHAq is warmed with phosphorus, H and inflammable P hydride are given off.—8. Heated with sodium, Na₂O and H are formed.—9. Moist NaOH withdraws carbon dioxide from the air, forming NaHCO₂.—10. Reacts with acids to form Na salts.—11. NaOHAq ppts metallic hydroxides, or oxides, from solutions of many metallic salts; in some cases the hydroxide dissolves in excess of NaOHAq-e.g. AlO₃H₂, ZnO₂H₂-12. Molten NaOH acts generally as an oxidiser; e.g. As, Sb, Fe, Pt, &c. form arsenate, antimonate, ferrate, and platinate of Na. Salts are generally decomposed by molten NaOH, giving Na salts, and setting free the bases .- 13. According to Schöne (A. 193, 241), addition of hydrogen peroxide to NaOHAq produces Na₂O₂.8H₂O (v. Sodium dioxide, p. 482).

Pickering (priv. comm.) gives the following data:—

P.c. NaOH	S.G. NaClAq at 15° (water at 4° =1)	P.c. NaOH	S.G. NaOlAq at 15° (water at 4° =1)	P.c. NaOH	S.G. NaClAq at 15° (water at 4° =1)
0	0.999180	17	1.88707	34	1.373453
1	1.010611	18	1.199783	35	1.383815
2	1.021920	19	1.210861	36	1.894092
3	1.033109	20	1.221933	37	1.404279
4	1.044317	21	1.233062	38	1.414363
5	1.055463	22	1.244119	39	1.424353
6	1.066602	23	1.255134	40	1.434299
7	1.077733	24	1.266092	41	1.444161
8	1.088856	25	1.277063	42	1.453929
9	1.099969	26	1.287990	43	1.463623
10	1.111069	27	1.298877	44	1.473249
11	1.122165	28	1.809708	45	1.482850
12	1.133250	29	1.320496	46	1.492406
13	1.144353	30	1.331213	47	1.501927
14	1.155450	31	1.841879	48	1.511412
15	1.666538	82	1.352472	49	1.520868
16	1.177619	33	1.362991	50	1.530282

Combinations.—1. With water to form hydrates. The hydrate 2NaOH.7H₂O was obtained by Hermes (B. 3, 122) by exposing NaOHAq S.G. 1'365 to the cold of a severe winter; monoclinic crystals, S.G. 1'405, melting at 6°; in vacuo gave off 3H₂O. Göttig (B. 20, 543) obtained a dihydrate, NaOH.2H₂O, by heating NaOH in 96'8 p.c. alcohol very gradually to 100°. By cooling NaOHAq, Pickering (C. J. 63, 890) obtained the following hydrates, with the freezing-points noted:—

Hydrate	Freezing-point
NaOH.H.O	64.30
NaOH.2H.O	12.5
NaOH.3·11H ₂ O (sic)	2.73
NaOH.8.5H,O	1 5·55
aNaOH.4H,O	7 ·57
BNaOH.4H,O	- 1·7
NaOH.5H.O	-12.22
NaOH.7H.O	-23.51

2. With carbon dioxide to form NaHCO₂.
Sodium, iodide of, NaI. Mol. w. not known with certainty, but formula NaI is probably Voz. IV.

molecular. Melts at 628° (Carnelley, C. J. 33, 278); at 650° (V. Meyer a. Riddle, B. 26, 2443). S.G. 3·45 (Filhol, A. Ch. [3] 21, 415); 3·654 at 18·2 (Favre a. Valson, C. R. 77, 579). S.H. (26° to 50°) ·0881 (Schuler, P. 136, 70); (16° to 99°) ·08684 (Regnault, A. Ch. [3] 1, 129). Kremers (P. 108, 120) gives the following data for solubility in water:—

Temp.	S.	Temp.	s.
0°	158.7	80°	803
20	178.6	100	812.5
40	208.4	120	322.5
60	256.4	140	888.8

S. = $264 \cdot 19 + \cdot 3978t$, when t varies from $64 \cdot 7^{\circ}$ to $138 \cdot 1^{\circ}$ (de Coppet, A. Ch. [5] 30, 411). Gerlach (Fr. 8, 285) gives following:—

S.G. NaIAq	P.C. NaI	S.G. NaIAq	P.O. NaI
1.04	5	1.36	35
1.082	10	1.432	40
1.128	12	1.51	45
1.179	20	1.6	50
1.234	25	1.7	55
1.294	80	1.81	60

Easily sol. alcohol. Saturated NaIAq boils at $141\cdot1^{\circ}$. H.F. [Na,I] = 69,080; [Na,I,Aq] = 70,300 (Th. 3, 232).

Formation.—1. When Na and I are fused together only very small quantities combine (Merz a. Weith, B. 6, 1518).—2. By decomposing BaI₂Aq, or CaI₂Aq, by Na₂CO₃ or Na₂SO₄, filtering, and evaporating.—3. By neutralising HIAq by Na₂CO₃, and evaporating.—4. I is added to water and iron filings till the iron is almost all dissolved, the solution is filtered, and Na₂CO₃ is added so long as FeCO₃ ppts.; the liquid is filtered (if alkaline it is neutralised by HIAq) and evaporated, any Fe₂O₃ which separates being filtered off (Baup, J. Ph. 9, 37, 122).

Preparation.—1. Iodine is added to mode-

rately conc. NaOHAq till a yellow colour is produced; finely-powdered charcoal, equal to c. 100 of the weight of I used, is added; the liquid is evaporated to dryness and the residue is heated in a covered crucible to dull redness for some time; when cold, the residue is dissolved in water, the liquid is filtered (neutralised by HIAq if alkaline) and crystallised at 40°-50°. (For more details v. Potassium iodide, Preparation, p. 304.)—2. A quantity of NaOHAq is divided into two equal portions; I is added to one part till a yellow colour remains, an equal quantity of I is then added, and then the other portion of NaOHAq is saturated with SO₂ and added, and the whole is evaporated till NaI crystallises out; the salt is purified by recrystallisation from water at 40°-50° (Stephani, J. Ph. [3] 26, 450).

Properties.—Crystallises from aqueous solutions at 40° to 50° in cubes (Mitscherlich, P. 17, 385); the hydrate NaI. 2aq separates at ordinary temperatures. Melts at 628°; volatilises less readily than KI, but at a lower temperature than NaCl (Mohr, A. 21, 66). Deliquesces in moist air, becoming rose-coloured with separation of some I and formation of some Na₂CO₂ (Girault, J. Ph. 27, 890).

Reactions and Combinations.—1. Heated in air some I is separated and O absorbed, and the reaction becomes alkaline (Berselius, Lehrbuch

II

[5th ed.] 8, 215).—2. Heated with charcoal in contact with air much Na, CO₂ is formed (Girault, l.c.). Most of the reactions of NaI are similar to those of KI (v. Potassium iodide, Reactions, p. 304).

Combines with most iodides of less positive metals. Forms a compound with arsenious oxide NaI.As,O₆ (Rüdorff, B. 21, 3051). The dihydrate NaI.2H₂O oxystallises at the ordinary temperature from NaIAq in large monoclinic prisms; S.G. 2·448; melting at c. 65° and giving NaI; effloresces in dry air, and deliquesces in moist air.

Sodium, manganicyanide of, v. vol. ii. p. 342.

Sodium, manganocyanide of, v. vol. ii. p. 342.

Sodium, nitride of, Na₂N. A dark-grey solid; formed by heating NaNH₂ (v. Sodium AMIDE, p. 475) to redness in absence of O. Resembles potassium nitride (q. v., p. 304) (Gay-Lussac a. Thénard, A. Ch. 65, 325).

Sodium, nitroprusside of, v. vol. ii. p. 341.

Sodium, oxides of. Two oxides have been isolated, Na₂O and Na₂O₂.

Sodium onder Na₂O. (Sodium monoxide.) It is doubtful whether this compound has been obtained pure. Davy (T. 1808. 24) said that when Na is fused with dry soda there is 'a division of the oxygen between the alkali and the sodium.' Na does not combine with O when exposed to quite dry air at the ordinary temperature (von Bonsdorff, P. 41, 296). According to Beketoff (v. B. 12, 856; 16, 1854), NaOH and Na do not react at a red heat. Beketoff (J. R. 1883. [i.] 277 (v. B. 16, 1854)) allowed Na to drop into a red-hot copper cylinder, then passed in a mixture of 1 vol. O and 4 vols. air, and heated by a blowpipe; in this way he obtained Na₂O, but always mixed with some Cu and traces of Na; considering the conditions under which Na₂O₄ is obtained (v. infra, Sodium dioxide) it is likely that B.'s Na₂O contained Na₂O₂.

Na₂O is described as a greyish solid; Beketoff's preparation was rose-coloured, but it contained Cu. Na₂O dissolves in water to form NaOHAq; B. gives [Na²O,Aq] = 55,500; and trom this and other data Thomsen (Th. 3, 232) calculates [Na²,O] = 99,760. According to B. (l.c.), hot Na₂O reacts with H to form NaOH and Na; heated to c. 300° in CO it gives Na₂CO₂ and Na (B., l.c.). Combination with CO₂, to form Na₂CO₃, occurs at c. 400°. By heating Na amalgam with HgO, B. (l.c.) obtained the compound Na₂HgO₂ = Na₂O.HgO. Comey and Jackson (B. 21, 1589; Am. 11, 145) describe various compounds of Na₂O with ZnO and H₂O.

Sodium dioxide Na₂O₂. (Sodium peroxide.) The formation of an oxide with more O than Na₂O, by heating Na in air, was noticed by Gay-Lussac a. Thénard (A. Ch. 65, 325). Pure Na₂O₂ was obtained by Vernon Harcourt (C. J. 14, 267) by heating Na, in a flask of hard glass filled with N, till the Na melted, then sending in a slow stream of dry air, and heating till the metal was changed to a yellowish-white solid, and heating this for some time in dry O. Na₂O₂ is a white solid, becoming yellowish on heating. It deliquesces gradually in air, and then is gradually changed into solid Na₂CO₂. Soluble water,

with production of much heat; on heating O is evolved; Na₂O₂Aq deposits crystals of the hydrate Na₂O₂.8H₂O on standing over H₂SO₄ (v. infra, Hydrates). Na₂O₂ acts as an energetic oxidiser when heated; C, I, P, S, Sn are oxidised, more or less rapidly; heated with CO, Na₂CO₃ is formed; with N₂O the products are NaNO₂ and N (v. Harcourt, i.c.).

Hydrates of sodium dioxide. The octohydrate, Na₂O₂8H₂O, was obtained by Harcourt (l.c.) by evaporating a solution of Na₂O₄ in water over H₂SO₄. Fairley (C. J. [2] 16, 125) obtained the same hydrate by adding H₂O₂Aq to NaOHAq and ppg. by alcohol. Schöne (A. 193, 241) also obtained the octohydrate by the action of H₂O₂Aq on NaOHAq, evaporating in vacuo, drying on a porous tile, washing with 90 p.c. alcohol, and drying between filter paper. By using a considerable excess of H₂O₂Aq, Schöne (l.c.) obtained crystals to which he gave the composition Na₂H₁O₈.4H₂O = Na₂O₂.2H₂O₄.4H₂O; on drying over H₂SO₄, Na₂O₂.2H₂O₂ remained. The dihydrate Na₂O₂.2H₂O is obtained by drying the octohydrate for some time over H₂SO₄.

Sodium, phosphide of. A compound of Na and P was supposed by Gay-Lussac a. Thénard, and also by Davy, to be formed by heating the elements together in N. Vigier (Bl. [2] 3, 7) recommends to place a piece of Na in rock oil, boiling at c. 120°, in a retort, to add P little by little (much heat is produced and some oil distils) till there is an excess and some crystallises on the cold parts of the retort, to distil off the oil from the black phosphide formed, to wash with CS₂, and dry in a stream of CO₂.

The phosphide may be kept in dry air; in moist air, water, or acids it gives off inflammable P hydride (v. also Lüpke, C. C. 1890. [ii.]

Bunsen (A. 138, 292) obtained a phosphide of Na by strongly heating a mixture of Na₂HPO₄ and Na in a narrow glass tube.

Sodium, platinosyanide of, v. vol. ii. p. 344. Sodium, platinosulphocyanide of, v. vol. ii. p. 351.

Sodium, salts of. Compounds formed by replacing H of acids by Na. These compounds belong to the type NaX where X is a monovalent acidic radicle. The chief salts of Na derived from oxyacids are antimonate, arsenate, bromate and hypobromite, carbonates, chlorate, perchlorate, chlorite and hypochlorite, chromate and dichromate, indate and periodates, manganate and permanganate, molybdates, nitrate and nitrite, phosphates, phosphites and hypophosphite, selenates and selenites, silicates, sulphates, sulphites, thiosulphate and thionates, tellurates and tellurites (v. Carbonates, Nitrates, Sulphates, &c.).

Sodium, selenides of. Two selenides have been isolated, Na₂Se and Na₂Se₂, corresponding with two of the sulphides and with the two oxides.

SODIUM MONOSELENIDE Na.Se. Obtained by mixing well-cleaned Na rubbed to powder with the proper proportion of Se (Rosenfeld, B. 24, 1658). Uelsmann (A. 116, 127) says Na.Se is obtained in large colourless crystals, by saturating NaOHAq with H.Se, then heating in a stream of H and allowing to cool; it seems probable that these crystals were a hydrate of Na.Se (s.

infra). Fabre (C. R. 102, 613) obtained Na₂Se by heating 2Na₂Se.9H₂O (v. infra, Hydrates) in a current of N to not above 400°; F. describes Na₂Se as a hard, white, crystalline, deliquescent solid, becoming reddish-brown when fused; owing to its action on glass, F. did not get Na₂Se free from silica and alumina.

Hydrates of sodium monoselenide. By passing a rapid stream of H₂Se into NaOHAq (1 part NaOH in 4 water) Fabre (l.c.) obtained Na₂Se.16H₂O, melting at 40°, giving Na₂CO₃, a little Na₂SeO₃, and Se on exposure to air. By using more conc. NaOHAq (3:1), and keeping the temperature down, F. obtained Na₂Se.9H₂O; and when very conc. NaOHAq was used, he obtained 2Na₂Se.9H₂O.

For data regarding heats of formation of Na₂Se and hydrates, v. Fabre, C. R. 102, 703.

Sodium diselenide Na₂Se₂. Wöhler and Dean (A. 97, 1) obtained a selenide of K by heating K₂SeO₃ with charcoal; Rathke (A. 152, 211) showed that the selenide was K₂Se₂, and Jackson (B. 7, 1277) using Na₂SeO₃ and charcoal obtained the corresponding selenide of sodium.

Sodium, seleno-antimonate of, v. vol. i.

p. 286.

Sodium, selenocyanide of, v. vol. ii. p. 348.
Sodium, silicofluoride of, Na₂SiF_s. Prepared by adding H₂SiF_sAq to conc. NaClAq, washing, and drying. Gelatinous when ppd., but becomes crystalline on drying. S.G. 2·7547 at 17·5° (Stolba, Fr. 11, 199). S. ·65 at 17·5°; 2·46 at 100° (S., Lc.). Insol. alcohol. Melts at red heat, giving off SiF₄ (Berzelius).

Sodium, sulphides of. Several compounds of Na and S exist, but there is considerable doubt as to the compositions of some of the substances that are described as definite sul-

phides of Na.

Sodium monosulphide Na2S. Obtained by passing H2S over dry NaOH, the heat produced suffices to remove H2O formed (Kircher, A. 31, 339). Rosenfeld (B. 24, 1658) says that Na_2S is formed when 1 part thoroughly clean Na is rubbed to powder and mixed with 8 parts NaCl and '7 part S; the combination is sudden, and light is produced; pressure must be avoided in mixing the substances. By heating 100 to 105 parts Na₂SO₄ with 20 parts charcoal, a fleshcoloured mass of Na2S with some C is obtained; the reduction may also be effected in H. By dissolving the product of reduction in water and crystallising, the hydrate Na₂S.9H₂O is obtained; by dissolving the crystals again, heating with Cu (to decompose polysulphides), and evaporating to dryness in a stream of H, Na₂S remains (Priwoznik, A. 164, 69).

Na₂S is also obtained, fairly pure, by gently heating Na₂S.9H₂O in a stream of H (Weyl, P. 123, 362; cf. Sabatier, A. Ch. [5] 22, 5).

Na₂S is an amorphous, flesh-coloured, deliquescent solid, with a strongly alkaline reaction; easily sol. water, solution being colourless if polysulphides are quite absent. According to Weldon (v. Lunge's Soda-industrie, 2, 311), Na₂S is not fusible if it be quite free from polysulphides, thiosulphate, and NaOH.

Na SAq becomes yellow in moist air from formation of polysulphides (Kolb, A. Ch. [4] 10, 106); when air is blown into Na SAq, NaOH

and Na₂S₂O₂ are formed, and then Na₂SO₄ (Lunge, *l.c.*, p. 531). Na₂S is decomposed by acids, even by CO₂, with evolution of H₁S (v. Stromeyer, A. 107, 372). Heating Na₂SAq with NaHCO₃ or NH₄·H.CO₃ produces complete decomposition to Na₂CO₂ (Lunge, *l.c.* p. 300). Boiling with AlO₂H₁, or heating Na₂S with Al₂O₃, forms Na aluminate (v. W. J. 1863. 713; 1865. 332). CaO is said not to decompose Na₂SAq (v. Kolb, A. Ch. [4] 10, 106). Several metallic oxides decompose Na₂SAq, forming NaOHAq, sometimes (e.g. with CuO) Na₂S₂O₂Aq is also formed. Na₂S combines with various metallic sulphides (v. Schneider, P. 138, 302; 151, 446; Völcker, A. 59, 35; Berthier, A. Ch. [2] 22, 245).

Hydrates of sodium monosulphide.
The hydrates Na₂S.9H₂O, Na₂S.10H₂O, and
Na₂S.5H₂O have been formed. The hydrate with
9H₂O is most easily obtained by saturating one
half of a solution of NaOH with H₂S, adding
the other half, and evaporating, when large
colourless crystals separate (Rammelsberg, P.
128, 172); S.G. 2·471 (Filhol, A. Ch. [4] 28,
529). The pentahydrate is formed similarly
to the hydrate with 9H₂O, but using alcoholic
in place of aqueous NaOH (Böttger, A. 223, 335;
v. also Finger, P. 128, 635; Lemoine, C. R.
98, 45). For the preparation of Na₂S.10H₂O
v. Damoiseau (C. C. 1885. 36). Göttig (J. pr. [2]
34, 229) by partially saturating NaOH in alcohol
with H₂S got hydrates with 5, 5₁, and 6 H₂O.
Sodium disulphide Na₂S₂. This compound

Sodium disciplina Na₂S₂. This compound is supposed by Sabatier (A.Ch. [5] 22,5) to exist in the solution obtained by heating Na₂SAq with solution of polysulphides formed by digesting Na₂SAq with S. Böttger (A. 223, 335) obtained the hydrate Na₂S₂5H₂O by dissolving S in an alcoholic solution of No.

alcoholic solution of Na2S.

SODIUM TRISULPHIDE Na₂S₂ is formed, according to Schöne (J. 1867. 190), mixed with Na₂SO₄, by strongly heating Na₂CO₂ with excess of S (cf. Sabatier, l.c.). For the hydrate Na₂S₂.3H₂O v. Böttger (l.c.).

SODIUM TETRASULPHIDE Na₂S₄ is said to be obtained by heating Na₂S₅ (Chapman Jones, C. J. 37, 461; cf. Sabatier, l.c.). For hydrates v. Schöne (P. 131, 380) and Böttger (l.c.).

The existence of a pentasulphide is doubtful (v. Schöne, l.c.; Sabatier, l.c.; Geuther, A. 226,

232; Chapman Jones, l.c.).

Soda liver of sulphur. This name is given to the brown solid obtained by heating S with Na₂CO₂ in a closed vessel; it contains various sulphides of Na, along with Na₂S₂O₃, Na₂SO₄, and generally some Na₋CO₋.

and generally some Na,CO_s.

Sodium, sulphocyanide of, v. vol. ii. p. 352.

Sodium, sulphydrate of, v. Sodium hydro-

SULPHIDE, p. 479.

Sodium, telluride of. According to Rosenfeld (B. 24, 1658), well-cleaned Na in powder readily combines with Te.

Sodium, thio-arsenates and thio-arsenites of, v. vol. i. pp. 316, 317.

Sodium, thio-carbonate of, v. vol. i. p. 708. M. M. P. M.

SODIUM ETHIDE NaC,H,. This compound is not known in the free state. Sodium dissolves in cold ZnEt,, separating zinc. The solution, cooled to 0°, deposits ZnEt,NaEt in trimetric tables [27°], sol. benzene, and decomposed by water. CO₂ acting on ZnEt,NaEt

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forms sodium propionate (Wanklyn, A. 107, 125; 108, 67). Etl attacks ZnEt, NaEt, forming ethane and O.H. (Frankland, A. 110, 107). Sodium methide may be obtained in like manner (Wanklyn, A. 111, 234)

SOJA OIL. The oil from Soja hispida contains tripalmitin and tristearin (Meisel a. Böcker,

M. 4, 365)

SOLANINE C₁₂H₂₈NO₁₈ (?) (Firbas, M. 10, 541); C₄₂H₇₁NO₁₈; C₄₂H₃₇NO₁₈ (Hilger, A. 195, 817); C₂₁H₃₈NO, (Kletzinsky, Z. [2] 2, 127). [244°]. Occurs in the berries of the deadly nightshade (Solanum nigrum) (Desfosses, J. Ph. 6, 374), in berries of S. verbascifolium (Payen a. Chevallier, J. Chim. Med. 1, 517), in small quantity in the tubers and green parts of the potato (S. tuberosum), in the shoots of sprouting potatoes (Baup, A. Ch. [2] 81, 109; Otto, A. 7, 150; 26, 232), to the extent of '03 p.c. in diseased potatoes (Kassner, Ar. Ph. [3] 25, 402), in the flowers, stalks, and berries of the woody nightshade (S. Dulcamara), and in the root of Scopolia japonica (Martin, Ar. Ph. [3] 13, 336).

Preparation.—1. The berries are pressed,

the juice ppd. by ammonia, and the pp. re-crystallised from alcohol.—2. Fresh potato sprouts (200 kilos.) are digested with 2 p.c. acetic acid; the filtrate is ppd. by ammonia; and the pp. extracted with boiling 85 p.c. alcohol. To the hot alcoholic filtrate NH, Aq is added until turbidity appears. On cooling, a mixture (125 g.) of solanine and solanein is deposited. These bases are separated by fractional crystallisation from hotspirit (Firbas; cf. Reuling, A. 30, 225; Zwenger a. Kind, A. 109, 244; 118, 129; Kromayer, Ar. Ph. [2] 116, 114; Missaghi, B. 9, 83; Gmelin, A. 110, 167).

Properties.—Colourless needles, v. sol. hot 85 p.c. alcohol, insol. CHCl, and ligroin, sol. HClAq, almost insol. water and ether. The crystals are C₅₂H₅₅NO₁₈4½aq (Firbas). Poisonous. Not affected by alcoholic potash. Sublimes at 190° (Blyth). Gives an orange colour with I and H₂SO₄. On heating with 2 p.c. with I and H.SO4. On heating with 2 p.c. HClAq it yields solanidine and a dextrorotatory sugar that reduces Fehling's solution, $[a]_p = 28.6^\circ$, yields with phenyl hydrazine an osazone [199°], and is oxidised by HNO, to mucic and saccharic acids; lævulose is not present. Solanine colours a mixture of alcohol and H2SO, rose red. With ammonium selenite (1 g.) in conc. H2SO, (20 c.c.) it gives a canaryyellow colour turning to brown and, after three hours, to violet-red (Ferreira da Silva, Bl. [3] 6, 87; C. R. 112, 1267). A solution of ammonium vanadate in H₂SO₄ is coloured brown by dry solanine, the colour becoming red, and finally disappearing on dilution (Mandelin) Salts.—CaH,1NO16HCl. Got

Got by adding ether to an alcoholic solution of the base and HCl. Gelatinous pp. which dries up to a gummy mass.—B'₂H₂PtCl₄: yellow flocculent pp.—B'H₂SO₄: amorphous, v. sol. water.—B'₂H₂C₂O₄7aq; crystalline crusts.

Acetyl derivative C42HasAc6NO18. Needles

(from ether), insol. water (Hilger).

Solanein C. H. NO., 4aq. [208]. Obtained as above (Firbas). Amorphous. More sol. 85 p.c. alcohol than solanine. Decomposed by HClAq into the same products as solanine. Colours Mandelin's reagent red.

Solanidine $C_{40}H_{81}NO_{2}$. [191°]. Formed as above. Colourless needles (from ether). Affects Mandelin's reagent in the same way as solanin. Ac₂O at 140° gives a diacetyl derivative (Firbas). Converted by fuming HClAq in the cold into amorphous yellow solanicine C.H.NO (?), which yields B'HCl and B'2H2PtCl (Zwenger, A. 123, 341).

Salts. B'a(HCl), aq: crystalline powder.

B'4H2SO, 8aq: colourless plates [247°].

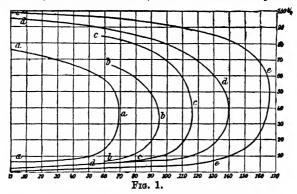
SOLUTIONS. Inasmuch as many of the questions connected with the state of bodies in solution are not yet fully elucidated, and inasmuch as two very different conceptions are put forward regarding the compositions of bodies present in aqueous solutions of salts, the editor of that portion of the Dictionary to which the subject of solutions belongs thought it advisable to ask the originator of the electrolytic dissociation hypothesis of saline solutions to write a short article on solutions in general, and to invite one of the leading upholders of the hydrate hypothesis of saline solutions to summarise the arguments in support of that hypothesis.

SOLUTIONS I. A solution is a homogeneous mixture of two or more bodies in the liquid state. A distinction is made between the dissolved substance and the solvent; the substance present in greater quantity is generally spoken of as the solvent. In some cases the solvent is taken to be the substance with the lower temperature of solidification. In the cases of substances miscible in all proportions—a mixture, for instance, of equal parts of alcohol and water -there is no way of determining which should be regarded as the solvent and which as the dissolved substance. There are many instances of two bodies that cannot be mixed in all and any proportions; the solubility of such bodies is said to be limited. If the 'dissolved body' in such a case is present in excess, either as gas, liquid, or solid, a condition of equilibrium is finally attained wherein the solution can take up no more of the dissolved body as long as the external conditions (temperature and pressure) remain unchanged; such a solution is said to be saturated. The attainment of saturation is hastened by shaking.

Saturated solutions of gases. When a gas is placed over a liquid, such as water or alcohol, a portion of the gas dissolves in the liquid. There is a constant relation at constant temperature between the quantity of gas remaining per unit volume and the quantity dissolved by the liquid per unit volume (Henry's law). Inasmuch as the quantity of the gas, per unit volume, is pro-portional to the partial pressure of the gas, the law may be stated by saying that the quantity of the gas dissolved in the liquid is proportional to the partial pressure of the gas above the liquid. As all the gases that have been examined dissolve with production of considerable quantities of heat, the solubilities of these gases decrease as temperature increases, in accordance with the second law of thermodynamics. The measurements made by Bunsen (Gasom. Methoden [Braunschweig, 1877]) show that the solubilities of some gases are independent of temperaturefor instance, the solubility of H in water and of O and CO in alcohol. Henry's law holds good, as indicated by theory, only so long as the quantity of the gas per unit volume, both undissolved

and in solution, is small, and the molecules of the dissolved and the undissolved gas are of equal magnitude. Van't Hoff has made use of the law of Henry to prove the equality of the molecules of the same gas in the gaseous state and in dilute solution (Z. P. C. 1, 489 [1877]). When NH₂, HCl, and SO₂ are dissolved in water in large quantities these gases do not follow the

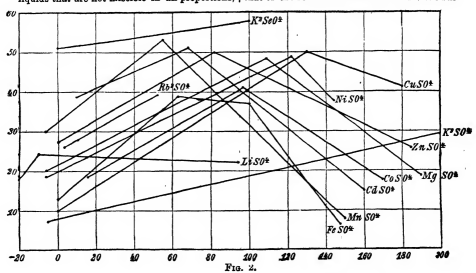
temperature, which corresponds to the critical temperature for the liquids, is passed, the two liquids dissolve in all proportions. This behaviour has been more fully examined by Alexengeff (W. 28, 305 [1886]) and has been represented by him in curves, some of which are reproduced here (fig. 1). The curve a represents the behaviour of water and phenol, b that of water



law of Henry; neither does the law hold good for solutions of acetic acid in benzene or ether in water, because the molecular magnitudes of these compounds are not the same in the gaseous state and in solution, as is proved by determinations of the boiling-points of the compounds in question (Nernst, Z. P. C. 7, 97 [1891]).

Mutual solutions of liquids. When two liquids that are not miscible in all proportions,

and salicylic acid, c that of water and benzoic acid, d that of water and aniline phenolate, and e that of water and aniline. The abscissæ represent temperature, and the ordinates percentages of the dissolved substances in the solutions. To each temperature correspond two points in the curve showing the percentage composition of the solution of the substance in water, and that of the solution of water in the molten sub-



such as ether and water, are shaken together, two layers are formed, one floating on the other; one of these contains much water and little ether, while the other contains much ether and little water. The compositions of the two layers change with change of temperature. Generally speaking, the compositions become more alike as temperature rises, until a temperature is resched whereat they are identical; when this

stance; thus, for phenol and water (curve a), at 20°, the solutions are composed of 92.5 p.c. water and 7.5 p.c. phenol, and 28 p.c. water and 72 p.c. phenol respectively. The two liquids are miscible in all proportions above 69°.

change with change of temperature. Generally speaking, the compositions become more alike as temperature rises, until a temperature is reached whereat they are identical; when this

SOLUTIONS.

solution of di-ethylamine in water—the convexity of the curve is turned to the left.

Saturated solutions of solids. Most of the measurements that have been made deal with solutions of salts in water. The example of Gay-Lussac has generally been followed, and the solubility of the salt has been taken as the weight thereof dissolved by 100 parts by weight of the solvent. Étard and Engel have given solubility-curves for very large variations of temperature (C. R. 98, 993, 1276, 1432; 104, 1614; 106, 206, 740 [1884-88]). Some of these results are shown in fig. 2, p. 485; the abscisse represent temperature and the ordinates parts by weight of the salts in 100 parts of the solution (not in 100 parts of the solvent). The change of solubility with change of temperature depends on whether the process of solution is accompanied by disappearance or production of heat; in the former case solubility increases, in the latter case it decreases, as temperature rises. Inasmuch as different hydrates of the same salt are dissolved with the production of different quantities of heat, these hydrates give different solubility-curves. Sodium sulphate is the salt that has been most carefully examined in this respect. The hydrate Na₂SO₄.10H₂O, that exists up to 83°, dissolves in water with disappearance of heat; hence the solubility of this hydrate increases as temperature increases. On the other hand, Na2SO4, stable above 33°, dissolves with production of heat, and its solubility decreases with increasing temperature. The so-called curve of the solubility of sodium sulphate consists, therefore, of two parts, one rising till the temperature reaches 38°, and one falling above that temperature. Looked at accurately, the first part of the solubility-curve holds good only for Na₂SO₄.10H₂O₅ and the second part only for Na, SO4. For the solubility of a chemical compound is determined only when the composition of the solid compound that is in contact with the solution is definitely stated. The compound CaSO, is considerably more soluble than the crystalline hydrate CaSO₄.2H₂O. The measurements made by Etard show that many other sulphates-MnSO. and CdSO4, for instance—behave similarly to Na₂SO₄ (see fig. 2).

The solubilities of most salts increase as temperature rises; the solubility of NaCl varies very little with temperature. The solubilities of many compounds of calcium decrease with increasing temperature, e.g. hydroxide, acetate, isovalerate, isobutyrate, succinate, and iso-succinate of calcium. Barium succinate and SrSO, behave similarly, but not the other corresponding salts of Ba and Sr. Some calcium salts show maximum solubilities at certain temperatures-e.g. CaSO₄.2H₂O at 35°; others show minimum solubilities—e.g. propionate at c. 55°, isobutyrate at 65°, also normal butyrate, valerate, iso-octoate, citrate, and benzoate (Allen, C. N. 57, 236 [1888]). Attempts have been made to express the connection between the solubilities of salts and temperature by a formula. The following expression holds good in many cases: S = a + bt (t = temperature, and a and b are constants); i.e. solubility changes linearly with temperature—for instance, for KCl, BaCl, NaNO, and several sulphates, especially at high temperaadd a third or a fourth term, and to adopt the formula $S = a + bt + ct^2 + dt^3$ (c and d are constants as well as a and b). The formula log. S = a + bt + ... + ... often gives a better representation with a smaller number of coefficients.

Supersaturated solutions. When a salt whose solubility increases with temperature is dissolved at a high temperature in water (or other solvent), and the solution is cooled, a temperature is reached whereat the solution is saturated with reference to a definite hydrate of the solid salt; but if the solution is further cooled salt does not separate, provided there is no crystal of the hydrate in question in contact with the solution. Such a solution is said to be supersaturated. A solution may be supersaturated with reference to one hydrate, but not supersaturated with reference to another hydrate. For example, Na₂SO₄.7H₂O is more soluble in water than Na₂SO₄.10H₂O₅, and so a solution may be obtained, by dissolving Na2SO4, that is supersaturated as regards the decahydrate but not as regards the heptahydrate; that is to say, if a crystal of Na₂SO₄.10H₂O is brought into the solution, at a certain temperature, the decahydrate crystallises out, but at the same time the solution is able to dissolve the hydrate with 7H₂O. If temperature falls again the solution, of course, becomes supersaturated with reference to both the hydrates.

At one time the supposition was often held that the dissolved substances were present in supersaturated solutions in a state different from that in which they existed in ordinary solutions. More accurate investigations of the physical properties on both sides of the point of saturation have shown that there is no essential difference between the solution before and after saturation, but that the relation of a supersaturated to a saturated solution is exactly the same as that of a saturated solution to the solution before saturation.

Colloidal solutions. Certain substances, such as gelatin, absorb unlimited quantities of water and become swollen thereby; when much water has been taken up, liquids are formed which do not part with the dissolved body on cooling, as ordinary solutions do, but set to jellylike substances. All compounds that form colloidal solutions have very large molecular weights; such compounds are acids of Mo, Si, Sn, Ti, and W; oxides of Sb, Fe, and Mn; sulphides of Sb, As, Bi, Cd, Co, Au, Fe, Pb, Hg, Ni, Pd, Pt, Ag, Tl, and W; many organic substances, such as caramel, dextrin, eggalbumen, tannin, &c.; and also some elements, such as Se and Ag. Some of these solutions are characterised by the fact that the addition of small quantities of foreign substances—as salts, acids, and bases--causes coagulation, whereupon the bodies in the colloidal solutions become insoluble. For that reason it is generally supposed that colloidal solutions are intermediate between emulsions and true solutions. Emulsions - e.g. a fine deposit of alumina-possess the special property that the emulsionised bodies are thrown down much more quickly in presence of electrolytes, and also of other foreign substances, than and several sulphates, especially at high tempera in absence of these (Barus a. Schneider, Z. P. C. three (Etard). In other cases it is necessary to 8, 291 [1891].) Colloidal substances having molecular weights under 30,000—gelatin or gum, for instance—are characterised by the fact that if their solutions are solidified by cooling they again become liquid and transparent when heated. Colloids having molecular weights greater than 30,000 do not share in this property, but behave, in this respect, like very supersaturated solutions (Sabanejeff, J. R. 1891. [1] 80; Z. P. C. 9, 89 [1892]). When colloids take up water, and swell but do not dissolve, heat is produced; when they dissolve heat disappears (Wiedemann a. Lüdeking, W. 25, 145 [1885]).

Reasons for assuming the existence of hydrates in solutions. When a dissolved substance is isolated, by various methods, from its solvent the solid generally separates in combination with water of crystallisation. It has, therefore, been generally supposed that a pre-ponderating quantity of this hydrate must be present in the solution. This conclusion is not, however, tenable; for if only a minimum quantity of the compound that separates out were present in the solution this might suffice to cause the separation. The following consideration has been brought forward as an argument in favour of the existence of hydrates in solutions. A solution of 57 parts by weight HI and 43 parts water boils at 127°, and the distillate has the same composition as the residue; this lends some countenance to the view that this mixture behaves like a definite chemical compound. On adding water to this solution and distilling a more dilute acid at first passes over, and at last the acid with 57 p.c. HI distils off. If the solution contains more than 57 p.c. HI to begin with, the distillate at first contains a more conc. acid, and, later, acid of 57 p.c. This behaviour is simply explained in the following way. A solution of 57 p.c. HI possesses a minimum vapourpressure, so that both less dilute and more dilute solutions at the same temperature have larger vapour-pressures. Speaking generally, the vapour from a solution does not contain both components in the same proportion as the solution itself. In the case under consideration the vapours from the more dilute solutions contain more water (in proportion to HI) than the solutions, and the vapours from the less dilute solutions contain, relatively, more HI. The dilute solutions increase in concentration as distillation proceeds, the boiling-point rises, and there is at last produced the 57 p.c. solution which possesses the lowest vapour-pressure at an equal temperature—i.e. the highest b.p. (127°) at an equal pressure. If, on the other hand, distillation is commenced with a more concentrated solution proportionally more HI passes over, and an approach is gradually made, from the other side, to the solution with highest b.p. There is, therefore, no binding reason for regarding this solution as a definite chemical compound. For, indeed, the composition of the solution changes according to the pressure under which the distillation is conducted; and this could not well be the case were the solution really a definite chemical compound.

In investigating the freezing-points and vapour-pressures of solutions it was found that in many cases the deviation from the freezing-point and vapour-pressure of the pure solvent

was proportional to the quantity of substance (salt) in solution. But in other cases this proportionality seemed not to be maintained unless the supposition were made that a portion of the water had combined with the dissolved substance (Wüllner, P. 103, 529; 105, 85; 110, 564 [1858, 1860]; de Coppet, A. Ch. [4] 23, 366; 25, 502; 26, 98 [1871-2]; Rüdorff, P. 114, 63; 116, 59 [1861-2]; 145, 599 [1870]). It was, therefore, supposed that such compounds as HI, HCl, BaCl₂, CaCl₂, NaBr, &c., were combined in aqueous solutions with definite quantities of water of crystallisation. But the recently-discovered laws which express these phenomena lead to very different methods of explanation, so that the conclusion that these compounds exist as hydrates in solutions seems, in this respect, to be entirely without foundation.

respect, to be entirely without foundation.

Mendeléeff not long ago examined the changes in the S.G., accompanying changes in the composition, of aqueous solutions of alcohol and of sulphuric acid (Z. P. C. 1, 273 [1887]); he thought he had found abrupt irregularities in the changes of S.G. He supposed, without any theoretical foundation, that at the concentrations whereat these occurred the solutions corresponded to perfectly definite hydrates. Pickering (Z. P. C. 6, 10 [1890]) showed, as the result of more accurate investigations, that Mendeléeff's conclusions rested on inaccurate observations. Pickering, however, accepted Mendeleeff's idea, and supposed that the higher derivatives of the S.G. in reference to percentage composition showed similar irregularities. He treated the freezing-points and the electrical conductivities of solutions in the same way, and he supposed that in all these cases he had discovered such irregularities as indicated the existence of definite hydrates. It is absolutely impossible to reconcile the numbers for the electrical conductivities of dilute solutions of acids and bases (e.g. for acetic acid) with Pickering's views; nor do the most recent investigations on the freezing-points of very dilute saline solutions (Jones, Z. P. C. 11, 534; 12, 623 [1893]) in any way agree with these views. Moreover, the method used by Pickering is not free from objection under any conditions.

From an extended and systematic investigation recently made into the constitution of ammoniacal metallic compounds, and compounds analogous therewith, Werner (Zeit. f. anorg. Chem. 3, 267 [1893]) concluded that in electrolytically conducting salt solutions the metallic atoms of the salts might be expected to be generally accompanied by six molecules of water, and that those salts which readily combine with water of crystallisation would be the best conductors. This statement is not, however, in any way in keeping with the experi-mental results. The salts of ammonium, K, Rb, and Cs conduct better than any others that have been examined; after these come the salts of Na, Li, and the metals of the earths; and then, much behind these, come the salts of the heavy metals; but water of crystallisation combines most readily with the salts last mentioned, and least readily with those mentioned first.

Conclusions regarding the existence of hydrates of substances in solution have also been drawn from the results of investigations of other

physical properties, such as contraction of volume, or production of heat, during solution, or viscosity, &c.

In general, it has been supposed that where these properties exhibited a maximum or minimum, or other purely mathematical characteristic, there existed a definite hydrate of corresponding composition. Attention may, however, be called to the fact that these points generally shift with changes of temperature, so that the hypothetical hydrate must have a different, and constantly changing, composition at different temperatures—a conclusion which is not in keeping with the representation of the hydrate as a definite chemical compound.

It is, indeed, in the highest degree likely that when definite hydrates separate from a solution the same hydrates were previously present in the solution, perhaps only in small quantity; but we have as yet no accurate knowledge as to the magnitude of the quantity. The methods that have been used in investigating this most interesting question have scarcely brought to light a single new result, although many and renowned investigators have carried out a very large amount of work in this direction.

The solubility of a salt, at constant temperature, is to a small extent dependent on pressure. Sorby was the first to carry out detailed investigations on this subject (Pr. 12, 538 [1863]). The following statement may be demonstrated by the use of the dynamical theory of heat. When the total volume of salt and solution is diminished (or increased) by the taking up of salt into the solution, then the solubility increases (or diminishes) with increased pressure. The researches of Braun (W. 30, 272 [1887]) have confirmed the theoretical conclusions.

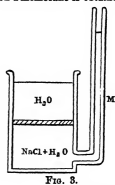
Dilute solutions. Osmotic pressure.—If a quantity of the pure solvent is floated on a solution, a movement of the dissolved substance takes place and the substance strives to distribute itself throughout the whole of the solvent. This occurrence is completely analogous with another; if a cylinder is filled with CO2 and the mouth is covered with a membrane, another cylinder is filled with H and placed mouthdownwards on the first, and the membrane is withdrawn, the two gases begin to diffuse into each other (notwithstanding the action of gravitation). The H is driven downwards by its partial pressure, and the CO2 is driven upwards. The velocity of diffusion is proportional to this propelling force, otherwise it is dependent only on the freedom of motion of the molecules among each other. The inverse value of this freedom of motion is called molecular friction. The following statement expresses the results of experiment: the velocity of diffusion is proportional to the difference between the partial pressure of one of the two gases in the upper and lower parts, and is inversely proportional to the friction of the molecules. Solutions behave in a similar The substance in solution in the under layer strives to distribute itself into the upper layer; this striving corresponds to the pressure of a gas, and, as will be shown immediately, it is equal to the so-called osmotic pressure. The is equal to the so-called osmotic pressure. solvent is driven downwards by the same force. In this case also the velocity of diffusion is proportional to the osmotic pressure, and is inversely

proportional to the friction of the molecules of the dissolved substance against the solvent. The only difference between this case and that of gases is that molecular friction is very much larger in liquids, a fact that is explained by the great number of molecules against which a molecule of the dissolved substance collides during its movements.

The amount of molecular friction is known in certain cases, viz. for electrolytes. Let a cubical trough (sides 1 cm.) be filled with the solution of an electrolyte (e.g. NaCl), and let an electric current be passed through the trough by the help of two electrodes A and B. The positive electricity is carried by the metallic parts (Na) of the electrolyte in the direction of the current (towards B); the negative electricity travels with the negative radicle (Cl) of the electrolyte towards A. If the Na and Cl atoms—or ions, as they are called in this case—are impelled by unit electrical force, the Na ions will move with a velocity v_1 and the Cl ions with the velocity v_2 . These velocities can be determined, partly indirectly from the conductivity and the migration numbers of Hittorff, partly directly by chemical Putting the corresponding frictions analysis. as r_1 and r_2 , then $v_1 = \frac{1}{r_1}$ and $v_2 = \frac{1}{r_2}$. The frictions of a large number of ions have been determined in this way. If we now have a layer of pure water floated on a solution of NaCl, in a vessel, then the velocity of diffusion, D, is equal to the quotient of osmotic pressure, O, and the frictions $r_1 + r_2$ of the salt. We have, therefore, $D = \frac{1}{r_1 + r_2}$. Nernst (Z. P. C. 2, 618 [1888]) has completely established the accuracy of this

formula.

Osmotic pressure can be measured directly. Let us suppose that in the foregoing example we had a division separating the solution of NaCl from the water, and that the division allowed water, but not NaClAq, to pass through it (a so-called semipermeable membrane); v. fig. 8. Such a membrane is obtained by impreg-



nating a plate of porous burnt clay with ppd. Cu₂Fe(CN)_e. The water will then be driven downwards, and will be compressed in the lower part of the apparatus; the pressure in this division will increase; this may be confirmed by using a manometer (M, fig. 8). Equilibrium is attained after a time, and the manometer then indicates a certain pressure, which is called the

osmotic pressure of the solution. When no more water is driven downwards, then, in the case in point, the partial pressures of the water in the upper and lower parts of the apparatus are equal. The excess of pressure in the lower partthat is, the osmotic pressure—is then equal to the partial pressure of the NaCl molecules. Measurements of this kind have been made by Pfeffer, Tammann, and Adie (Osmotische Untersuchungen, Leipzig, 1877; W. 84, 229 [1888]; C. J. 59, 844 [1891]).

Semipermeable membranes are found in the lining of living cells which incloses the contents of the cell. De Vries made observations with cells of Tradescantia discolor and Begonia manicata; Donders and Hamburger used bloodcells. If a cell is immersed in water the cell swells because of the entrance of water; if a soluble substance is added to the water, in constantly increasing quantity, the water will at last be driven equally inwards and outwards; if this limit is overpassed, the water passes out of the cell and the protoplasmic contents of the cell shrink together. The whole process may be \mathbf{T} he followed by the help of the microscope. limit reached immediately before shrinkage begins is characterised by the attainment of equality between the osmotic pressure of the cell-contents and that of the surrounding solution. By bringing similar cells into solutions of different substances it is possible, in this way, to determine the concentration that must be attained by two solutions in order that they may possess equal osmotic pressures. Using the results of Pfeffer and de Vries, van't Hoff made an examination of the magnitude of osmotic pressures. He found that the osmotic pressures exerted in their solutions by non-conductors of electricity—such as cane sugar, urea, glycerin, &c .- were exactly the same as the pressures which these substances would exert, in accordance with Avogadro's law, were they present as gases in the same volume as the volume occupied by the solutions. The osmotic pressure is therefore proportional to the concentration and the absolute temperature (law of Boyle and Gay-Lussac); this was confirmed by Pfeffer's investigations (van't Hoff, Handingar der Stockh. Ak. 21 [1886]; Ar. N. 20 [1885]; Z. P. C. 1,481 [1887]).

An explanation of gaseous pressure is found in representing it as the result of the bombardment of the walls of the vessel by the gaseous molecules. Similarly, osmotic pressure is thought of as resulting from the knocking of the molecules of the dissolved substance against the walls of the containing vessel. The gaseous laws hold good in all respects for osmotic pressure.

Vapour-pressures of solutions. Let there be a solution, say of cane sugar, in a trough, separated by a vertical semipermeable membrane from pure water, and let the membrane extend above the solution; the air above the liquid behaves exactly like the semipermeable membrane, inasmuch as it allows the water to pass through (as water-vapour), but it does not allow the cane sugar to pass through (because of the non-volatility of the sugar). The water has, therefore, a tendency to pass across through the air as well as through the semipermeable membrane from the water in the solution. This circumstance may be expressed by saying that the pressure of the water-vapour is greater over the water than over the solution. In like manner it follows that solutions which have equal osmotic pressures have also equal vapour-pressures. By making use of the second law of thermodynamics it can be shown (van't Hoff, l.c.) that a solution containing n molecules of dissolved substance to N molecules of solvent has a vapourpressure, P, which is given by the formula

$$P = \left(1 - \frac{n}{N}\right)p,$$

 $P = \left(1 - \frac{n}{N}\right)p,$ where p = the vapour-pressure of the pure sol-(The molecular weight of the solvent must be taken as that of the substance in the gaseous state at the same temperature.) formula was originally experimentally established by Raoult; it agrees well with experience (C. R. 103, 1125 [1886]).

Boiling-points of solutions. According as the vapour-pressure of a solution is less than that of the solvent (assuming that the dissolved substance possesses no marked vapour-pressure), so the solution begins to boil, under a definite external pressure, at a higher temperature than the pure solvent. The following formula may be deduced from the dynamical theory of heat:

$$\mathbf{E} = \frac{0.2 \, \tau^2}{\mathbf{L}} \cdot n,$$

where E is the increase in the boiling-point of the solvent brought about by dissolving n molecules of the substance in 100 grms. thereof, τ is the absolute boiling temperature, and L is the latent heat of vapourisation of 1 grm. of the solvent. Beckmann has contrived an apparatus for determining E, and therefore for finding n—that is, for determining the molecular weight of the dissolved substance when the quantity thereof in 100 grms. of solvent is known. He has shown that the formula gives results which agree closely with experience (Z. P. C. 5, 76 [1890]).

Freezing-points of solutions. When a solution is caused to freeze, in most cases only the pure solvent separates as a solid. The solid substance is in equilibrium with the solution at the freezing-point; hence, the vapour-pressures of the solvent over the solid and over the solution must be equal, as otherwise distillation would take place from the one to the other, and equilibrium would not be attained. For the sake of simplicity let us take water as the solvent. Ice and water have, therefore, the same vapour pressure at the freezing-point of the latter (\hat{O}°) ; but an aqueous solution has a smaller vapour pressure at O° ; hence at O° ice cannot have the same vapour pressure as an aqueous solution, but this equality occurs only at a lower temperature. In other words, the freezing-point of a solution is lower than that of the solvent. Van't Hoff (l.c.) has deduced the following expression from the dynamical theory of heat, $E = \frac{.02 \, \tau^2}{W}$. **, where E is the difference between the freezing-points of the pure solvent and a solution therein, which solution contains n gram-molecules of the dissolved substance in 100 grms. of the solvent, \(\tau \) is the absolute temperature, and W is the latent heat of fusion of

1 grm. of the solvent. This formula is of much

importance, for the molecular weights of very

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many substances have been determined by its help. It is immaterial in this case whether the dissolved substance exerts a considerable vapourpressure or not, for only the pure solvent freezes out. The following table gives some values calculated by the formula placed side by side with the values obtained by experiment, for the case where n=1:

		E observed	E calculate
Water	,	18.9	18.9
Acetic acid .		38.6	38.8
Formic acid		27.7	28.4
Benzene .		50.0	53.0
Nitrobenzene		70.7	69.5

Electrolytic dissociation. When determinations of molecular weights are made, by the foregoing methods, based on osmotic pressures (de Vries), freezing-points (van't Hoff, Raoult), depressions of vapour-pressures (Raoult), or increments of boiling-points (Beckmann), the values obtained for all those substances whose solutions conduct electricity are much smaller than would be expected. For instance, the results obtained by working with a half-normal solution of NaCl lead to the value 31.6 for the molecular weight of the salt, a number which is 1.85 times smaller than the calculated value, 58.5 (NaCl). From this and many other considerations Arrhenius drew the conclusion that the molecules of electrolytes are largely dissociated in aqueous solutions; about 85 p.c. of the NaCl, for instance, in an aqueous solution of this salt is regarded as dissociated into Na and Cl. This conclusion is completely analogous with those which are drawn regarding the dissociation of the molecules of ammonium salts and other substances, in the gaseous state, from observations of the abnormal vapour densities of these substances. The development of the theory of electrolytic dissociation, based on these considerations, and confirmed in the fullest way by experience, has lent much support to the theory of solutions which has already been sketched. This part of the subject is treated by Ostwald in the article ELECTRICAL METHODS, to which reference should be made (this vol., pp.

Some of the deductions regarding diffusion, solubility, and affinity must, however, be mentioned here.

Diffusion of mixtures. When a solution of HCl is placed in contact with pure water, the HCl gradually diffuses into the water. Inasmuch as most (almost all) of the HCl molecules are dissociated, one would expect the diffusion to result in a separation of the H and Cl from each other, just as the alums and the double salts of the sulphates of the alkalis and the magnesium metals, which are partially decomposed in aqueous solution, can be separated into their constituent salts in this way (Graham, T. 1850. 1, 805; Marignac, A. Ch. [5] 2, 546 [1874]). But this separation does not occur. The H moves into the water more quickly than the Cl, and, because of the positive electrical charge of the H ions, the water becomes positively and the solution negatively charged. If this charging is not carried off by metallically connected unpolarisable electrodes placed in the solution and in the water (v. this vol. p. 212), it prevents the separate diffusion of the H and the Cl. But there is another way whereby the restrain-

ing force of the electrical charge on the diffusion of the H may be diminished—namely, by addition of chlorides. For instance, the velocity of diffusion of HCl (properly speaking, of H) is increased in the ratio of 1:2-24 by adding 25 times the quantity of NH,Cl (Arrhenius, Z. P. C. 10, 51 [1892]). Similar, although not so strongly marked, characteristic phenomena, which find an explanation in the dissociation theory, have been observed by Graham and Marignac.

Decrease of solubility by addition of foreign salts. Nernst (Z. P. C. 4, 372 [1889]) was the first to draw attention to the conclusion from the theory that a slightly soluble salt, e.g. silver acetate, must be more soluble in pure water than in a solution that contains other silver ions (e.g. AgNO₃) or acetate ions (e.g. NaC₂H₃O₂). Similar relations are shown by gases which partially decompose (e.g. NH4SH); these exhibit smaller vapour-pressures when one of the two components (NH, or H2S) is added. This lowering of solubility can be calculated from the laws of mass-action, with results which agree well with experience (Noyes, Z. P. C. 9, 603 [1892]).

Division of a dissolved substance between two solvents. Berthelot and Jungfleisch examined the distribution of succinic acid between ether and water, and found that the concentrations of the acid in the aqueous and in the ethereal solution were in a constant proportion, which appeared to be independent of the absolute concentration. Many other substances were found to behave in the same way (A. Ch. [4] 26, 396, 408 [1872]). This result corresponds completely with Henry's law for gases. It is readily deduced from the theory; but in doing this it is assumed that the dissolved substance has the same molecular weight in both solutions. If this is not the case—as, for example, in the distribution of benzoic acid between benzene and water-altogether different laws express the phenomena (these laws, also, are in keeping with experi-

mental results; Nernst, Z. P. C. 8, 110 [1891]).

Reaction velocities. There are many socalled catalytic processes that are brought about by the H ions of acids-for instance, the inversion of cane-sugar, saponification of esters, &c. According to the theory, the velocity with which these reactions take place must depend only on the number of H ions that are present, and not on the nature of the reacting acid. This con-clusion is confirmed by experience (Arrhenius, Z. P. C. 4, 226 [1889]; cf. Affinity, vol. i.,

especially pp. 77-81).

Division of a base between two acids present in equivalent quantities. Thomsen made investigations regarding the quantity of a base (NaOH) that reacts to form salts with two acids (e.g. HCl and HF) added in equivalent quantities. He found that, in the case quoted, 95 p.c. NaCl and only 5 p.c. NaF were formed in dilute aqueous solution. It may be deduced from the theory that the quantities of NaOH combining with the two acids (HCl and HF) are in the same ratio as the extents of dissociation of the acids at the same dilution. The observations of Thomsen and of Ostwald agree extremely well with this deduction from the theory (Arrhenius, Z. P. C. 5, 14 [1890]; cf. AFFINITY, vol. i. p. 81).

Influence of temperature on rate of solution. If a finely and equally powdered substance is shaken with a solvent at two different temperatures, for a very short time, the rapidity of solution is found to increase rapidly with the temperature. For instance, the rates of solubility in water of cream of tartar and benzoic acid increase between 0° and 17° in the ratios 1:3.8 and 1:3.1.

So-called solid solutions. Palladium is able to take up much H, but the mixture retains the solid form. There is, therefore, here no proper solution; nevertheless, van't Hoff has shown that the laws which hold good for solutions are valid for this phenomenon also (Z. P. C. 5, 322 There is formed at first an alloy, [1890]). Pd₂H, which possesses a definite dissociation pressure. If the pressure of the H is increased more H is absorbed, and the quantity absorbed is in direct proportion to the excess of pressure of the H. Henry's law, therefore, holds good. On this ground van't Hoff called mixtures similar to this 'solid solutions.' Solid solutions of thiophene and benzene have been examined by van Bijlert (Z. P. C. 8, 343 [1891]).

Heats of solution. A considerable quantity of heat is generally produced during the solution of a substance in a solvent. The heat of solution of a substance is defined to be the quantity of heat that appears during the solution of a gram-molecule of the body in much water. The following table gives the heats of solution of some of the commoner substances. Most of the numbers are taken from Thomsen

1----

(Th. 3, 195).

Gases.				
Chlorine Cl			48	70
Carbon dioxide CO.			588	32
Ammonia NH.			84	30
Hydrogen fluoride HF			1180	00
" chloride HCl			173	10
" bromide HBr			199	10
iodide HI		_	192	10
" sulphide H ₂ S			45	
Sulphur dioxide SO ₂	•	•	77	
Liquids.				
Methyl alcohol CH ₃ .OH			200	
Ethyl " C ₂ H ₃ .OH			25	40
Propyl " C.H.,OH			30	50
Ether (C.H.),O .			59	40
Acetic scid C ₂ H ₂ O.OH			4	20
Sulphuric acid H2SO4			178	50
Solids.				12500
Caustic potash KOH .	• • • • • • • • • • • • • • • • • • • •	•	•	-80
" " hydrate KOH Lithium chloride LiCl .	.ZD ₂	•	•	8440
	•	•	•	
Sodium ,, NaCl .	•	•		-1180
Potassium , KCl .	•	•	•	-4440
Sodium bromide NaBr .) ot	÷ ^	•	-190
,, hydrate Nal	or.zr	120		-4710
Potassium sulphate K ₂ SO ₄		_	-	 6380
Mercuric chloride HgCl		•	-	0000
	•		-	-3300
Sodium acetate NaC ₂ H ₃ O ₂	•	:	-	4200
Sodium benzoate NaC,H,O,	•	:	•	4200 800
Sodium benzoate NaC, H, O ₂ Benzoic acid C, H, CO, H	•	•	•	4200 800 - 6700
Sodium benzoate NaC, H ₃ O ₂ Benzoic acid C, H ₃ .CO ₂ H Silver chloride AgCl	•			4200 800 6700 15800
Sodium benzoate NaC,H,O ₂ Benzoic acid C,H,CO ₂ H Silver chloride AgCl , bromide AgBr	•		-	4200 800 6700 -15800 -20200
Sodium benzoate NaC, H ₃ O ₂ Benzoic acid C, H ₃ .CO ₂ H Silver chloride AgCl	•		-	4200 800 6700 15800

Volume changes accompanying solution. In most cases the solution of a substance in water is accompanied by a considerable amount of contraction. For instance, a mixture of 100 c.c. alcohol with 100 c.c. water measures only 192.8 c.c. at 18°. The contraction may be so great that the volume of the solution is less than the volume of the water used as solvent: for instance, when Na₂CO₂, NaOH, or LiOH is dissolved in much water. The following table shows the volume change, in c.c., that takes place when one gram-equivalent of some of the most important compounds is dissolved in much water—e.g. in 10 litres:—

	\mathbf{H}	Na	ĸ	NH.
$\mathbf{o}\mathbf{H}$	18	- 5.8	3.6	
Cl.	18.3	16.6	26.9	37.4
NO,	29.0	28.0	38.5	48.2
₽SO.	16.2	6.4	15.7	$24 \cdot 2$

A mixture of 40 grms. NaOH with 10 litres water occupies, therefore, a volume of 9994.2 c.c.

When the solvent is other than water an increase of volume sometimes takes place; for instance, a mixture of 100 c.c. alcohol with 100 c.c. CS₂ occupies about 202 c.c.

Specific heats of solutions. The water value of a solution of a salt in water is not equal to, but is generally less than, the sum of the water values of the water and the salt. In a few cases this decrease is so great that the water value of the solution becomes less than that of the water used as solvent. The following data show the water values of one gram-equivalent of the chief salts in extremely dilute solutions (according to the determinations of Thomsen):-

The water value of a solution of 40 grms. NaOH in 10 litres water, for instance, is not only not greater than that of the water alone (10,000), but is distinctly smaller (9,973).

Deviations shown by concentrated solutions. The laws that have been stated for the osmotic pressure, the lowering of freezing-point, the raising of boiling-point, and the lowering of vapour-pressure, brought about by substances in solution, are valid, strictly speaking, only for very dilute solutions. Deviations from these laws occur when the solutions are more concentrated. Sometimes the observed values are smaller than those theoretically deduced. In such cases it is supposed that molecular aggregates are formed. This occurs, for instance, with solutions in benzene of oxims, alcohols, phenols, and fatty acids, and also with fairly concentrated aqueous solutions of the sulphates of Mg, Cu, Zn, and Cd, and CdI₂ (Beckmann, Z. P. C. 2, 787 [1888]; Arrhenius, Z. P. C. 2, 496 [1888]). On the other hand, most concentrated aqueous solutions give values greater than the calculated values; this is made especially evident by Tammann's investigations into the vapour-pressures of salt solutions at 100° (Mem. de l'Acad. de St. Pétersbourg [7] 1, 85 [No. 9, 1887]). An explanation of this behaviour has 800 been sought for in the attraction between the *olvent and the dissolved substance (Arrhenius, Z. P. C. 10, 40 [1892]).
S. A.

SOLUTIONS II. The view that hydrates exist in aqueous solutions, and analogous compounds in non-aqueous solutions, is one which has long been held by many chemists; it is only in the last few years, however, that the hydrate or association theory has assumed a precise form, and that definite experimental evidence in support of it has been accumulated. In giving some account of the present position of this theory it will be convenient to collect the evidence on which it depends under the headings 'general' and 'special,' and to discuss separately the bearing which recent work on dilute solutions has upon it. Aqueous solutions have naturally been more studied than solutions in other solvents, but these other solutions have been investigated sufficiently to show that they are similar to aqueous solutions in every respect, except as regards electric conductivity. Although, therefore, to simplify matters in the present article, in most instances aqueous solutions alone are mentioned, it must not be understood that the arguments do not apply equally to other cases.

General. The changes accompanying dissolution seem to be in every respect similar to the recognised accompaniments of chemical combination: evolution of heat, a more or less profound alteration in the nature of the reagents, and an irregular variation in the properties of the resulting solution with regular variations in the proportions of dissolved substance and solvent. For reasons which will be mentioned below, these irregularities may often not be very pronounced, but it may safely be stated that every attempt which has been made to express the properties of any series of solutions by a simple expression agreeing, within the limits of experimental error, with the observed values throughout a wide range of concentration has been unsuccessful.

The affinity which a large number of substances possess for water, as evidenced by the many solid compounds which they form with it, and by the heat evolved in their formation, renders it prima facie extremely improbable that these substances should exist in presence of excess of water without combining with it. It is, moreover, those very substances which exhibit the greatest tendency to form solid hydrates which are generally most soluble. A more careful study of the thermal phenomena of solutions places this argument beyond the range of mere speculation. From the known heat of fusion of water and of a few anhydrous salts we can obtain a very close estimate of what the heat of fusion of any hydrated salt would be, if no change beyond the mere passage from the solid to the liquid condition occurred; with hexahydrated calcium chloride, for instance, the heat of fusion under such circumstances would be 11,000 to 12,000 cal., and on the most exaggerated estimate could not exceed 14,000 cal., whereas if decomposition as well as mere liquefaction occurred the heat absorbed would be at least 25,000 cal., since the heat of fusion of the 6H2O alone is 9,500 cal., and the heat of combination of CaCl₂ with 6H₂O (both solid) is 14,500 cal. The observed heat of fusion, however, is only 11,417 eal., a value which disproves that any considerable change beyond mere liquefaction has occurred during the fusion—i.e. the combination which existed in the solid must exist also in the liquid. The data in the case of sulphuric acid afford more striking evidence of a similar character. The heat of combination of solid water with the solid acid to form the solid monohydrate is 6,550 cal., and if, when the reagents are mixed in the liquid condition, they do not combine chemically, far less than this amount of heat would be evolved; whereas the heat actually evolved is found to be almost identical with the above, namely 6,667 cal.

The separation of a crystalline hydrate from a solution is an argument to which due weight has hardly yet been attributed in favour of the existence of that hydrate in the solution, although not necessarily in large quantity. If there are no molecules of the hydrate existing as such in the liquid, these must have been deposited in the solid form at the moment of their formation, whereas we know that whenever deposition occurs simultaneously with formation as in the precipitation of an insoluble salt, the substance is deposited in the amorphous and

not in the crystalline condition.

If, as would appear to be the case, it is necessary to admit the existence of hydrates in concentrated solutions, it is necessary to admit their existence in dilute solutions also. The mass action of an excess of one or other of the constituents of any composite fluid is well known, and is universally accepted: it must operate with hydrates as much as with other substances, and must do one of three things: either (1) it must increase the stability or amount of the particular hydrate present, or (2) it must combine with it to form a higher hydrate (if it is the water of which excess is added), or (3) it may decompose the hydrate by reacting with the non-aqueous constituent of it to form new compounds, as when free alkali and acid are produced (a decomposition, however, which can be proved in many cases to occur to but a negligible extent), but in no case could excess of water decompose the hydrate taken so as to liberate the dissolved substance in the anhydrous condition. The thermal phenomena of solution afford important evidence in this case also; to interpret them properly, however, it must be remembered that the dissolution of a substance in excess of solvent entails the separation of the particles of that substance from each other to the same extent as if it were vapourised, and that to effect this separation the same amount of heat must be absorbed in either case. The heat evolved due to the reaction of a liquid with excess of solvent is, therefore, the observed heat of dissolution minus the heat absorbed in vapourisation, and, in the case of a solid, the heat of fusion as well as that of vapourisation will have to be subtracted. Making allowance for these so-called physical changes, we find that the formation of every concentrated solution—that is, a solution which contains hydrates—is accompanied by the evolution of heat, and inasmuch as, with the same allowance, dilution is always accompanied by a further evolution of heat, the action of the diluent cannot be regarded as a reversal of the initial reaction, but rather as an extension and completion of it, the hydrates present in the concentrated solutions being increased in amount or in complexity thereby. When matters are simplified by dissolving a gas, instead of a liquid or solid, and adjusting the pressure of the gas or the quantity of solvent so that the former occupies the same volume after as before dissolution, we still find that a very considerable evolution of heat occurs; with the haloid acids this is as much as 17,000 cal., and though this may be glossed over by some of the advocates of the physical theory by stating that the decomposition of the acids into their ions, which they consider occurs, evolves a very large amount of heat, no such explanation can be offered in cases where no such decomposition is imagined, and it has been proved that the dissolution of gaseous non-electrolytes both in water and in other solvents is accompanied by the evolution of as much as 5,000 to 14,000 cal. In the face of such a considerable loss of potential energy, it is impossible to maintain that the substances losing it remain unchanged, and that the dissolved substance can, in any true sense, be still gaseous and uncombined with the solvent. It is important to note that van der Waals, through a different line of reasoning, has arrived at the same conclusion as to the necessity of recognising some action evolving heat between the substance and solvent, even when the former is in the so-called dissociated condition.

Amongst the general considerations favouring the view that dissolution consists in the formation of liquid compounds, not the least important is that this gives us some intelligible reason for dissolution occurring at all, whereas if the solvent is regarded as being inert and only playing the part of so much empty space (as some of the advocates of the physical theory have maintained), we have no explanation of why dissolutions occur; the presentation of empty space to a stable solid does not make the latter fill the empty space, and even attributing dissolution to the possession of a certain 'solution pressure ' by the dissolving substance would appear to amount to no more than stating that a substance dissolves because it does do so.

Special. Berthelot determined the heats of dissolution of various series of solutions of different concentrations, and concluded that his results when plotted out showed the presence of changes of curvature or inflections at certain points, indicative of the existence of hydrates in the solutions. In many cases he adduced various special considerations in support of the existence of the hydrates thus indicated, but his determinations were not sufficiently numerous or accurate, nor were his methods of examining them sufficiently precise, to lead to more than vague conclusions. Thomsen's investigations on the same subject were equally unsatisfactory; he sought to disprove the existence of any such changes of curvature by finding empirical equations to represent each series of results, but in the four cases which he thus investigated the equations deduced express but a small portion of the whole series, leaving in the remainder errors ten and a hundred times greater than the experimental error. Mendeléeff next took up the question from a different point of view. Theoretical considerations led him to conclude that solutions consisted of different hydrates according to the

amount of water present, and that, if not more than two hydrates were ever present together in the same solution, there would be definite changes of curvature in the figures representing the properties of the solutions at the concentrations corresponding to these hydrates; that is, that the rate of change in the property with change of concentration would be different when the solutions consisted of the hydrates A and B from what it would be when they consisted of B and C, so that there would be a change at a concentration corresponding to the composition of B, and further, he concluded that these rates of change (first differential co-efficients) in the case of the densities would be rectilinear functions of the percentage composition between the points A and B, B and C, &c. (It should be noticed in passing that the idea which lies at the root of Mendeléeff s conception-and, to a certain extent, of Berthelot's also—is that the water in any solution is nearly entirely combined with the dissolved substance, and not merely that the solution consists of a simple hydrate mixed with excess of free That the concentration-rate of change water.) of the densities is representable within experimental error by a series of straight lines is probably true in many cases, but in the two cases on which Mendeleeff chiefly relied for proof these first differences are conspicuously curvilinear throughout. The reason of this, doubtless, is that solutions of a given concentration generally contain more than the two hydrates which he postulated. Roozeboom's work on ferric chloride, and Pickering's on sodium hydroxide, render it probable that five or six hydrates may co-exist in the same solution; at any rate, as many as four have actually been obtained from certain solutions. Pickering next attacked the subject from a purely experimental point of view. The distinctive feature of his work is the search for changes of curvature (breaks) by the application of a flexible lath to his plotted results. A lath bent by the application of two couples near its extremities forms, for all practical purposes, a curve of a very simple nature; and it has been proved that the use of such a lath is tantamount to, and leads to precisely the same conclusions as, the application of ordinary parabolas with three or four constants deduced mathematically from the experimental values; and in several respects, besides expeditiousness of application, this method is superior to the mathematical method. The mere fact that a certain figure may be represented accurately by a series of parabolas is by itself no proof that it really consists of these independent curves, any more than the existence of breaks can be disproved by finding an artificial equation which will bridge them over; the correctness of any particular form of representation can be measured only by the results to which it leads, and in this respect the discontinuous expressions would appear to be highly satisfactory. The positions of the breaks are not dependent on the taste of the draughtsman: the figures examined seem to split up naturally into certain sections only, and when drawn in these sections it was found, in some dozens of series of experiments, that the apparent error of the points agreed within 5 or 10 per cent. with the known experimental error (though in a great number of the cases the true-

magnitude of the latter was not ascertained till after the drawings were made), whereas drawings of a similar character, but placing the breaks at other points, or drawings obliterating the breaks altogether, represented the error of the points to be far larger, often 10 or 100 times larger, than the known experimental error. It must be remarked that, once the magnitude of the experimental error is known with certainty, no drawing can be accepted unless it agrees closely with it, and Pickering has devised a method by which the experimental error can be accurately determined independently of any considerations other than the examination of the final results themselves; he has also devised a means of obtaining a numerical estimate of the acceptability of any drawing, by combining together the various factors which are usually taken as affording a criterion of acceptability. Again, when sections of increasing lengths are taken, and these are represented by single parabolas, or bent-lath curves, there is little or no increase in the apparent error of the points till the sections extend beyond a point where one of the supposed breaks exists, but as soon as they do so there is a large and sudden increase, indicating that some real change at the point in question exists. Moreover, it has been shown that in a case where two parabolas will represent a series of results perfectly, a single parabola will not do so, even if it contain as many constants as the two parabolas together contain. The strongest argument, however, in favour of the reality of these breaks is that the various properties of any series of solutions, although they form figures differing from each other widely in general form, are yet all thoroughly concordant as to the positions at which the breaks occur. The properties of sulphuric acid which were investigated were the densities at four temperatures and the contraction on mixing deduced from them, the heat of dissolution, the thermal expansion, the electric conductivity (Kohlrausch's values), and the thermal capacity; subsequently also the freezing-points and van der Willigen's values for the refractive indices were investigated. Perkin also found indications of two of the breaks in his determinations of the magnetic rotations, and still more recently Féry has recognised some other of the breaks in his own measurements of refractive indices. Three different properties were also examined by Pickering in the case of calcium chloride and nitrate, and were found to show a similar concordance.

That the breaks are really due to, and indicate the presence of, compounds in solution is shown by the fact that, not only in the cases above mentioned, but also in many others where other solvents besides water were used, they always occur at points which correspond within small limits with definite molecular proportions -wherever, that is to say, the proportion of substance to solvent is sufficiently simple to permit of any conclusions at all being drawn in the matter. Further evidence on this point was also afforded from a study of the freezing-points of fourteen of the alkyl amines, where, with one exception, hydrates of the very compositions indicated by breaks in the case of some of the amines were isolated in the crystalline condition in the case of the others. Finally, as a more striking, though perhaps not more cogent, argument, we have the isolation in the crystalline condition of no less than six new hydrates, the existence of which had been foreshadowed in the properties (chiefly densities) of the solutions—namely, H₂SO₄·4H₂O, HBr.3H₂O, HBr.4H₂O, HCl.3H₂O, HNO₄·H₂O, and HNO₅·8H₂O—the last three being especially conspicuous examples, for the only marked breaks which the densities showed coincided with the only hydrates which were subsequently isolated.

Two important features of the conclusions drawn from the above-mentioned work on sulphuric acid should be mentioned: firstly, the large number (18-20) of hydrates of which indications were obtained; secondly, the complex nature of hydrates in very dilute solutions. Any improbability which might be considered to attach to the former conclusion (though, in reality, no data exist on which to found any estimate of the probabilities in such a case) is removed by the fact that a still larger number of hydrates (when equal ranges of concentration are compared) have been isolated in the crystalline condition from solutions of sodium hydroxide. The complexity of the hydrates indicated extends up to compounds with hundreds and even thousands of molecules of the solvent, but, with the explanation to be mentioned below as to their possible constitution, these hydrates can scarcely be branded as improbable; at any rate, the breaks which indicate their existence appear to be precisely similar in nature to those which indicate the presence of the simpler hydrates. Pickering has investigated the freezing-points of many dilute solutions besides those of sulphuric acid, and has found that nearly all of them show indications of similar changes, and those recognised in the cases of sodium chloride and sulphuric acid have received further confirmation from a series of results made by H. Jones; a confirmation of special importance, since Jones's determinations were made with the view of disproving the existence of breaks.1 Complex hydrates, though of a comparatively small degree of complexity, would appear to exist in regions other than those of very dilute solutions; two such were indicated in the neighbourhood of the monohydrate of sulphuric acid, and a similar one, perhaps another also, has been isolated in the case of soda. These are probably compounds of two different hydrates.

It is important to note that the changes of curvature here described need not necessarily be absolutely abrupt. On the strength of experimental evidence only, where experimental error necessarily exists, it is obviously impossible to prove or disprove the abruptness of any change, and no satisfactory mathematical theory has yet been formulated to lead to any views on the subject. It is sufficient for the purpose that comparatively abrupt changes exist. Nor should it be expected that these changes would necessarily be of a very marked character, for a large proportion of the factors determining the properties of solutions must be of a purely physical or even mechanical character; the densities of sulphuric acid solutions, for instance, must rise more or less gradually from 1 to 1.85, and it is only in

¹ It should be noted that Jones himself asserts that his results entirely disprove Pickering's 'breaks' (c. B. 26, 547, 1633).—M. M. P. M

the variations from the 'more' to the 'less' that we can expect to find evidence of inter-

mediate compounds. Application of the hydrate theory to explain the properties of dilute solutions. Diversity and irregularity are some of the most marked features of the properties of concentrated solutions, and though in dilute solutions these irregularities are much less conspicuous (as, indeed, we should expect them to be when the hydrates present become excessively complex and, consequently, excessively unstable), they are still recognisable, and it is only in the most extreme dilution, where the experimental error amounts to a large fraction of the total quantities measured, that the results can be expressed within the limits of this error as an apparently regular function of the concentration. It may safely be stated that any purely physical theory which confines itself to regions of extreme dilution, and which ignores some of the most marked features of dissolution, as well as the mass of direct evidence showing the existence of compounds in solution, cannot be accepted as a complete theory; nor, on the other hand, can the hydrate theory of solutions be accepted as satisfactory unless it can be shown to be consistent with the two main characteristics of weak solutions—the apparent quasi-gaseous independence of the dissolved substance in dilute solutions, and the approximately accurate results which are obtained when calculations are based on the assumption that electrolytes in dilute solutions are dissociated into their ions. However important the service rendered to science by the gaseous and dissociation theories of solution, there are many fundamental objections which militate against their universal acceptation. They offer no satisfactory explanation of why substances dissolve at all, and still less why water alone, of all ordinary solvents, should resolve salts, &c. into their ions. These ions are represented to be atoms charged with enormous electric charges, but no explanation is given of the origin of these charges, or of the peculiarities which they are supposed to exhibit; two atoms oppositely electrified are represented as being less combined than when not charged at all; the charged atoms are represented as swimming about in the water without decomposing it, and without giving up their charges to their oppositely electrified companions, although they can part readily with them to an electrode; and to explain the thermal phenomena of solution, it has to be boldly assumed that the decomposition of molecules (elementary or composite) into charged atoms often evolves a considerable amount of heat, and consequently that the charging of an ordinary atom must be a process which evolves heat also. The only crucial experiment which has so far not been susceptible of an opposite interpretation seems to disprove any real dissociation of electrolytes. The lowering of the freezing-point of a solvent is accepted as affording a measure of the number of acting units (molecules or ions) in any substance added to it, and when, for instance, to some acetic acid a weak solution of sulphuric acid is added, the depression produced proves that the latter contains fewer acting units than the water and sulphuric acid do separately, instead of more, as

would be the case if the sulphuric acid molecules had to be split up into independent ions.

In a paper in the Berichte, Pickering has offered an explanation of the behaviour of dilute solutions based on the hydrate theory and on the views held by him as to residual affinity. The explanation may be rendered more intelligible by representing chemical attraction, as we represent electrical attraction, to be due to 'charges' on the surfaces of the attracting matter, but inalienable from the matter, owing to a repulsive force between the atoms similar to that which produces elasticity preventing the atoms ever coming close enough together to allow of the charges combining. The charges are represented as being always self-attracting. Each atom possesses one or more unit affinitycharge according as it is a monad or a polyad: with a free atom the charge would be uniformly distributed over its surface, whereas when two atoms are combined the greater part of the charges would be drawn to those sides of the atoms facing each other; but, as in the case of electric charges, a certain amount would still be left on the further sides capable of acting on, and being acted on by, other atoms or molecules in like condition, this residue, or unbound charge, forming what appears as residual affinity. As in the case of electrical charges, the nature of the matter composing different atoms will cause different degrees of mobility in the charges on their surfaces, and hence different pairs of atoms will be held together with different degrees of firmness, and different molecules will exhibit different amounts of residual affinity. When a molecule A B, possessing a certain amount of residual affinity, is surrounded by a number of other molecules CD possessing residual affinity also, the two residual charges will react on each other and increase the amount of the charges on the external portions of the molecules, leaving a smaller amount of charge on the portions of A and B which face each other; in the same way the charge on C will be partially withdrawn from the surface facing D, and will be able to retain a smaller amount of D's charge on the portion of D which is next to it, so that D will exhibit more residual affinity than formerly, and will be able to react in a similar manner on its neighbours, C' D'. Thus, by a process precisely analogous to electric induction, the number of molecules of C D which are acted on by, and are more or less combined with, A B may be very large, and far greater than the number which are capable of coming into its immediate vicinity. The existence of the very complex hydrates which have been recognised in dilute solutions is thus easily explicable. It is evident that a molecule when thus surrounded by, and combined with, a large number of solvent molecules will be attracted equally in every direction, and as the magnitude of the force exerted on it by each individual solvent molecule must be very small, it will be able to move easily in any direction, especially as any of the solvent molecules from which it parts during the process will have their places supplied at once by other similar molecules, of which, owing to continuous dissociation and recombination, there is an abundant supply in the liquid. Thus, into whatever new position the

sentral molecule moves, the status quo ante is immediately reproduced, and, consequently, the molecule will move almost unfettered by the presence of the solvent, and will act almost as if it were in the gaseous state. We thus obtain at once an explanation of why the application of the ordinary gaseous laws to dilute solutions yields very nearly true results; we should, indeed, expect that they would not be absolutely true, for certain definite numbers of solvent molecules would be capable of a more symmetrical arrangement about the central molecule than others, and such arrangements (definite hydrates) would be more stable than others, and the tendency to form them might have an appreciable effect on the regularity of the results: small irregularities of this sort are exactly what we find in the case of those dilute solutions which have been fully examined.

It is obvious that as the quasi-gaseous freedom of the molecule is dependent on its being combined with the solvent, this freedom obtains only so long as the molecule remains within the solvent, and would not admit of the possibility of its leaving the liquid, or of exerting an external gaseous pressure; for the same reason it would be impossible for it to pass through any crevice so small as not to allow it to retain as many solvent molecules as it was combined with in the solution. This gives us an explanation of the action of semipermeable diaphragms and all the phenomena of osmotic pressure; it, moreover, removes a difficulty which must be experienced in accepting the explana-tion given by the supporters of the physical theory of the action of these diaphragms-why it is that the water molecules (which are admittedly combined in great part into complex aggregates) can so easily pass through interstices which are impervious to what are held to be the single free molecules and ions of the dissolved substance. Direct experiment would seem to decide conclusively in favour of the explanation of osmotic pressure given above, for, when a suitable solution in a porous pot is immersed in either of the substances which compose the solution, osmosis through the pot towards the solution occurs in both cases, showing that osmotic pressure is not due to the impermeability of the pot to either of the constituents of the solution, but to its impermeability to the solution as a whole.

In a case where the atoms of the dissolved molecule possessed a considerable amount of residual affinity, the action and reaction between them and the solvent molecules would, in every respect, be proportionally great, and, consequently, those portions of the whole charges which are utilised in uniting the component atoms of the dissolved substance would be reduced, and, in extreme cases, might be reduced to the lowest possible limits—that is, till the amount of affinity utilised in holding them together was equal to that utilised in attaching them to the solvent molecules, in which case there would be an equal distribution of the affinity charges over the atoms. The atoms composing a molecule in such a condition would have a great amount of freedom of motion within the molecule; being attracted equally in all directions, they would be at liberty to start to

move in any direction, and their movements might attain considerable amplitude under suitable conditions as to the rate at which the forces between them varied with the distance: for to explain the apparent independence of the atoms, and therefore the abnormal osmotic pressure of a salt molecule, 'it only requires,' according to Fitzgerald, 'the space within which [the atoms] are bombarding about to be small compared with the space rate of variation of the force between [them].' The atoms of the two molecules in such a condition would obviously change partners very readily during collision, and this gives us an explanation of why such an interchange does take place when two different salts, &c., are mixed. This ready exchange, moreover, would take place continuously, and in a definite direction, under the influence of any external force, such as a charged electrode, a Grothüss' chain being formed, and we should consequently expect to find that substances in this condition—i.e. those which give abnormally large osmotic pressures—were electrolytes; this is so: and since, moreover, the amount of electrolysis occurring in a given time under a given force would be proportional to the number of molecules in the condition postulated, it follows that this number-and, therefore, the osmotic pressure—could be calculated, at any rate approximately, from the conductivity. In short, whatever calculations hold good on the assumption that the atoms in the molecules are absolutely independent will also hold good if they are assumed to be in the peculiar condition of combination here described—a condition which is brought about by their great tendency to combine with the solvent, and not by a tendency to part company for no assignable reason. On the present view also it is possible to understand why it is that salts and acids are generally electrolytes, and why water is the solvent which makes them electrolytes, for both salts and water are composed of elements which have a strong attraction for elements of an opposite character, and in such a case the residual affinity of both the elements (basylous and chlorous) constituting the salt would be acted on by that of the two opposite elements constituting the water, and we would get an amount of mutual reaction which would be impossible in the case of other substances.

The memoirs referred to in this article are: Berthelot, Méc. Chim. 1, 394; 2, 142; Thomsen, Thermochem. Untersuch. 3, 1-216; Mendeléeff, The Investigation of Aqueous Solutions according to their Specific Gravities (1887); also The Principles of Chemistry; and C. J. 52, 779; Roozeboom, Z. P. C. 10, 477; Féry, C. R. 115, 1809; Van der Waals, Z. P. C. 8, 214; Perkin, C. J. 63, 70; Jones, Z. P. C. 11, 112, 536; 12, 623; Pickering, C. J. 53, 865; 55, 14; 57, 16, 881; 63, 99, 141, 436, 890, 998; P. M. [5] 29, 427; 30, 400; 82, 20, 90, 478; 33, 132, 436; 34, 35; 35, 127; 36, 111; Z. P. C. 6, 10; B. 24, 277, 1579, 3817, 3826, 3629; 25, 1099, 1814, 1589, 1854, 2011, 2518, 3484; 26, 277, 121, 1977, 2307, 2766; 27, 80, 67; C. N. 57, 116; 63, 290, 305; 64, 1, 311. S. U. P.

SONNENSCHEIN'S REAGENT. Prepared by adding phosphoric soid to a warm solution of ammonium molybdate in nitric soid, boiling the pp. with squa regis to destroy NH, evaporating to dryness, and dissolving in 10 p.c. nitric acid (A. 104, 45). This reagent gives yellow pps. with alkaloids.

SOPHORIN. S. 009 in the cold; 5 at 100°. S. (alcohol) 3 in the cold; 7 at 78°. Obtained from Chinese yellow berries, the undeveloped flower buds of Sophora japonica (Stein, J. pr. 58, 399; 85, 351; 88, 280; Spiess a. Sostmann, J. 1865, 587; Förster, B. 15, 216). Decomposed by boiling dilute H.SO₄ into isodulcite and yellow sophoretin, which resembles quercetin.

80PHORINE. An alkaloid obtained from the pods of Sophora speciosa (Wood, Ph. [3] 8, 1047). Liquid, forming a crystalline hydrochloride, which gives a deep-red colour with FeCl₃. Poisonous. A poisonous alkaloid may also be obtained from the seeds of Sophora to-

mentosa (Greshoff, B. 23, 3539).

SORBIC ACID C.H.O. i.e. CH₃.CH:CH.CH.CH.CO₂H. Mol. w. 112. [184·5°]. (228°). H.C. 730,000 (Ossipoff, J. R. 20, 650). Heat of neutralisation: Gal a. Werner, Bl. [2] 46, 802). Occurs in the juice of unripe mountain-ash berries (Hofmann, C. J. 12, 43; A. 110, 129). The juice is partially neutralised with milk of lime, decanted from acid calcium malate, and distilled after addition of H₂SO₄. Needles (from dilute alcohol), v. sol. alcohol and ether, m. sol. hot water, volatile with steam. Partially decomposed on distillation (Barringer a. Fittig, A. 161, 307). Has no odour. Br in CS, forms tetra-bromo-hexoic acid [179°] and di-bromo-hexenoic acid [90°-95° Fuming HBr forms di-bromo-hexoic acid [68°] (Stahl, B. 9, 120). Very dilute alkaline KMnO. at 4° gives aldehyde, racemic acid, oxalic acid, and CO_2 (Doebner, B.~23,~2376). Sodium-amalgam forms hydrosorbic acid.

Salts.-BaA'2. - CaA'2: silvery scales. -

AgA': white crystalline pp.

Ethyl ether EtA'. (195.5°). Liquid. Chloride C_eH,OCl. Converted by NH₃ and aniline into crystalline C_eH₇O.NH₂ and C_eH₇O.NHPh respectively.

Hydrosorbic acid v. Hexenoic acid.

Isosorbic acid v. Hexenoic acid.

Reference.—Oxy-sorbic acid.

SORBINOSE C₆H₁₂O₆ i.e. CH₂(OH).(CH.OH), CO.CH₂OH (?). S.G. ¹⁶ 1·654. $[a]_j = -47^\circ$ at 7° . S. 200. An unfermentable sugar obtained from the fermented juice of mountain ash berries (Pelouze, A. Ch. [3] 35, 292; Byschl, J. 1854, 664; Delfis, C. N. 24, 75). The juice is left to stand for a year, and then filtered and evaporated. Trimetric crystals, as sweet as cane sugar. Lævorotatory. V. e. sol. water, sl. sol. hot alcohol. Forms with NaCl a compound crystallising in cubes. Its aqueous solution dissolves CaO, PbO, and Cu(OH)2. Not attacked by Br. Yields tri-oxy-glutaric acid on oxidation by HNO, (Kiliani a. Scheibler, B. 21, 3276), but no mucic or saccharic acid (Tollens, A. 249, 222). HIAq and P yield hexyl iodide. Reduced by sodium-amalgam to sorbite (Vincent a. Delachanal, C. R. 111, 52). Not turned brown by hot alkalis. Reduces Fehling's solution. Bromine water yields glycollic acid (Hlasiwetz a. Habermann, A. 155, 120). On heating sorbinose (1 pt.) on the water-bath with a solution of phenyl-hydrazine hydrochloride (2 pts.) and NaŌAo (8 pts.) in water (10 pts.) it | Vol. IV.

yields the osazone $C_{1s}H_{2z}N_4O_4$ [164°], which crystallises in yellow needles, sl. sol. hot water, v. sol. alcohol, v. sl. sol. ether (Fischer, B. 17, 582; 20, 827).

SORBITANNIC ACID. Obtained from the berries of Sorbus aucuparia (Vincent a. Delachanal, Bl. [2] 47, 492). Gives a yellow colour with alkalis; reduces silver salts in the cold; is not ppd. by alum or gelatin. Potash-fusion forms protocatechuic acid and phloroglucin.

SORBITE $C_0H_{14}O_6$. Mol. w. 182. [111°]. $[\alpha]_D = -1.73^\circ$. Occurs in mountain-ash berries (Boussingault, A. Ch. [4] 26, 376); in plums, cherries, and the fruits of nearly all the Rosacea Vincent a. Delachanal, C. R. 108, 147; 109, 678; 111, 52). Formed on reduction of sorbin and of l-gulose (Fischer a. Stahel, B. 24, 2144). Formed also, as well as mannite, by reducing glucose (dextrose) with sodium-amalgam in the cold (E. Fischer, B. 23, 3684; Meunier, C. R. 111, 49). Pearly crystals (containing aq). Melts at 75° when hydrated. V. sol. water and hot alcohol. Neither ferments with yeast nor reduces Fehling's solution. Lævorotatory, becomes dextrorotatory on addition of sodium borate (Vincent a. Delachanal, C. R. 108, 354). HIAq yields (β)-hexyl iodide (Hitzemann a. Tollens, B. 22, 1048). Boiling with Ac₂O and some ZnCl₂ forms a hexa-acetyl derivative. Benzoic aldehyde and a little hydrogen chloride give $C_8H_{12}O_5(C_7H_8O)$, crystallising in rectangular prisms [c. 175°]; if more acid is present the insoluble compound $C_8H_{12}O_4(C_7H_8O)$, [162°] is formed (Meunier, C. R. 108, 148; 110, 577; A. Ch. [6] 22, 423). In like manner valerio aldehyde and HClAq form, on shaking, $C_6H_{12}O_4(C_5H_9O)_2$, crystallising in prisms, sol. alcohol and ether. Sorbite may be exidised to glucose by heating in sealed tubes with brominewater at 60° and treating the product with Pb

and PbO (Vincent a. Delachanal, C. R. 111, 53).

SORDIDIN C₁₂H₁₂O₂. [180°]. A neutral crystallisable substance occurring in the lichen Zeora sordida (Paterno, J. 1875, 863). Insol.

water, v. sol. alcohol and ether.

SPARTEINE C₁₁H₂₂N₃. Mol. w. 234. (311° at 723 mm.) (Bamberger, A. 235, 868). [a]_B = -14·6° at 26° (Bernheimer, G. 13, 451). Occurs in the common broom, Spartium scoparium (Stenhouse, A. 78, 1; Mills, C. J. 15, 1). Prepared by extracting the plant with water acidulated with H₂SO₃, evaporating, and distilling with NaOHAq. The distillate is acidified by HCl; evaporated to dryness; and the residue distilled with solid KOH. The base is finally rectified over sodium. Heavy oil, with peculiar odour and bitter taste, sol. alcohol and ether, insol. benzene and ligroin (Houdé, Fr. 25, 568). Lævorotatory. Turns brown in air. Alkaline in reaction. Narcotic poison.

Reactions.—1. Yields (γ)-methyl-pyridine on distillation with lime (Ahrens, B. 21, 826).—2 HIAq at 200° yields MeI and C₁₁H₂₄N₂ (276°), which forms a platinochloride [230°], an aurochloride [157°], and a nitrosamine.—3. On passing the vapour through a red-hot tube it gives pyridine, (γ)-methyl-pyridine, C₂H₄, propylene, NH₃, HCy, and other bodies.—4. H₂O₃ forms a

base C_{l.}H_{.s}N_{.s}O_{.r}
Salts. - The hydrochloride is amorphous. - B'H.PtCl, 2aq. Decomposed by boiling K K

water (Oechsner De Coninck, Bl. [2] 45, 131).— B"2HAuCl. Decomposed by boiling water.-B"HI: tables, m. sol. cold water.—B"H, ZnI... B"I_s: green needles (from alcohol), insol. cold water.—B"H,804: prisms.—B"2C,H,N,0,.

Methylo-iodide B"MeI. Trimetric plates;

a:b:c = .899:1:1.601.

Ethylo-iodide B"Etl. Prisms (from water), v. e. sol. water and alcohol, insol. NaOHAq.—B"EtIHI. Got by heating sparteine with Etl and alcohol at 100°. Prisms (from cold alcohol). Not coloured by alcoholic potash (De Coninck, C. R. 104, 513). Yields (De Coninck, C. R. 104, 513). B"EtClHClPtCl.

Dihydride C₁₆H₂₈N₃. (283°). Got by reducing sparteine with tin and HCl (Ahrens, B. 20, 2219). Colourless oil.—B"HCl: very deliquescent needles.-B"H_PtCl. Blackens at 239°.-B"HAuCl. Blackens at 180°.-Picrate.

[125°]. Needles, decomposing at 215°.—B"HgCl₂.

Oxysparteine C₁₅H₂₄N₂O. [84°]. Formed by oxidising sparteine, and extracting the solution with chloroform (Ahrens, B. 24, 1095; 25, 3607). White hygroscopic needles, v. sol. water, alcohol, and ether. Its solution is strongly alkaline. Reduces hot Fehling's solution.—B"H,Cl, 4aq. Needles, v. sol. water. — B'HClaq. — B", H. PtCl, 4aq. Decomposed at 209°. — B"H. PtCl, aq. Needles: decomposed at 223°. — B"HAuCl, — Mercury double chloride [58°]. — B'HBr 2; aq. — B'HBr 3 aq. -B'HI aq. -B'HNO, aq. - Picrate [178°].

Methyl-iodide Methyl-iodide of oxysparterne B'MeI. [193].—B'MeCl.—B'MeClHClPtCl, aq. [229°]. POCl. at 150° converts oxysparteïne into an oily base C18H22N2 which yields C18H22N22HAuCl4

Di-oxy-sparteine C₁₈H₂₆N₂O₂. [129°]. Formed by dissolving sparteine in aqueous H2O2 (Ahrens, B. 20, 2220; 25, 8609). Prisms, v. e. sol. water and alcohol, sl. sol. benzene, insol. ether. Alkaline in reaction.—B"H_PtCl_{*}. Blackens about 235°.—B"HAuCl_{*}. [c. 146°]. Needles.—B"HI. [137°].—B'HBr. [147°].—B'H,HgCl_{*}. [193°]. Tri-oxy-sparteine Cl₅H₂₄N₄O₅. Formed by

action of H₂O₂ on oxy-sparteine. Crystalline, v. sol. water and alcohol.—B'2H2PtCl 3 aq.

B'HAuCl, [137°]

SPECIFIC GRAVITY v. DENSITIES, BELATIVE,

vol. ii. p. 870.

VOLUMES. SPECIFIC The molecular weights of substances when divided by their respective specific gravities, taken under comparable conditions, furnish values which have been termed atomic, molecular, or specific volumes. The specific volume is, therefore, the space occupied by aggregates of atoms, including the interstitial spaces, whose weights are proportional to the molecular weights of the anbstances.

Objection has been raised against the term 'specific volume' on the ground that, as specific gravity is the weight of unit volume, specific volume should by analogy be the volume of unitweight; hence of late years the term 'molecular volume' has been preferred. The term 'specific volume ' has, however, acquired by definition and use a distinctive meaning. The adoption of the term 'molecular volume' is almost certain to be attended with confusion, owing to the different sense in which it is frequently employed by physicists and chemists. The specific gravity of solids and liquids, referred to water at 4°. is the weight in grams of the unit volume. If, then, the molecular weight be expressed in grams, we may define the specific volume of a solid or liquid substance as the number of cubic centi-

metres occupied by this mass.

The specific gravity of a gas is usually referred to hydrogen as unity. It follows, then, from the law of Avogadro that all gases should have the same specific volume. There are a few exceptions to the universal truth of this statementarising from so-called abnormal vapour densities, but the specific volumes of the exceptional substances stand in the same simple relation to the normal value that their molecular weights stand to what analogy would indicate as the normal molecular weight.

Inasmuch as the specific gravity of a substance is affected to a greater or less extent by heat, it is obviously necessary to determine this constant under comparable conditions of temperature—that is, at temperatures at which heat may be supposed to exercise the same effect on the substance. In the case of liquids Schröder suggested that temperatures at which the vapour pressures of the various liquids are the same—as, for example, their boiling-points under a standard atmosphere-should be regarded as comparable.

In the case of solids the effect of temperature is of course much less marked, and hence the specific gravity of these substances is usually taken at the ordinary temperature of the air.

Specific volumes of solids.

The first attempt to trace relationships between the chemical nature of substances and their equivalent volumes appears to have been made by Le Royer and Dumas in 1821 (J. Ph. 92, 408). They sought to determine the equivalent volumes of the elements by dividing their atomic weights by their respective specific gravities, the values so obtained being termed by them atomic volumes. They were led to infer that these volumes formed an arithmetical series a supposition which was hardly warranted by the facts then known, and which has since been completely disproved by more accurate observations. This idea of combination among solids in definite volumetric proportion was further developed in 1824 by W. Herapath (P. M., Nov. 1824), who sought to prove that the volume of the oxygen in a metallic oxide bears a simple ratio to that of the metal with which it is combined. Almost simultaneously the same problem was attacked by Karsten (S. 65, 894), and subsequently, in 1830, by Boullay, but with no definite general result.

Ammermüller, however, in 1840, concluded (P. 49, 841; 50, 406) that the specific volumes of compounds containing the same elements in different proportions are either identical or stand to one another in rational proportions. in his 'Introduction to the Study of Molecular Chemistry,' recognised that equivalent amounts of many bodies of analogous composition occupy the same volume, and he inferred that the specific volumes of all substances are multiples of one and the same number, a conclusion also drawn by Le Royer and Dumas, but which is not supported by facts.

Kopp (P. 47, 138), in 1889, first attempted to

dedace general formulæ from which he was able to calculate the specific gravities of certain oxides and haloid salts, with results which showed, in general, a fair agreement with the observed values. In the same manner he calculated formulæ for other anhydrous salts-such as sulphates, carbonates, and nitrates-on the supposition that such salts consist of combinations of oxides and acids, or that they are made up of a radicle, acid plus oxygen. By means of these formulæ he inferred that it is possible to draw conclusions concerning the specific gravity of metals for which this constant is unknown. Kopp, in this memoir, used the term specific volume for the first time, and he defined it as the molecular weight (Mischungsgewicht) of a substance divided by its specific gravity. He found that the specific volumes of similarly reactive elements—as, for example, chlorine, bromine, and iodine; tungsten, molybdenum, chromium, iron, manganese, nickel, cobalt, &c.—are equal or nearly equal. In other cases—as silver and gold, potassium and sodium-the specific volumes stand to each other in simple relations. Elements which, like barium and strontium, form isomorphous compounds have the same specific volume. Strictly speaking, this law can hold only for those substances which are perperfectly isomorphous. The more nearly the crystalline forms of isomorphous substances are identical, the more nearly will their specific volumes be the same. This is made evident by a comparison of the axial ratios of witherite, strontianite, arragonite, and cerussite; and also of the carbonates of zinc and magnesium (mesitene), the carbonates of iron and manganese, and dolomite and calc-spar. It is seen that there is a direct comparison in the case of the latter compounds between the length of the principal axis a and the specific volume V, such that a 4.789 =0.0127671V, from which it is possible, of course, to deduce the specific gravity of the substance from its crystalline form. It also follows that an increase of specific volume is occasioned by an increase in the length of the axis a. If we heat one of these crystals the density decreases; the axis a must therefore increase in length, while the angle R becomes less obtuse. This fact, indeed, was discovered by Mitscherlich, who found that the specific gravity of calc-

spar decreased in the ratio of 1 to $\frac{1}{1-001961}$ when

calc-spar was heated through 100°. The specific gravity of calc-spar is 2.7220, when a = 0.8544 and $R = 105^{\circ} 5'$. By heating calc-spar through 100°, the specific gravity becomes 2.7167, or the specific volume changes from 36.73 to 36.80. If we determine the length of the axis a by means of the above formula, we find it to be 0.85672, corresponding to an angle R of 104° 57', or a difference of 8', which closely agrees with that actually observed by Mitscherlich (P. M. [3] 18, 255). Schröder (P. 1840. 553), starting from the observation of Ammermüller, that equal volumes of the two oxides of copper contain the same amounts of copper and multiple amounts of oxygen, assumed that the volume of the copper, as of the oxygen, is equal in the two substances, but that the amount of the oxygen in the cuprous exide stands to that in the cupric oxide as I to 2. Hence Schröder drew the general conclusion that the same element can have different specific volumes in different compounds, but that the several values for the specific volumes stand in simple relations to each other. He saw in this hypothesis not only an explanation of the condensation which accompanies chemical union, but also a rational basis for the belief that the specific volume of a compound is equal to the sum of the specific volumes of its components.

Schröder found that if in a series of analogous bodies, AO, BO, CO, the specific volumes of which are known, we subtract from these values the primitive atomic volumes of A, B, and C respectively, we obtain a constant remainder. This he found to be the case with the oxides of lead, cadmium, and zinc, and hence he inferred that the metal in these oxides retains its primitive atomic volume. Kopp assumed that this is equally true of the salts of the heavy metals, but with the salts of the metals of the alkalis and alkaline earths this is impossible, as the specific volumes of the salts are, as a rule, smaller than the primitive atomic volumes of the component metals. He had consequently to assume for these metals a special atomic volume, which however, remains the same in all the salts. He determined these values as follows: Suppose M+R to be a compound of a heavy metal, and m+R the analogous compound of a light one; suppose A to be the known specific volume of M+R, and a that of m+R, B the primitive atomic volume of M, and b that of m.

Then, M + R = A, and M = B.

Therefore the atomic volume with which **R** is contained in the compound is A-B, say x.

It is assumed that R retains its value in m+R.

and since m + R = a, and R = x;

therefore b, i.e. the atomic volume of m = a - x. Kopp also showed that the densities of the hydrated oxides, and of a number of hydrated salts, may be calculated with considerable accuracy by assuming certain definite values for water in a state of combination. It ought to be stated, however, in this connexion, that subsequent researches have indicated that Kopp's conclusions respecting the specific volume of water of crystallisation must be slightly modified. Schiff, many years ago, showed that the members of certain classes of hydrated salts have practically the same specific volume. Thus, all the alums have a specific volume of about 277; double sulphates of the form M2M"(SO4)2.6H,O have a common volume of 207; and all the vitriols—that is, salts of the form M"SO,.7H,Owhether isomorphous or not, have the specific volume 146.

Thorpe a. Watts (C. J. 37, 102, [1880]) have shown that the volumes occupied by the several molecules of water vary with the degree of hydration of the salt. In the case of the so-called magnesian sulphates, the first molecule of water, the constitutional water, or 'water of halhydration' of Graham, occupies considerably less bulk than the remaining molecules; its mean relative value is 10.7. Each additional molecule appears to occupy a gradually increasing volume. The difference between the specific volumes of

the monohydrate and dihydrate is 13·3; between the volumes of the dihydrate and trihydrate the difference is 14·5; between the trihydrate and the tetrahydrate it is 15·4; and between the hexhydrate and heptabydrate it is 16·2. These observations are so far in harmony with Kopp's general conclusions that in the compounds containing only a small number of water molecules (1 to 3) the specific volume of the water is 12·4; in others containing a larger number of molecules of water (2 to 7) it is 13·4; whereas, in a third class, containing the largest number of molecules of water (from 3 to 10), its mean value is 15·3.

Specific volumes of liquids.

Methods. The specific gravity of the liquid at its boiling point may be determined either directly or indirectly. Ramsay (C. J. 35, 463 [1879]) devised a simple method by which the weight of a known volume of liquid at its boiling-point may be directly ascertained with approximate accuracy. The vessel containing the liquid consists of a thin glass lemon-shaped bulb of about 10 c.c. capacity. At the upper end of the bulb is a capillary tube, bent into the form of a hook; the lower end is provided with a similar hook of solid glass. The capacity of the bulb is ascertained by weighing it full of water at a known temperature. The bulb is filled with the liquid to be examined; in the usual way, by warming and dipping the capillary neck of the bulb beneath the surface of the liquid. It is not necessary to fill the bulb completely; two or three c.c. of the same liquid are then introduced into the wide tube, and the bulb is suspended within it by thin platinum wire attached to a piece of glass rod passing through a hole in the cork. The wide tube is then heated until the liquid within it boils violently, and it is kept in ebullition so long as liquid drops from the end of the capillary neck of the bulb. The lamp is now removed, and the whole allowed to cool. When cold the bulb is dried and weighed. The specific gravity of the liquid is given by the formula

Sp. gr. =
$$\frac{W'}{1 + (0.00015 \times TW)}$$

in which W' is the weight of the liquid; W, that of the water filling the bulb at 0°; and T the temperature at which the liquid boils; 0.00015 is an empirically-determined co-efficient required in order to make the volume correspond with the real volume at T.

A somewhat similar method of determining the specific gravity of a liquid at its boiling-point is described by R. Schiff (A. 220, 78). The liquid is introduced into a flask of about 7 or 8 c.c. capacity, provided with a long narrow neck on which is a graduated scale. The capacity of the flask up to the zero point is accurately determined by weighing with mercury, and the value of the scale divisions in fractions of a c.c. is also carefully estimated. The flask is then suspended in the boiling tube; a few c.c. of the liquid under investigation are placed in this tube and heated to boiling. After a few minutes the position of the liquid in the graduated neck is noted, and the flask is with-drawn, wiped, and weighed. The specific gravity

at the boiling-point compared with water at 4° is given by the equation

$$D_4 = \frac{P}{V_t[1 + K(t-4)]}$$

in which P = corrected weight of the liquid in the flask,

and $\nabla_t =$ the apparent volume of the liquid at t° ,

K = the co-efficient of expansion of the glass.

It will be noticed that Schiff assumes that the liquid in the flask actually acquires the temperature of the vapour by which the flask is surrounded. Lothar Meyer (v. Neubeck, Z.P.C.1, 652) has slightly modified Ramsay's apparatus. The bulb is best made of Jena glass, and has the capacity of about 2.5 c.c., and its weight, capacity, and thermal expansion are accurately known. It is filled with the liquid, and is suspended by means of a bent platinised-nickel wire within the flask, the neck of which can be closed by a cork. The flask contains a few c.c. of the liquid under investigation, which, when heated, may if necessary be caused to boil at any desired temperature below the ordinary boiling-point by connecting a side tube with a condenser, pressure-regulator, and fall-pump. As soon as the liquid within the bulb has acquired the temperature of the boiling liquid, and no further expansion occurs, the source of heat is removed and the bulb is weighed when cold.

The specific gravity of a liquid at its boilingpoint is, however, most accurately ascertained
indirectly, although this method demands far
more time and apparatus, and a greater number
of experimental operations, than the direct
method. The method consists in accurately
determining (1) the boiling-point of the liquid;
(2) the specific gravity of the liquid at some
convenient temperature—best at 0°; and (3) the
rate of thermal expansion of the liquid from 0°
up to the neighbourhood of its boiling-point.
From these data the specific gravity of the liquid
at the boiling-point can be readily calculated.

1. Determination of the boiling-point. This is best made in an apparatus so arranged that the inner tube, containing the thermometer, is surrounded by a jacket of vapour from the boiling liquid. The boiling liquid should contain a few scraps of platinum foil or a spiral of platinum wire; or, what is even better, a few short lengths of fine capillary tubing. If the amount of the liquid is very small the bulb of the thermometer should be surrounded with a little fibrous asbestos, as suggested by Ramsay a. Young (C. J. 47, 42). The thermometer reading must, if necessary, be corrected for the emergent column, either by the tables of Rimbach (B. 22, 3072) or by that of Thorpe (C. J. 87, 159), and should be reduced to normal pressure by the method of Crafts (B. 20, 709). The correction to standard atmosphere may, however, be ascertained with approximate accuracy by the formula

$$\theta = (p - 760) \frac{273 + t}{8200}, \text{ or } \theta = (p - 760) \frac{273 + t}{10000},$$

in which θ is the correction, t the observed boiling-point, and p the barometric pressure reduced and corrected.

The first correction is applicable to the

greater number of liquids, the second to water and the lower alcohols (Ramsay a. Young, P. M. 1885, 515).

- 2. Determination of specific gravity. This is most accurately effected, at least for substances which are liquid at ordinary temperatures, by Perkin's modification of the Sprengel apparatus. For liquids which are alterable on exposure to air a bottle with a graduated stem is most convenient. The bottle should be previously filled with dry nitrogen, and a special apparatus, analogous to that used by Thorpe (l.c.) must be employed to transfer the liquid to the bottle. A form of bottle for very viscid liquids has been described by Brühl.
- 3. Determination of thermal expansion. the various modes of ascertaining the thermal expansion of a liquid the so-called dilatometrical method is, on the whole, the most convenient. It consists in inclosing the liquid in a vessel of known capacity, shaped like a thermometer, and provided with a graduated and accurately calibrated stem. The instrument is placed in a bath of suitable liquid, the temperature of which can be raised to the desired point, as ascertained by a thermometer; and the height of the liquid in the stem of the dilatometer, and hence its apparent volume at the temperature of observation is noted. A series of such observations at various temperatures up to the neighbourhood of the boiling-point is thus made, from which an expression, say of the form $V_t = a + bt + ct^2 + dt^3 \dots$ may be calculated. This expression must now be corrected for the expansion of the glass of the dilatometer (obtained by observations with mercury in the usual manner), and from the corrected expression the volume, and hence the specific gravity, at the boiling-point of the liquid may be deduced.

For details of the mode of carrying out these observations v. Kopp (A. 96, 1 et seq.); Thorpe (C. J. 37, 141).

Thorpe (C. J. 63, 262 [1893]) has devised a modification of the ordinary dilatometrical method, which permits the thermal expansion of a liquid to be determined with much greater ease and rapidity than has hitherto been possible. It obviates the use of large baths, and, by reducing the size of the dilatometers to the smallest limit consistent with the proper degree of accuracy, as determined by the error of a thermometer reading, it avoids the necessity for long stems and the consequent correction for the cooled 'emergent columns.'

Results.—The first accurate determinations of the various physical constants needed to ascertain the specific volumes of liquid substances were published by Kopp in 1855 $[A.\ 96\ [1855]\ 1_36, 153_185, 303_335; A.\ 100\ [1856]\ 19_38; v.$ also $P.\ 72\ [1847]\ 1_62, 176, 223_293).$ Kopp concluded, from the results of a long series of observations, that:—

(1) The selection of the temperature of equal vapour-pressures as a basis of comparison seemed to be warranted by the fact that regularities are thereby made evident which otherwise are not apparent.

(2) Differences of specific volume are proportional to differences in chemical composition.

(3) Isomeric liquids of the same chemical type have equal specific volumes.

(4) The substitution of hydrogen for an equivalent amount of oxygen only slightly affects the specific volume.

(5) One atom of carbon can replace two atoms of hydrogen without altering the specific volume of the compounds.

Kopp further found that the specific volume of a liquid compound was conditioned, not only by the composition, but also by the constitution of the compound. Thus the relative position of an oxygen atom in a molecule affects its specific volume; carbonylic oxygen and hydroxylic oxygen have two very different values. Sulphur, in like manner, would appear to have two specific volumes, depending on its position or mode of combination in a molecule.

Definite values for the specific volumes of carbon, hydrogen, and oxygen were obtained from the following considerations. An increment of CH₂ corresponds to an increase of specific volume of 22. Since C and H₂ occupy the same volume, we have C = 11 and $H = 5 \cdot 5$. The replacement of H₂ by carbonylic O is attended by a slight increase in the specific volume. Kopp found that the most probable value for O in this form of combination was 12 \cdot 2. For hydroxylic oxygen it is 7 · 8; obtained by subtracting the value for H₂ (11) from the specific volume of water, $18 \cdot 8$. Hence the specific volume of a compound $C_a H_b O_c O'_d$, where O is carbonylic oxygen and O' is hydroxylic oxygen, may be expressed by the formula

V = 11.0a + 5.5b + 12.2c + 7.8d.Determinate values for the specific volumes of the halogens-and, with less precision, for phosphorus, arsenic, and antimony, for silicon, titanium, and tin-were also obtained by Kopp. As regards nitrogen it was found that in the amines $N=2\cdot3$. The group CN=28; the group NO₂=33. If, then, carbon and oxygen preserved their ordinary values in these radicles N must possess at least three different values depending on the mode of combination. There is. however, no evidence to disprove the supposition that the values for the carbon and oxygen atoms are not equally affected in these groups. This, indeed, suggests the possibility that compound radicles like CO, HO, NO, CN, &c. may possess definite specific volumes which are not necessarily the sum of the specific volumes of the component atoms as ordinarily ascertained.

H. L. Buff (A. Suppl. 4, 129) sought to show that the specific volume of carbon, like that of oxygen and sulphur, is affected by its mode of combination-or, in other words, that carbon in unsaturated compounds has a greater specific volume than in saturated bodies—from which he surmised that the specific volume of an element is in general determined by its particular atomic value. Thorpe (C. J. 37, 392) found that isomeric liquids have not invariably the same specific volumes. There is a well-marked difference, for example, between ethylene and ethidene chlorides; indeed, ethylene compounds in general appear to have smaller volumes than those calculated by Kopp's values. Städel has shown that in the series of chlorinated and brominated ethanes and ethylenes, the isomeride of higher boiling-point, i.e. the etaylene derivative, has invariably the lower specific volume. Since these compounds are all saturated, and the only variable constituent is a monovalent element (Cl or Br), it would appear probable that the specific volume of the halogen is also variable. Isomeric hydrocarbons manifest similar differences, whence it is obvious that either one or both of the elements must have a slightly variable volume, depending on grouping or mode of combination; it may be that the iso-group, like the groups carbonyl, hydroxyl, nitryl, &c., has a special volume, which is not necessarily the sum of the volumes of the component atoms as deduced from Kopp's values. Kopp himself found that the volumes of isomerides were in a number of cases only approximately equal, and in others quite unequal. The term 'chemical type,' used in the sense in which Gerhardt employed it, is not sufficiently distinctive to denote the differences, say, between the normal and iso-compounds, or between aniline and the picolines, and it is questionable whether Kopp would have considered such cases as coming within his rule.

The observed specific volumes of the aromatic compounds are frequently lower than the calculated values. Indeed, our views as to the constitution of the aromatic compounds would lead us to expect that the specific volume of benzene, and the volumes of the derivatives which contain the benzene grouping, would probably be different from the values deduced from observations made for the most part on compounds of totally different constitution. Kopp (A. Suppl. 5, 303 [1867]) showed from Louguinine's observations that while benzene has an abnormally low specific volume, its homologues show the constant increase of 22 for an increment of CH, which is what might be anticipated, since these homologues are produced by the addition (substitution) of methyl, ethyl, &c., to the benzene group. Jungfleisch's observations on the specific volumes of the chlorine substitution products of benzene also seem to show that the positions of the chlorine atoms affect, in a very marked manner, the specific volume of the product (C. R. 64, 911).

Further observation has shown that Kopp's conclusion that liquid elements and radicles have the same volume in combination as in the free state is well founded. Thus the observed volume of NO₂ = 32.0, calculated = 31.5; observed volume of Br = 53.6, calculated = 53.4; observed volume of CN = 28.9, calculated = 28.9. The observed specific volume of Cl from Knietsch's determinations of the specific gravity of liquid chlorine is 22.8; the mean calculated value is 22.7. Kopp surmised that members of the same chemical family would be found to have the same specific volume; observation shows, however, that the specific volumes gradually increase with the increase of atomic mass (Thorpe, l.c.).

Schiff (A. 220, 71 [1883]) has concluded that while it is generally true that isomeric compounds have slightly different specific volumes, it is almost invariably the case that the substance possessing the higher boiling-point has also the higher specific volume (compare Städel). In the case of the metameric esters of the fatty acids, it is found that, as a rule, the specific volumes increase with the diminution of the number of carbon atoms in the acidic radicle and with the increase of the carbon atoms in the

alcoholic radicle. At the same time, it would appear that the differences between the observed and calculated values are mainly due to the alcoholic radicle, the acidic radicle apparently having but slight influence. This is in conformity with Lossen's observations, that while the ethers and acids give experimental values which are almost in exact accordance with Kopp's values, the aldehydes and alcohols show wider variations, methyl alcohol giving too great an observed value, while the others give smaller and smaller values as the amount of carbon increases. It is, however, noteworthy that the differences between the aldehydes and derived alcohols remain almost constant, which is not the case with the aldehydes and acids, where the difference appears to increase with the molecular weight. Hence the differences between the homologous aldehydes are very nearly equal to those between the corresponding homologous alcohols (Lossen). The mode in which carbon is combined in an organic compound has, according to Schiff, a distinct influence on its specific volume; like Buff, he finds that a doubly linked carbon atom occupies a smaller volume than when singly linked. It is, however, very doubtful whether the facts at present known are sufficient to establish this conclusion.

There is, however, good reason to believe that what we call 'specific volume' is not a purely additive property. The specific volumes of substances are, in all probability, affected by many more conditions than those we have hitherto taken cognisance of. The value $CH_2 = 22$ has no other significance than as expressing the average increment in volume in successive members of a homologous series. Indeed, as the physical data increase it becomes doubtful whether even this mean value is correct. It would seem that the value augments as the series is ascended. The relation C = 2H no longer applies to carbon compounds in general. What is true of carbon and hydrogen is equally true of oxygen, whether as carbonylic or as hydroxylic oxygen. No definite or uniform values can be assigned to oxygensuch that the specific volume of a liquid compound containing this element can be accurately calculated. The values given by Kopp and others are simply mean values, but the actual volumes are affected by conditions of which, as yet, we have no very precise knowledge and which we have no certain means of measuring. The values for the other elements are, of course, affected by these considerations. Thus the specific volume of chlorine is obtained on the assumption that the values for carbon and hydrogen are constant.

Lossen (A. 254, 42) has devised formulæ which take note, or express the measure, of the influences which affect the uniformity in the values of specific volumes of organic compounds. These formulæ can only be considered as first approximations, but their value will be evident from the fact that they serve to reproduce the observed values with a greater approach to accuracy than has hitherto been possible. Out of the 407 compounds which furnished the experimental material on which these formulæ are based, the observed molecular volumes of 352 differ by less than 2 p.c. from the calculated volumes. Comparatively few of these differences are to be ascribed to experimental errors. In

the main they are caused by influences of structure and composition which as yet we have no certain means of measuring, such as the effect of substituted chlorine, or the special effect of iso-grouping, or of the ortho-, meta-, or paraposition, &c.

According to Lossen, the specific volumes of the greater number of compounds containing carbon, hydrogen, and oxygen which have hitherto been determined, may be calculated by the formula

Sp. vol.
$$C_n H_m O_p(\mu) = (10.24 \pm x0.5)(n+p) + (5.12 \pm x0.25m) \pm \frac{1}{4}(n-2)^2 \pm 1.4\mu$$

In this equation x denotes a number between 0 and 1, which varies with different homologous series but which is constant for the members of the same series. The sign μ represents the number of hydrogen atoms required to convert the formula into that of a saturated compound. The term $\frac{1}{4}(n-2)^2$ is introduced to compensate for the increase in the value corresponding to CH₂ as the molecular weight increases. Gartenmeister (A. 233, 804) having shown that in the case of the fatty esters the mean increase corresponding to CH₂ is 0.5, Lossen adds the term $\left(\frac{1}{2} + \frac{2}{3} + \frac{2}{3} + \cdots + \frac{n-3}{2}\right)$, which he erroneously states is equal to $\frac{1}{2}(n-2)^2$,

which he erroneously states is equal to $\frac{1}{2}(n-2)^2$, a number always subsequently employed, but which, as comparison shows, serves to give good agreement with observed values.

For the majority of fatty compounds the formula

 $C_nH_mO_p=10\cdot45(n+p)+5\cdot225m+\frac{1}{4}(n-2)^n+1\cdot5\mu$ gives results agreeing with the observed values. For the alcohols, however, the expression becomes

$$C_nH_mO_p=10\cdot 1(n+p)+5\cdot 05m+\frac{1}{2}(n-2)^n+1\cdot 85\mu$$
.

If it be supposed, as seems highly probable, that for all the members of a series of similarly constituted compounds the same influences are existent, whereby the specific volumes exhibit variations from the additive quantities assumed by Kopp, then we may regard the variable x as a term which takes account, and may be regarded as the measure, of these disturbing influences, and which, though constant for the members of the same series, should, as before stated, vary for other homologous series of compounds (Thorpe a. Jones, C. J. 63, 289).

Schröder regards the specific volume of an element in combination as variable within limits determined by the nature of the chemical compound. In any one compound, however, all the elementary atoms are regarded as occupying either equal or multiple volumes. Hence every specific volume is a multiple of a certain space-unit or stere, the value of which may vary between 6.7 and 7.4. depending on the number, nature, and mode of union of the atoms. Schröder thus obtains his determinate values. The specific volumes of formic, acetic, and propionic acids increase about 22 units for each increment of CH2; in the case of alcohols the increase is about 20 units. The volume of formic acid is 41.8, i.e. 2 × 20.9; that of methyl alcohol is 42.8 or 2×21.4 . Hence in formic acid, CH₂O₂, the O₃ occupies the same volume as CH₂; and in methyl alcohol, CH₂O, the HO has the same volume as CH. In like

manner ethyl alcohol, C_2H_2O , has the volume 62·1, i.e. 8×20 -7. Since $2\times CH_2=2\times 20$ -7, then $H_2O=20$ -7. Acetic acid has the volume 63·6, i.e. 8×21 -2; $2CH_2$ is 2×21 -2 and $O_2=21$ -2. The volume of acetaldehyde is 56-9, i.e. about 6·5 smaller than that of acetic acid; this would indicate that the substitution of OH by H lowers the volume by 6·6. Since $H_2O=21$ -4, it follows that the hydrogen and oxygen in hydroxyl each occupy one space-unit or stere. As $CH_2=21$, we find that carbon also occupies one stere or space-unit. Now, from the volume of formic acid which contains 6 steres, it is found on subtracting 3 steres for CH_2 and 1 stere for hydroxylic oxygen, that the carbonylic oxygen must occupy 2 steres.

We obtain, then, the following rule: The number of space-units or steres of the saturated compounds of carbon, hydrogen, and oxygen corresponds to the number of the atoms increased by as many units as there are atoms of carbonylic oxygen present.

If we calculate by means of this rule the steres of the saturated compounds, we find that they vary within narrow limits, and for the most partinorease with increasing molecular weight. In the greater number of cases the values range between 6.9 and 7.2. Ostwald has determined the value of the stere for a large number of saturated fatty compounds, with the following results:—

Hydrocarbons 6:89, 6:99, 6:82, 7:11, 7:23.

Alcohols 7.12, 6.91, 6.77, 6.88, 6.78, 6.78, 6.81, 6.74, 7.09

Acids • 6.97, 7.06, 7.11, 7.19, 7.24, 7.24, 6.85.

Esters • 7.04, 7.08, 7.05, 7.14, 7.26, 7.43, 7.45, 7.47.

Aldehydes . 7:11, 7:05, 7:18, 7:01, 7:18, 6:93,

In the series of the hydrocarbons, the acids, and the esters, the steres in the cases of the normal compounds increase regularly with increasing molecular weight; in the series of the alcohols the steres decrease up to the third member and then increase. The secondary and tertiary compounds have, as a rule, smaller steres than the normal compounds. Kopp concluded from the approximately equal volumes of the alcohols and corresponding acids that H2 and O are volumetrically equivalent; and from the equivalence of the volumes of benzyl and amyl compounds he inferred that C, and H, are mutually replaceable without alteration of volume. Hence he assumed that the hydrogen atom occupies only half the volume of the oxygen or carbon atom. Schröder established the volume equivalence of CH2, HOH, and O'O, and he inferred from the difference in volume between alcohol and aldehyde that hydroxylic oxygen has the same volume as hydrogen and carbon, while carbonylic oxygen has twice the volume. While, then, Kopp assumes approximately $H_2 = C = 0$, Schröder makes H = C = 0 (Ostwald, Lehrbuch, [2nd ed.] vol. i. p. 388).

In the case of unsaturated and aromatic compounds, Schröder assumes that each double linkage is attended with an increase of volume amounting to one stere; hence the above rule has to be modified in this sense when applied to compounds of this class. The value of the stere

in a number of unsaturated compounds is found to be:-

6.87, 7.09, 6.93, 6.99. Hydrocarbons6.72. Alcohols

Esters and ethers . 7.14, 7.13.

These values vary, practically, within the same limits as in the saturated compounds.

As regards aromatic compounds, it would seem to follow from the identity in the volumes of benzoyl and amyl compounds, as indicated by Kopp, as well as from the corresponding relation between the isobutyl and phenyl compounds, that C.H. comprises the same number of steres as C4H9—that is, 13. Of these, 5 are occupied by hydrogen; so that the carbon group C. occupies 8 steres.

The values of the stere in a number of aromatic compounds are as follows:

6.85, 6.94, 6.98, 7.00, 6.95, 7.04, Hydrocarbons 7.04, 7.04, 7.06, 6.84

Other compounds 6.91, 6.87, 7.05, 6.97, 7.16, 7:26, 7.50, 7.28, 7.14, 6.96, 7.07.

The value of the stere here also varies within the usual limits; it is comparatively small for the hydrocarbons (6.8-7.0), larger in the case of the alcohols, and still larger in that of the esters (7.2-7.5) (Ostwald, *l.c.*).

There is one consideration which is vital to the whole question, and to which, therefore, a brief reference must be made. It relates to the choice of conditions under which the values we term specific volumes are really comparable. Although Horstmann and Lossen have advanced reasons against the practice, contending that at any other temperature, say 0°, relations similar to those now established are made manifest, it has been the custom, in accordance with Kopp's direction, to compare the specific volumes of liquids at the temperatures of their respective boiling-points under a standard atmosphere. Whether, however, the temperature of the boiling-point, under these circumstances, is a truly comparable condition is open to question. It has been urged by Horstmann that, since what we call atomic volume is the space not merely filled by an atom but also that in which it moves, it is not a priori probable that at temperatures which differ, say by 300° —as, for example, in the case of C_4H_{10} (boiling-point 1°) and $C_{18}H_{28}$ (boiling-point 317°)—these volumes will be the same. Moreover, as pointed out by Bartoli, the boilingpoint cannot in the nature of things be a strictly comparable condition, since it is affected by pressure to a different extent in the case of different liquids. Objections of even greater weight may be urged against the suggestions of Tschermak and Krafft to take the melting-point as a comparable state.

No doubt, theoretically speaking, a valid condition should be when pressure, volume, and temperature are expressed in terms of their critical values. But that certain regularities in the molecular volumes at the boiling-points have, in spite of this, been discovered may be explained, as Guldberg has shown, when we compare the values of T, the absolute boiling-point, with those of T, the absolute critical temperature; in those cases in which these two perature; in shoot constants are known, the ratio $\frac{\mathbf{T}}{\mathbf{T}_i}$ approximates to $\frac{2}{8}$ Hence it follows that qualities like molecular volumes, which alter only slowly with temperature, are comparable at the ordinary

boiling-points (Z. P C. 5, 374).

It ought, perhaps, to be stated that subsequent observations show that the so-called 'corresponding temperatures' deduced from Van der Waal's generalisations have not that degree of validity as temperatures of comparison which they were originally assumed to possess. Indeed, the present condition of knowledge warrants the statement that Kopp's original method of comparison is as valuable as any yet indicated (Thorpe, C. J. 68, 775 [1893]). T. E. T.

SPECTROSCOPIC ANALYSIS v. PHYSICAL METHODS, section Optical methods, subsection

Spectroscopic methods, this vol. p. 239.

SPELTER. A commercial name for sinc.

SPERGULIN (C,H,O,)n. Occurs in the seedcoverings of Spergula vulgaris and S. maxima (Harz, C. C. 1879, 24). Amorphous. Its alcoholic solution exhibits dark-blue fluorescence, which is destroyed by sunshine. A small quantity of potash or Na CO, added to the alcoholic solution causes it to fluoresce green.

Conc. H2SO4 forms a dark-blue liquid.

SPERMINE $C_{10}H_{24}N_4$ (Poehl, B. 24, 359); $C_4H_{10}N_2$ (S.). Occurs as phosphate in the spermatic fluid, in calves' liver and heart, and in some pathologic preparations that have been kept under alcohol (Schreiner, A. 194, 68). It is not identical with pyrazine hexahydride (Majert a. Schmidt, B. 24, 241; cf. Ladenburg, B. 20, 442; 21, 758; Poehl, C. R. 115, 515; Duclaux, C. R. 115, 155, 549). Crystals (from alcohol), v. sol. water, v. sl. sol. alcohol. Alkaline in reaction. Absorbs CO₂ from the air. Its aqueous solution is ppd, by phosphomolybdic acid and by potassium bismuth iodide. — C₁₀H₂₈H₄HCl: prisms, v. e. sol. water.—C₁₀H₂₈N₄2H₂PtCl₂.—C₁₀H₂₈N₄4HAuCl₄.—C₁₀H₂₈N₄4HAuCl₄.—C₁₀H₂₈N₄4HAuCl₄.—C₁₀H₂₈N₄4HAuCl₄.—C₁₀H₂₈N₄AHAuCl₄.—C₁₀H₄AHAuCl₄.—C₁₀H₄ $C_{10}H_{26}N_4(H_1PO_4)_2$ 6aq: [170°]; rosettes of pyramids, sl. sol. hot water.

SPIKE 01L. S.G. above 900. Slightly dextrorotatory (Schimmel, Ph. [3] 22, 329). [a]_B = 1° 24' (Voiry a. Bouchardat, C. R. 106, 551) or lævorotatory (Bruylants, J. Ph. [4] 30, 139). An essential oil obtained from the blossoms of Lavandula aspica latifolia (Lallemand, A. 114, 197; Sauer a. Grünling, A. 208, 75). Smells like lavender. Contains a terpene (175°) or (158°), which yields crystalline C₁₀H₁₈HCl, camphor, borneol, and a resin. According to Voiry and Bouchardat, oil of spike contains inactive spikol C10H18O [0°] and C10H18 $(155^{\circ}-160^{\circ})$ [α]_D = 24°, which yields a hydrochloride [129°] [α]_D = -2°.

SPIROGRAPHIN v. PROTEÏDS, Appendix C. SPONGIN v. PROTEÏDS, Appendix C.

STACHYDRIN C,H₁₂NO₂. [210°]. Occurs with glutamine, tyrosine, and stachyose in the tubers of Stachys tuberifera (Von Planta a. Schulze, B. 26, 939). Colourless, deliquescent crystals (from water). Behaves like betains with regard to alkaloidal reagents.—B'HCl. Prisms, v. sol. water, sol. cold water (difference from betaine).—B'₂H₂PtCl₂ 2aq. Trimetric crystals; a:b:o=*608:1:*828.—B'HAuCl₄. Small yellow prisms (from water).

STACHYOSE $C_{18}H_{22}O_{16}$ 3aq. [a]_D = 148°. Occurs in the roots of Stachys tuberifera (Planta a. Schulze, B. 23, 1692; 24, 2705). Tablets, with sweetish taste, v. sol. water. Dextrorotatory. Has no action on Fehling's solution until after boiling with mineral acids, by which it is converted into galactose and an isomeride (? glucose). HNO₃ forms mucic acid, galactose, glucose, and cane-sugar. Gives a red colour on heating with resorcin and HClAq. Gives no pp. with lead acetate until NH₂Aq is added.

STANNATES, and salts related thereto; v.

STAPHISAGRINE v. DELPHININE.

STARCH. Amylum. $nC_{12}H_{20}O_{10}$. The value of n has not been definitely settled; it is undoubtedly high. Brown and Morris (C. J. 55, 462), employing Raoult's method for determining molecular weights, show n for soluble starch (see below) = 100, i.e. molecular weight = 32,400. The molecular weight of starch cannot be less than this. From O'Sullivan's work (C. J. 35, 783) it would seem that n is not less than 36. Pfeifer a. Tollens (B. C. 1882, 775; A. 210, 295), from the composition of some sodium and potassium compounds prepared from starch, arrive at the value n=2; the substances are probably compounds, not of starch, but of some decomposition products thereof. Sachsse (C. C. [3] 8, 732) and Nägeli (A. 173, 218) proposed $6C_8H_{10}O_4 + H_2O$ as the formula. Other observers (Mylius, B. 20, 694; Salomon, J. pr. [2] 28, 82) have suggested different formulæ, but the evidence is not satisfactory; we may take it, however, that the molecule of starch is not less than n=100.

Occurrence.—Starch is present at some time or the other in all green plants. It is said to be found in almost all parts of the plant, but it is specially stored up in seeds, the pith of stems, in bulbs, tubers, rhizomes, and roots—generally those parts of the plant which serve as a store for reserve material. It is, however, a question whether the granules recognised as starch in leaves, sap, &c., outside the reserve organs, are chemically identical with the starch of these organs. The evidence, one way or the other, is unsatisfactory. Starch is not known to be an animal product. It is present in some fungi (Bourquelot, J. Ph. [5] 24,197).

Formation.—Starch has not been prepared artificially. It is produced in the chlorophyll cells of plants, light, carbon dioxide, and water being necessary; oxygen is eliminated at the same time. No doubt, intermediate products—amylan-like bodies, sugars, &c.—are at first produced, but the granules are the first visible products of the assimilation found in the leaves. The starch thus formed is transferred to the reserve organs as such, or, more probably, as some sugar or other transformation product or products, as leaves are known to contain transforming agents (Baranetzky, Die stärkeumbildenden Fermente in den Pflanzen, Leipzig, 1878; Brasse, C. R. 99, 878).

Preparation.—Starch of commerce is prepared from various sources—viz., amongst others, wheat rice, maize, potatoes, Maranta indica (American starch), Maranta arundinacea (arrowrect), the roots of Jatropha Manihot or Manihot utilissima (tapioca), the stems of several species of Sagus or Cycas (sago). For manufacturing methods see Thorre's Dictionary, art. 'Starch.' In the laboratory, starch can be prepared from any starch-containing material as follows:—

From materials that can be ground, such as the cereals, &c., wheat, barley, maize, rice, &c. The material is ground in a coffee-mill, and the meal steeped in a 0.6 to 1 p.c. sol. KHO or NaHO. After standing 24-36 hours the coarser portions are separated by straining, with rubbing, and a slight flow of water through a wire sieve of about 20 wires to the inch. The strained milky liquid is allowed to stand for a short time, when a layer of crude starch settles at the bottom of the vessel. The liquid, with the matter in suspension, is transferred to another vessel, and again allowed to stand, when a second deposit of crude starch takes place. The process may be repeated a third and a fourth time. The whole of the suspended matter is thus allowed to settle, when the fairly clear supernatant liquid is decanted off and rejected. The deposit is then rubbed through a fine hair sieve, with a slight flow of water, and allowed to deposit a layer of starch as before. This is repeated as long as a starch layer separates. The whole of the starch layers are then collected, again suspended in water, and allowed to settle. At times a layer contaminated with much brown matter falls out first; from this the supernatant liquid with the starch in suspension is decanted off and allowed to stand, when a fairly pure deposit of starch is obtained. Further crops can be got from the residue, but it is difficult to free them from fibre, &c.

From materials that cannot be ground, such as potatoes, bulbs, rhisomes, other roots, and pith. The well-washed material is rubbed down with a grater into water, to which afterwards the alkali is added. The first deposit in these cases contains, as a rule, earthy matter, but it is easily separated by allowing it to settle, which it does in a short time, and then decanting off the starchy liquid. Two or three depositions and strainings through a fine hair sieve give a clean starch.

In dealing with the cereals the starch can be also separated by what is known as the acid process. The meal is steeped in water and kept at a temperature of from 25° to 27° until acid is developed. On then stirring up with water the light flocculent cell walls and undissolved albuminoids separate and allow the starch to deposit on standing. By repeated suspensions, stirrings, and depositions, clean crops of starch are obtained.

These products, like the starches of commerce, are not pure, but, as a rule, contain ash, oil or fat, albuminoids, &c. Some of them are slightly alkaline, some acid, from the condition of the water used in the last washing. Purification is effected by treatment with dilute KHO solution (0.5 p.c.), then dilute HCl (up to 1 p.c.), then with strong alcohol, and finally with water. They then can be dried by exposure to the air on layers of filter-paper.

Structure.—The starch thus obtained varies much in appearance, from the glistening silky white of potato starch to the dead chalky white of rice or maize starch. It consists of microscopic granules, varying in size from about 0.2

The same plant always mm. to 0.002 mm. yields granules of specifically the same size and shape, with as little variation as the size and shape of the leaf, hence, from the microscopic appearance of the granules, it is always possible to tell the source of a sample of starch. If it is wished to determine the source of any sample of starch, the following classification of the starches will be of some use. If, under the microscope, the granules are

(a) large, rounded, and more or less marked with rings: the starch may be potato, tapioca,

sago, &c.

(b) rounded, very slightly ring - marked: barley, rye, wheat, &c.

(c) reniform: beans, peas, &c.
(d) intermediate in size, rounded, and irregular: crocus, tulip, and other bulbs.

(e) irregular in figure, bounded by surfaces more or less plain : maize, oats, rice, &c.

(f) small granules, rounded or irregular:

fern, chestnut, parsnip, &c.

It will be observed that some of these are round, oval, or reniform, while others are irregular figures bounded by surfaces more or less plain. The microscopic appearance does not always give the true form of the granule, but a form which is the result of the pressure of the cover glass; for example, the well-marked star with central spot in rye starch and the split and striated appearance of maize starch are produced by the pressure of the cover glass on the granules. The granules of some starches are marked with well-defined more or less concentric rings, well shown in potato starch; in others there are no markings, the granules appearing as a clear, transparent cell. It is not absolutely agreed upon whether the starch granule consist of a single body or is made up of two or more. Some hold that the properly purified granule is a single substance, the coating or layers of which differ from the contents simply in containing less water and being thus more dense. According to Nageli (Die Stärkekörner) and others, the dense portions consist of starch cellulose, while the less dense, transparent plasma is granulose. The stratified structure is the result of the mode of growth of the starch granule, the additions being made by intussusception, i.e. from within outwards. It is said that when the granule is ruptured the granulose dissolves in cold water, leaving the starch cellulose; the former gives the well-known blue reaction of starch with iodine, while the latter is only coloured yellowish (Jessen, P. 106, 497; J. pr. 105, 65; Brown a. Heron, C. J. 35, 610; Brukner, M. 4, 889). On heating the insoluble portion with water it also gives the characteristic blue reaction with iodine. Granulose may be separated from starch cellulose (a) by digesting the granules for several days with a saturated solution of NaCl containing 1 p.c. HCl—the cellulose remains undissolved (F. Schulze, Henneberg's Jour. Landwirth., new ser., 7, 214); (b) by digesting starch with saliva at 45°-55°, this dissolves the granulose (Nägeli, Die Stärkekörner, 110; (c) by the action of certain schizomycetes, which decompose the granulose of starch-paste and leave the cellulose untouched (Fitz, B. 10, 282); and (d) by the action of malt-extract on starch paste in the cold, the cellulose is left undissolved.

A. Mayer (C. C. 1887, 6; Bot. Zeit. 1886, 693) considers that the starch granule is a homogeneous substance, and that the terms 'granulose' and 'starch cellulose' must be abandoned, for he points out that the delicate transparent skeletons left when the gelatinised granules are acted on by saliva, dilute acids, &c., are produced by the action of the reagent on the starch, and are identical with amylodextrin. I am inclined to the view that starch granulose differs from starch cellulose in being less dense in consequence of containing less water of hydration. Starch granules act on polarised light, and when examined with the microscope between two Nicols, produce very pretty effects; v. Bailey (P. M. [5] 2, 123).

Properties.—Air-dried starch sometimes contains over 20 p.c. H₂O; this it loses, slowly towards the end, in a vacuum over sulphuric acid; by gradually raising the temperature to 100°, under these conditions, it soon becomes absolutely free from H2O. The specific gravity of air-dried starch varies very considerably, the variation being due in the main to the quantity of moisture. Dry starches, however, would appear not to be absolutely alike in specific gravity; that of anhydrous potato starch is 1.650, whilst anhydrous arrowroot starch is 1.5648 (Flückiger, Fr. 5, 305; Saare, J. 1884, 1654). Dry starch takes up water, with the evolution of much heat. It does not dissolve in H2O, and has neither taste nor smell.

(a) Action of heat. Dry starch is not coloured at 100°, and, indeed, the temperature can be increased considerably beyond that point without being changed. Starch containing water is, however, coloured by a moderately low temperature, and if the heat be increased to 160° a soluble product is obtained which is known as dextrin or British gum. It is a mixture of undescribed composition. The action of heat on dry starch has not yet been accurately recorded. When the heat is increased beyond 160°, and destructive distillation begins, carbon dioxide, gaseous hydrocarbons, water, acetic acid, and an empyreumatic oil are evolved, and finally a carbonaceous porous cinder is left.

(b) Action of water. As long as the granules are uninjured, starch is insoluble in cold water: when, however, the water is heated, the granules swell up, and a gelatinous, more or less transparent mass, known as starch-paste, is produced. This varies in transparency with the starch employed, as does the temperature at which the gelatinisation takes place. E. Lippmann (C. C. 1861, 859; J. pr. 83, 51) gives the following table on this point:

Source	Swelling-up temp.	Commencement of gelatinisation	Perfect gelatinisa- tion
Rye	45°	50°	55°
Maize	50°	55°	62·5°
Barley	87·5°	57·5°	62·5°
Potato	460	59°	62·5°
Rice	54°	59°	63°
Wheat	60°	65° ·	67·5°

If the paste is sufficiently dilute it can be filtered, but it is doubtful if the filtrate is a true colution Picton a Linder, C. J. 61, 156). The solution (Picton a. Linder, C. J. 61, 156). consistency or stiffness of starch-pastes, containing the same amount of dry starch, seems to

vary with the variety of starch employed, and, indeed, with the method of preparation, even with the same starch; but I feel inclined to think that if closer attention were given to the amount of dry starch and its purity much of this apparent difference would disappear. On the relative stiffness of flour or starch pastes, and the mode of estimating it, see Thomson (S. C. I. 1886, 143). If the paste be heated under pressure to temperatures above the boiling-point maltose and dextrin are said to be produced, but the change has not been accurately studied.

(c) Action of glycerol. Starch heated with glycerol to 190° is dissolved; alcohol ppts. soluble starch from the solution. If the heating be continued at 200°, dextrins are formed (Zulkowski, B. 13, 1395; 23, 3295; C. C. 1888, 1060).

B. 13, 1395; 23, 3295; C. C. 1888, 1060).
(d) Action of acids. Moderately strong HCl in the cold converts starch, in a few days, without changing materially its microscopic appearance, into a modification perfectly soluble in water. This body is identical with soluble starch prepared by the limited action of malt extract on starch paste (C. Lintner, J. pr. 34, 378; Brown a. Morris, C. J. 55, 450). Prolonged action of 12 p.c. HCl in the cold produces amylo-dextrin (Nägeli, Beiträge s. Kenntniss d. Stärkegruppe; Brown a. Morris, C. J. 55, 450). Boiling dilute acids convert starch, first into soluble starch, then into dextrin and maltose, intermediate products, amyloins, are said to be formed (Brown, Morris a. Moritz, E. P. No. 1809 of 1889), and finally dextrose. The complete conversion takes place the more quickly and perfectly the higher the temperature and the longer the period of the reaction (Allihn, J. pr. 22, 46; D. P. J. 250, 534). This is only accomplished by employing diluted acid. Carbonic acid, oxalic acid, &c., act like HCl and H₂SO₄, but less energetically. For the manufacture of dextrose (glucose) from starch, vide THORPE'S DICTIONARY. The action of acids has been studied chiefly by Kirchhoff, Guérin-Varry, Payen (see Gm. K.), Musculus (A. Ch. Vary, Fayen (see Gm. A.), Musculus (A. Ch. C.), C. (B.) 60, 203; J. pr. [2] 28, 496; Bl. 30, 4); O'Sullivan (C. J. 25, 581); Musculus a. Gruber (C. R. 86, 1459; Bl. [2] 30, 54); Bondonneau (C. R. 81, 972); Salomon (J. pr. [2] 25, 348; 26, 342; 28, 82 a. 122); Sachsse (C. C. [3] 8, 782); Schulze (J. pr. [2] 28, 311); Sostegne (G. 15, 376); Nijerli (Strategrappe, Leipnig, 1874). 1887, 376); Nigeli (Stärkegruppe, Leipzig, 1874, 33, 99). As a summary of this work it may be stated that dextrose is the final product, but that acids act on this, to some extent yielding products still imperfectly investigated; that intermediate substances, dextrin and maltose and compounds thereof, are first produced; that the rapidity of the change varies with the strength of the acid, with the temperature, and with the pressure at which the change is effected, the most complete and perfect production of dextrose resulting when the conversion is hastened under pressure in presence of a small percentage of acid, $1\frac{1}{2}$ to 2 p.c. or less, and the proportion of dry starch to dilute acid does not exceed 1 to 3. Gallasin (C. Schmitt a. Cobenzl, B. 17, 1000; Rosenbek, B. 17, 2456), a gummy body, is found in commercial glucose. This is identical with isomaltose obtained by Fischer (B. 23, 3687) by the action of strong HCl on dextrose (Scheibler a. Mittelmeier, B. 24, 801).

Action of diastase (malt extract). Diastase does not act on ungelatinised starch in the cold. (O'Sullivan, C. J. 80, 133; Brown a. Heron, C. J. 35, 596), but Kjeldahl has shown that this is not true of all starches. This is probably due to some condition of the starch connected with the state of ripeness of the material whence it was obtained. The action, and the products thereof, of diastase on starch paste has been the subject of much study, but as yet only the broad facts are agreed upon. When starch paste is heated to 60° or thereabouts, and a little prepared diastase solution or cold water extract of malt added, the pastiness begins immediately to disappear, the solution rapidly loses the power of giving a blue colour with iodine, and acquires, for a short time, the property of giving with that reagent a deep reddish-brown colour. This, too, it rapidly loses if the diastase is in sufficient quantity. The solution is then perfectly clear when some starches are employed, with others there is more or less flocculent suspended matter in a clear solution. In the cold this dissolution takes place slowly. So far the reaction can be followed with little trouble, but when it comes to a question of the products of the action, the subject becomes more difficult. Musculus (A. Ch. [3] 55, 203) states that when diastase dissolves starch paste at 70°-75°, the products consist of 1 mol. sugar and 2 mols. dextrin, and that no further action takes place. Payen (A. Ch. [4] 4, 286) asserts that more than 50 p.c. of the solid matter dissolved by the reaction is sugar, and says (A. Ch. [4] 7, 382) that four samples taken from an operation in the space of 14 hours contained 17.9, 20.9, 25.8, and 26.03 p.c. sugar on the total solids dissolved. Schwarzer (J. pr. [2] 1, 212) agrees with Musculus in finding equivalent quantities of dextrin and sugar in solution, but differs from him in supposing that dextrin is formed first, then sugar, and that the action ceases when definite equivalent proportions are produced. He says less sugar is produced at 65° than at lower temperatures; above 65°-70° the proportion of sugar to dextrin is as 1 eq.: 3 eqs.; below 60°, when the principal phase of the reaction is finished, the dextrin and sugar are in the proportion of 1 to 1. The change was considered complete when the solution no longer gave a colour with iodine. These investigators took the sugar as dextrose and estimated the dextrin by difference. O'Sullivan (C. J. 25, 581; 30, 137) showed that the sugar produced was not dextrose but maltose, and that the method of estimating the dextrin was wrong. He also demonstrated that dextrin and maltose were the invariable products of the transformation, and that by continuing the action the whole of the dextrin could be converted into maltose. It was, however, indicated in this work that another body-from which it was impossible to eliminate the cupric reducing power, and which seemed to be a mixture of maltose and dextrin-was amongst the products. Although this work laid the foundation of all that has since been done in the transformation products of starch, it received for a time no attention from the Continental chemists. Musculus (Bl. 22, 32) states that the saccharification of starch paste with diastase ceases when half the matter in solution is sugar. He attri-

butes his original impression that only one-third was converted into sugar to the varying structure of the starch granules, the coating of one variety offering a greater resistance to the action of the diastase than that of another. He still holds that the action is a molecular splitting-up of the starch molecule, attended with hydration, into dextrin and sugar. Bondonneau (C. R. 81, 1212) concludes that the action is not a splitting-up, but that the starch molecule must pass through four isomeric modifications, viz., amylogen $[a]_D$ = +216°, a-dextrin = +186°, b-dextrin = +176°, and γ -dextrin = +164°, before it arrives at the end-product, glucose. These bodies increase in cupric-reducing power as they descend to the final product. Petit (Bl. 24, 519) mentions, as a constituent of the transformation products, a sugar soluble in alcohol, fermentable, and without action on alkaline copper solution. This is, undoubtedly, to be traced to the fact that maltose does not reduce as much copper oxide as dextrose. O'Sullivan (C. J. 30, 125; 35, 770) showed conclusively when working under clearly-defined conditions that maltose and dextrin were the only products of the action, although, as stated, he pointed out the presence of a body which gave a reduction with copper oxide equal to 9-10 p.c. dextrose. The opticity of this body ($[a]_1 = +204^{\circ}-206^{\circ}$) was such as to indicate a mixture of 12 3-15 3 p.c. maltose from the reduction, and the remainder dextrin. He concluded that if the amount of reduction of the products be calculated as maltose, and the opticity due to this be subtracted from the total opticity observed for them, the remainder will agree with the activity of an amount of dextrin obtained by subtracting the determined maltose from the total products. He further showed that: (a) malt extract begins to dissolve starch at the temperature of gelatinisation, or a few degrees lower; (b) perfectly gelatinised starch is almost completely dissolved in the cold (10°-20°); (c) at 63° the action takes place according to the equation

A.
$$C_{18}H_{30}O_{18} + 0H_2 = C_{12}H_{22}O_{11} + C_6H_{16}O_8$$
; Starch Maltose Dextrin (d) at 64°-70° to

B. $C_{18}H_{30}O_{18}$ $\Big\} + 0H_2 = C_{12}H_{22}O_{11} + 4C_6H_{10}O_8$; Starch Maltose Dextrin (e) at 70°-71° (C. J. 1879) to

B'. $C_{72}H_{120}O_{50} + 80H_2 = 3C_{12}H_{22}O_{11} + 6C_6H_{16}O_8$; (f) and 71°-75° to

C. $C_{72}H_{120}O_{50} + 0H_2 = C_{12}H_{22}O_{11} + 10C_6H_{16}O_8$;

(g) and he stated that although the proportions of maltose and dextrin indicated by these equations are easily obtainable when the strict conditions are adhered to, it was only necessary to vary the proportions of malt extract and starch, and time and temperature, to obtain with ease products containing from 16 to 92 p.c. maltose. It is further pointed out that the final product, maltose, was very slowly converted into dextrose by some samples, at least, of malt extract. O'Sullivan (C. J. 35, 770) showed that soluble starch was the first product of the action, and that the changes afterwards brought about may be expressed:

 $0. \ \, C_{12}^{\ Starch} O_{\bullet \bullet} + OH_{2} \ \, = \ \, \begin{array}{c} Maltose \\ C_{12}H_{120}O_{11} + C_{\bullet \circ}H_{100}O_{40} \end{array}$

From the action of malt extract on these products he concluded that it is possible that only one β-dextrin exists. The theory of splitting up and breaking down of the starch molecules, as represented by the equations, does not account for all the facts eliminated. A theory which arranges all the molecules in solution in groups, dependent one on the other-and capable, therefore, of undergoing a simultaneous change, or of being moved at the same time-and the re-arrangement of these groups attendant upon the hydration of a definite proportion of the molecules in each of them, is more in accord with the eliminated facts. Märker (L. V. 22, 69) states that at 60° four molecules starch yield three of maltose and one of dextrin, at 65° there is less maltose, and at a higher temperature two molecules starch yield one of maltose and one of dextrin. Musculus and Gruber (Bl. 30, 54) look upon starch as a polysaccharide containing at least five times the group C12H20O10. When this is acted upon by diastase, or dilute acids, it is broken down with hydration into maltose and a dextrin containing a C12H24O10 group less than starch; that this dextrin is broken down in the same way, maltose and a dextrin containing a C12H20O10 group less than the first one, and so on through the series by successive stages, until finally the solution contains only maltose. They point to the existence of three dextrins giving no reaction with iodine, and call them α , β , and γ achroo-dextrins. From the optical activity and K they attribute to these bodies, they do not agree with O'Sullivan; these factors do not agree with those of a mixture of maltose and dextrin. Brown and Heron (C. J. 35, 596) have eliminated out of the possible varying proportions of maltose and dextrin indicated by O'Sullivan eleven distinct transformations: viz. soluble starch, erythro-dextrin α and β , achroo-dextrins α to η , and maltose. They agree with O'Sullivan that the dextrins are without action on alkaline copper solution, and that the dextrins and maltose are the only products of the reaction. They do not mention the body from which O'Sullivan could not eliminate the reducing power, and they did not observe dextrose amongst the final products with malt extract. They agree with him that the higher the temperature at which the transformation takes place the less maltose is found amongst the products. They represent the normal reaction taking place between 10° and 60° by the equation

 $10C_{12}\overline{H}_{20}O_{10} + 8H_2O = 8C_{12}H_{22}O_{11} + 4C_6H_{10}O_{20}$ Starch Maltose Dextrin

They consider the starch molecule to be at least $10C_{12}H_{20}O_{10}$, and that the hydrolysis of it begins by the formation of maltose and erythro-dextrin a, containing $9C_{12}H_{20}O_{10}$; that this, in its turn, yields maltose and erythro dextrin β , $8C_{12}H_{20}O_{10}$, and so on to the end—product, maltose. They thus agree with Musculus and Gruber as to the

character of the action, but disagree with them as to its quantitative nature.

Herzfeld (B. 12, 2120; 13, 8469) agrees with O'Sullivan and others that erythro- and achroo- dextrins are without reducing power on copper solution, but he points to the presence of a substance amongst the transformation products which seems to hold a position between dextrin and maltose; he names it malto-dextrin. $K = about \frac{1}{3}$ maltose, $[\alpha]_j = +171.6^{\circ}$; thus K corresponds to 33.3 p.c. maltose. A mixture consisting of one-third maltose and two-thirds dextrin would have an opticity $[a]_i = +199^\circ$; hence this contradicts the statement of O'Sullivan, confirmed by Brown and Heron, that, taking the reducing power as maltose, the opticity of the remainder of the transformation products corresponds to dextrin. Herzfeld considers that Bondonneau's γ-dextrin is maltodextrin.

Brown a. Morris (C. J. 47, 527) confirm the presence of malto-dextrin, but they show that Herzfeld was dealing with an impure body, and that when its true K and $[a]_j$, viz. $K = 21\cdot1$ and $[a]_j = +193\cdot1^\circ$, are examined it is found that they correspond with a mixture of maltose and dextrin. It is completely converted into maltose by malt extract at 50° to 60°. They do not agree with Herzfeld that it is a hydration product of dextrin, but hold that it is produced from starch and the polymeric dextrins by the fixation of a molecule of water upon the ternary group $(C_{12}H_{20}O_{10})_3$, thus $\left\{ \begin{matrix} C_{12}H_{20}O_{10} \\ C_{12}H_{20}O_{10} \end{matrix} \right\}_2$. They have discarded the series of equations, and consider that equilibrium is attained when a condition of things represented by the equation

 $10\ddot{\mathrm{O}}_{12}\dot{H}_{20}^{2}\mathrm{O}_{10} + 8\dot{H}_{2}\ddot{\mathrm{O}} = 8\dot{\mathrm{C}}_{12}\dot{H}_{22}\mathrm{O}_{11} + 2\dot{\mathrm{C}}_{12}\dot{H}_{20}\mathrm{O}_{10}$ Starch Maltose Dextrin Starch is arrived at. This is always the result when the change takes place at 50° to 60°, or when a higher transformation-product is degraded at that temperature. This degradation is due to the hydrolysis of the more complex polymeric dextrins and malto-dextrin. Bourquelot (C. R. 104, 576) looks upon the action of diastase on starch as a successive fixation of a water molecule with the production of maltose and a lower dextrin until the reduction of the degraded products is K=51.52 (about 82.5 p.c. maltose), but he does not describe his dextrin or dextrins, and he concludes, as has already been clearly proved, that heat alters, not alone the quantity of diastase, but

the quality.

Brown and Morris (C. J. 55, 462), from determination of the molecular weights of some of the transformation products, conclude, as suggested by O'Sullivan in 1879, that the dextrins are metameric, and not polymeric, as they hitherto considered them. They therefore abandon their former theory of the hydrolysis of starch by diastase, and, as far as I can see, come to a broad agreement with the theory exunciated by O'Sullivan. They mention another product, amylo-dextrin, $[a]_1 = +206^\circ$, $K = 9\cdot0$. Like malto-dextrin, it appears to be a compound of maltose and dextrin, $\{C_{12}H_{20}O_{10}\}_0$. Its op-

ticity and K agree with the body described by O'Sullivan in 1872, and mentioned above.

A new phase has been given to the trans-

formation by C. J. Lintner (Zett. ang. Ch. 1892, 263). He says he has isolated a body, which he calls isomaltose, from amongst the products. It is less fermentable and less soluble in alcohol than maltose, with $[\alpha]_D = +139$ and K=52, the opticity being nearly that of maltose and the reducing power about 84 p.c. thereof. Diastase converts it into maltose completely.

Since the publication of the above Morris a. Wells (Transactions of the Institute of Brewing, 5, 133), and Moritz (ibid. 4, 141) point to the presence of a whole series of amyloïns or malto-dextrins amongst what they call restricted starch-conversions, and they assert that isomaltose is a low amyloïn—i.e. one in which the maltose constituent largely predominates. These amyloïns begin with a high proportion of the dextrin residue, and end with a high proportion of maltose. Their optical activity and reducing power are such as would be yielded by mixtures of maltose and dextrin.

Schifferer (In. Dissertation, Basel) denies in toto the existence of the amyloins, and asserts that nothing is to be found amongst the products of the action of diastase on starch but achroo-dextrin, isomaltose, and maltose. His dextrin, however, is a reducing dextrin; even otherwise, the work will not bear criticism.

Action of other ensymes. Ptyalin (ptyalase). This enzyme of saliva liquefics starch paste (Nägeli, Die Stärkekbrner, p. 113; Lefberg a. Georgieski, Bl. [2] 25, 393; Dobroslavine, Bl. [2] 26, 452; Maercker, L. V. 22, 69; Watson, C. J. 1879, 539; Musculus a. De Mering, Bl. [2] 31, 105; Lea, J. Physiol. 11, 226). The products are maltose, a reducing, unfermentable dextrin, and a little dextrose. Ungelatinised starch is not acted upon by ptyalase; at a temperature slightly below the gelatinising point the starch is dissolved, and the action is most rapid when the ferment acts at 60° on previously-boiled starch paste (Bourquelot, C. R. 104, 71 a. 177).

Enzymes capable of dissolving starch have been observed in the pancreatic juice (Brown a. Heron, Pr. 1880, 394), in the small intestine (Hoppe-Seyler, Phys. Ch. 275; Brown a. Heron, Lc.), in the liver (Wittich, Pf. 7, 28; Bernard, C. R. 85, 519; Abele, Th. J. 6, 271), and in many other animal tissues (Ellenberger a. Hofmeister, Th. J. 12, 501; Paschutin, Th. J. 1,304). The stomach, pancreas, and peritoneal lymph of fishes contain an enzyme capable of dissolving starch (Richet, Th. J. 14, 359; Kruckenberg, Unter. Phys. Inst. Heidelberg, 1, 2). Blood contains a diastatic enzyme, for starch injected into the blood yielded maltose and dextrin (Bimmerman, Pf. 20, 201; Ploz a. Tiegel, Pf. 6, 249).

Starch-dissolving enzymes exist in the leaves and other parts of most plants (Baranetzky, Die stärkeumbildenden Fermente in den Pflanzen, Leipzig, 1878; Vines, Ann. Bot. 1891, 409).

Certain organisms, bacteria, moulds, &c., are capable of secreting an enzyme or enzymes having the power of dissolving starch. The nature of the dissolution products has, as yet, been very imperfectly studied; we have simply the general statement that they are sugar or sugars and dextrin (Wortman, Z. P. C. 6, 287; Fitz, B. 10, 282; Marcona, C. R. 954. 345 a. 856; Gayon a. Dubourg, C. R. 103, 885;

Atkinson, Pr. 82, 299; Takamine, E. P. 5700 a. 17374, 1891). Bacillus amylobacter yields dextrin and a small quantity of a crystalline body (Villiers, C. R. 112, 435 a. 113, 144 a. 536). Under the influence of Bacillus suaveolens, dextrin, glucose, alcohol, aldehyde, formic, acetic, and butyric acids are produced. Amylic alcohol is the result of the action of Bacillus amylozymicus, no doubt intermediate bodies; dextrin and sugar are at first produced. Atkinson states that the moulds grown on steamed rice in the manufacture of saké, in Japan, secrete an enzyme, which first converts starch into dextrin and maltose, and further acts on the maltose and dextrin, with the production of dextrose.

Some gums of the arabin group contain a starch-dissolving enzyme (O'Sullivan, C. J. 1891, 1061)

Maize, malted and raw, and other grains contain an enzyme capable of dissolving starch and yielding as a final product dextrose (Cuisinier, C. C. 1886, 614). Its action on starch is not very vigorous, but it acts more rapidly on dextrin and converts maltose very rapidly into dextrose (Geduld, Wochenschrift f. Brauerei, 8, 620; Lintner, Zeit. f. ges. Brau. 1892, 123). It would be interesting to determine in what respect this enzyme differs from that of the moulds, and to establish their separate existence.

Action of the halogens. Chlorine does not stain starch. Bromine colours it yellow. Iodine gives with it an intense blue. This reaction is a distinctive test for starch. The colour is destroyed by heating, but returns on cooling; but if the solution be boiled for some time, the colour does not reappear. The blue colour is also discharged by arsenious and sulphurous acids, by alkalis and carbonates thereof, and, indeed, alcohol can remove the iodine. A solution of iodine in strong alcohol does not colour dry starch. Some observers consider that the production of this blue colour is not due to a definite chemical combination of iodine with starch (Vogel, N. Rep. Pharm. 22, 349; 25, 565; Pellet, M. [8] 7, 988; Tomlinson, P. M. [5] 20, 168; Duclaux, A. Ch. [4] 25, 264), while others attribute a definite formula to the combination. According to Bondonneau (C. R. 85, 671), it is (C₆H₁₀O₅)₅I; Mylius (B. 20, 688) considers it to contain HI, and gives the formula (C₂,H₄₀O₂₀I),IH as probable. He finds that the HI can be displaced by metallic iodides. Stocks (C. N. 56, 212; 57, 183) and Seyfert (Zeit. ang. Ch. 1, 15) contradict this. Rouvier (C. R. 114, 128 a. 1366) attributes the formula (C.H.O.), I to the compound. Starch is oxidised by chlorine and by bromine to gluconic acid (Habermann, A. 172, 11; Herzfeld, A. 220, 864).

Action of alkalis. Weak solutions of the alkalis do not act on starch in the cold, but solutions containing over 8 p.c. real alkali cause the granules to swell up with the formation of a thick transparent paste, and, finally, a clear solution, a compound of starch with the alkali being formed (Schmidt, A. 51, 81; Ventzke, J. pr. 25, 65) which, according to the latter, is optically inactive, but this, no doubt, is in-correct, for Béchamp (C. R. 39, 653) gives the opticity [a], = +211° for the starch dissolved, and Thomsen (B. 18, 2168) shows that the ac-

tivity of dilute sods solutions is $[a]_D = +168$. On neutralisation this becomes much higher, corresponding, in fact, with Béchamp's number. The product does not reduce alkaline copper solution (Brown a. Heron, C. J. 35, 617). The potassium compound is obtained by ppg. the solution in dilute KHO with alcohol, pressing the pp., dissolving in H₂O, and re-ppg. with alcohol. This process repeated three or four times is said to yield a pure compound of the composition $C_{24}H_{52}O_{20}K$ (Pfeiffer a. Tollens, A. 210, 288). A sodium compound C₂₁H₂₉O₂₀Na has been obtained in the same way (Reichardt, Z. 1870, 404). These formulæ are, however, very improbable. Alcoholic soda does not act on starch (Dragendorf, J. f. Landwirthschaft, 7,

Starch heated with ammonia yields brown, amorphous, nitrogenous bodies (Thénard, C. R. 52, 444).

Fused with KHO, starch, like other carbohydrates, yields oxalic acid, acetic acid, and other products.

Action of alkaline earths. Barium, strontium, and calcium compounds similar to the sodium and potassium bodies have also been prepared. When solutions of soluble starch are ppd. with solutions of lime in sugar syrup, pps. are produced which are not very definite in composition, the percentage of lime showing a variation between C₂H₁₀O₃CaO and (C₂H₁₀O₃)₄CaO. Similar baryta compounds have been examined. When a solution of soluble starch is saturated with strontia and alcohol added, a strontium compound is ppd. (Lintner, Zeit. f. ang. Ch. 1888, 232). On distilling starch with lime acetone, mesityl oxide, isophorone, and ketones are produced (Harvat, C. C. 1887, 38).

On digesting starch with acetic anhydride, a triacetate is said to be formed $C_sH_1O_s(C_sH_2O_s)$, (Schützenberger a. Naudin, Bl. [2] 12, 110; A. Ch. [4] 21, 235; Michael, Am. 5, 359). This substance is amorphous; it is stained blue by iodine, and is decomposed by alkalis, with the

reproduction of starch.

Qualitative determination. The presence of starch is indicated by the granular appearance under the microscope, and starch granules are distinguished from all others by being stained blue by iodine solution and yellow by bromine. If the plasma in which the granules are contained is alkaline, it must be rendered slightly acid, or sufficient iodine solution must be employed to destroy the alkalinity. A solution of iodine in potassium iodide is usually employed, but an alcoholic solution answers the purpose. When the test is applied for starch in solution, the solutions must be cold and slightly acid and the reagent must be added in small quantities at a time. It is sometimes necessary to test a solution for soluble starch in presence of a-dextrini.e. the dextrin giving a reddish-brown colour with iodine. If this is in excess the reddishbrown colour covers the blue; ammonia added cautiously, drop by drop, discharges the reddishbrown, and if soluble starch be present the blue becomes definite and distinct. Care must be taken to avoid excess of ammonia (O'Sullivan). Small quantities of starch, which would otherwise not be observable, may be detected in the 'last runnings' of malt wort by adding a little

tannin and then alcohol to the solution. The starch is ppd. in this way, and on washing the pp. with water will yield the characteristic reaction with iodine (Burckhardt, Chem. Zeit. 1877, 1158). Starch, when moistened with an alcoholic solution of α -naphthol and a few drops of warm concentrated sulphuric acid added, acquires a deep violet-red colour (Ihl, Chem. Zeit. 11, 19).

Quantitative determination. There are very few materials containing starch of which a sufficiently accurate average sample can be obtained to render a definite estimation of the starch

therein of much value.

In some cases separation of the starch, by one of the methods given for its preparation, gives results of sufficient accuracy to satisfy the requirements.

In the case of potatoes, the percentage of starch is deduced from the specific gravity, a set of specially-constructed tables being used; but this method, on the face of it, cannot yield more than an approximation. The specific gravity of the washed potatoes is taken in the usual way, a balance constructed to meet the requirements of the case being employed. About 5 kilos. of the potatoes are weighed in a strong wire basket in air, and then in water; thence,

specific gravity = weight in air - weight in water.

According to the tables of Behrend, Märker, and
Morgan, we have from

G	Per cent. of		Per cent. of
Sp. gr.	starch	Sp. gr.	starch
1.080	13·9	1.120	22.5
1.090	16·0	1.130	24.6
1.100	18.2	1.140	26.7
1.110	20.3	1	

In cases in which fair average samples of the material can be obtained, several methods have been proposed, all depending on the conversion of the starch, or the starch transformation products, into dextrose by digestion with dilute HCl or H2SO4, the dextrose being estimated with Fehling's solution, volumetrically or gravimetrically, and the starch calculated therefrom according to the equation $C_6H_{10}O_6 + H_2O = C_6H_{12}O_6$ (v. Sachsse, C. C. [3] 8, 732; Märker, Handb. der Spiritusfabrication, 4 Auf., Berlin, 1886, 93; Soxhlet, Zeit. f. ges. Brau. 1881, 177; Faulenbach, Zeit. f. phys. Chem. 7, 510; Zip-perer, Rep. An. Chem. 6, 699). Dubrunfaut dissolved the starch by rubbing the material with concentrated HCl, diluting the solution with water to a certain volume, and determining the opticity. The starch was calculated from $[a]_1 = 192.4^{\circ}$

Effront (Bl. [2] 47, 5) dissolves the starch with malt extract, determines the optical activity of the solution, then heats it with ammonis, after which he treats it with sodium hypochlorite and with HCl, and again determines the opticity; but this method is open to many

objections.

These methods are unsatisfactory, inasmuch as it is difficult, if not impossible, to completely convert starch or starch products into dextrose, there being a probable error at one time from overconversion, at another from under-conversion; and, indeed, both may occur at the same time,

the amount of reduction of Fehling's solution taken as dextrose never being a correct measure of the starch transformed. Another objection to the method is due to the fact that dilute acids convert other substances besides starch into bodies capable of reducing copper oxide.

Girard (C. R. 104, 1629) titrates starch with iodine solution, 1 grm. of starch taking up 0.157

grms. iodine.

Asboth (Chem. Zeit. 11,785) proposes to add an excess of baryta water to the gelatinised starch and then alcohol. The dried pp. contains 198 p.c. BaO, the remainder is starch. This

method, too, cannot be of any use.

O'Sullivan (C. J. 45, 2) described a method by which starch in most materials can be estimated with accuracy. If a fair average sample of the material cannot be obtained by grinding and powdering alone, it is first dried in warm, dry air and then powdered. The following is the method as applied to the cereals, but it is also applicable to all materials when treated as just described.

Fivegrams, or thereabouts, accurately weighed -more if the material contains less than 40 p.c. starch—of the finely-ground material are introduced into a wide-necked flask of 100 to 120 c.c. capacity. To this sufficient alcohol of sp. gr. 0.82 is added to saturate it, and after a time 20 to 25 c.c. ether are introduced. The clear, ethereal solution is decanted off after standing for a few hours, and the residue again treated two or three times with ether. It is then extracted with alcohol, sp. gr. 90, at 35° to 38°, and treated with a large bulk of water, with which it is left in contact for at least 24 hours. If any sign of fermentation shows itself, a little salicylic acid solution may be used with the water. The residue, after being completely extracted with water, together with the filter, through which all the extracts should have been passed, is transferred to a beaker of about 100 c.c. capacity, and made up with water to about 40 to 45 c.c. This is heated to boiling for a few minutes in a water bath, care being taken, by continual stirring, to insure a homogeneous paste; then cooled to 62° to 63°, and 0.025 to 0.035 gram prepared diastase or its equivalent of malt extract added, the digestion being then continued at the temperature stated for a few hours. At the end of that time the contents of the beaker are boiled for a few minutes, thrown on to a filter, and the filtrate received in a 100 c.c. measuring flask The residue is carefully washed with small quantities of boiling water at a time, and the filtrate made up at 15.5° to 100 c.c. A determination of the reducing power calculated as maltose, and of the remainder of the optical activity as dextrin, gives the data for calculating the quantity of starch. This holds good even though the amyloin theory of the breaking down of starch be proved to be accurate.

Example.—5 grms. barley-flour treated as described, 008 grm. prepared diastase being employed, gave 100 c.c. solution of sp. gr. 1·01008 = 2·589 grms. solid matter. 9·178 grms. this sol. reduced 0·241 grm. CuO. Opt. act. in

1 If cold-water malt extract be used, a portion of it must be heated for the same time and at the same temperature as the assay; then bolled, and the opticity and iscluding power determined; these factors calculated on the quantity amployed must be allowed for.

200 mm. tube = +21.1 divs. (Soleil-Ventzke-Scheibler saccharimeter). Hence

0.241 + 0.7256 (K of maltose = 62.5) = 0.1748 grm. maltose;

9.178 grms.: 101.003 (the wt. of 100 c.c. sol.) ::0.1748:x;

x = maltose in the 100 c.c. sol. = 1.923; Opt. act. maltose $[\alpha]_j = +154^{\circ}$ and of dextrin $[\alpha]_j$ = + 222°

Hence 1 grm. maltose in 100 c.c. sol. in 200 mm. tube = 8.52 divs. of instrument mentioned, and 1 grm. dextrin under like circumstances = 11.56 divs. Then $1.923 \times 8.02 = 15.422$ divs. opt. activity of the maltose, $21 \cdot 1 - 15 \cdot 422 = 5 \cdot 678$ divs. opt. activity of the dextrin, and 5.678 + 11.56 = 0.491grm. dextrin in the 100 c.c. sol. Dextrin is derived from starch without any increase of weight: 100 grms. starch yield 105.5 maltose; hence

1.055 : 1.923 :: 1 : x starch = maltose;

x = 1.822 grms. starch.

1.822 + 0.491 (as dextrin) = 2.313 starch; in

the 5 grms. taken = 46.26 p.c.

It is clearly shown in the paper quoted that if starch is not estimated in the way indicated no reliance can be placed on the results. It happens when dealing with some varieties of material that the aqueous extract contains soluble starch (blue colour with iodine). This cannot be looked upon as starch; it must be estimated in the solution as the soluble modification.

soluble starch may be prepared (a) by triturating starch with sharp sand or powdered glass so as to disintegrate the granules and extracting with cold water; (b) by the limited action of malt-extract or of acid.

Delffs (P. 109, 648) prepared soluble starch by triturating starch with sand and water; the solution gave a dark-blue colouration with iodine. Flückiger (Z. 1861, 104) prepared a similar solution by acting on starch with a concentrated solution of calcium chloride and treating the resulting gummy mass with water, when, on filtration, a solution is obtained which exhibits all the characteristics of soluble starch. Musculus (Bl. [2] 22, 26; A. Ch. [5] 2, 385) does not consider this a true solution; he prepares the body by boiling starch with very dilute sulphuric acid, saturating the solution with chalk, and evaporating to a syrup. This deposits small granules which gradually increase in size, are soluble in hot water, and may be purified by precipitation with alcohol. It possesses no reducing power; its rotatory power is four times that of glucose. donneau (C. R. 80, 671) has prepared soluble starch by this method, but does not find it altogether soluble under all conditions. Soluble starch is undialysable. Fuming nitric acid converts it into a mononitro-derivative, C,H,(NO,)O,; dilute nitric acid oxidises it to carbonic and oxalic acids; bromine and silver oxide to gluconic acid (Reichardt, B. 8, 1020; 7, 424).

Nägeli (Beiträge z. näheren Kenntniss der Stärkegruppe, Leipzig, 1874, p. 33, 99; A. Ch. 173, 218) gives the following method: 1 kilo. potato starch is allowed to stand 6-8 weeks with 6 litres hydrochloric acid, S.G. 1.06; this is not soluble starch (v. above). Zulkowski (B. 13, 1395) prepares soluble starch by heating dry potato starch with glycerol at 180° 190° for half an hour. The solution is cooled, precipitated by alcohol, and the precipitate purified by solution in water and reprecipitation by alcohol.

Salomon (J. pr. [2] 28, 82) finds that soluble starch is the first product of the action of dilute acids on starch; it does not reduce Fehling's solution, and has an optical activity $[\alpha]_D - 211.5^{\circ}$. O'Sullivan (C. J. 1879, 772) prepares soluble starch by dissolving starch paste at 73°-74° with the least possible quantity of cold water extract of malt, boiling the solution as soon as it becomes clear, filtering, and concentrating. The soluble starch falls out on cooling as a white precipitate, which is purified by dissolving. in hot water and allowing to cool when it separates out again. It has a reducing power, 3.5-0.78, and an optical activity $[\alpha]_i = 219.5$ -222.0, the reducing power being probably due to a small quantity of maltose; v. also Action of C. O'S.

acids on starch, above.

STEARIC ACID C₁₈H₃₆O₂. Mol. w. 284. [69°]. (232° cor. at 15 mm.) (Krafft, B. 17, 1629); (359°-383°) (Carnelley a. Williams, B. 102); (359-353) (Catheries & Windins, B. 12, 1360). H.F. 126,000 (Von Rechenberg). S.G. (liquid) \$\mathbb{2}\$ \cdot 845; (solid) \$\mathbb{1}\$ \cdot 101; \$\mathbb{1}\$ \cdot 100. S.V.S. 332-6 (R. Schiff, A. 223, 264). S. (alcohol) 2.5 in the cold. S. (benzene) 22 at 23°. S. (CS₂) 30 (Vogel, J. 1866, 892). Occurs as glyceryl stearate in very many fixed animal and vegetable fats and oils (Chevreul, A. Ch. 88, 225; [2] 2, 354; 23, 19; Braconnot, A. Ch. 93, 250; Redtenbacher, A. 35, 46; Bromeis, A. 35, 86; 37, 303; Stenhouse, A. 36, 57; Erdmann, J. pr. 25, 497; Francis, A. 42, 256; Gottlieb, A. 57, 35; Laurent a. Gerhardt, A. 72, 272; Hardwick, C. J. 2, 232; Crowder, P. M. [4] 4, 21; Berthelot, A. Ch. [3] 41, 216, 432; 47, 297; Pebal, A. 91, 138; Heintz, A. 92, 295; Johnston, C. J. 29, 8).

Formation.-1. By saponification of cetylacetoacetic ether, obtained from cetyl iodide and sodium acetoacetic ether (Guthzeit, A. 206, 351). 2. By heating cetyl-malonic acid at 150°.—3. From ricinoleic acid by treatment with water, P, and I, followed by zinc and HClAq (Claus, B. 9, 1916).-4. By heating oleic acid with iodine (1 p.c.) for several hours at 275° and distilling the product in a current of superheated steam (De Wilde a. Reychler, Bl. [3] 1, 295).

Preparation .- Suet or cacao fat is saponified by NaOHAq, the acids ppd. by H2SO, and crystallised from alcohol. An alcoholic solution of the impure stearic acid (4 pts.) saturated at 0° is heated to 60° and mixed with a boiling alcoholic solution of Mg(OAc), (1 pt.). The ppd magnesium stearate is boiled with HClAq and the stearic acid recrystallised from alcohol (Heintz). Stearic acid can also be readily obtained from shea-butter which contains no other solid fatty acid (Buff a. Oudemans, J. pr. 89, 215).

Properties.—Pearly plates, insol. water, sol. ohol and ether. Tasteless and inederous. alcohol and ether. May be distilled in vacuo and, in small quantity, under atmospheric pressure, but in this case it is partly decomposed with formation of hydrothen purified by solution in hot water; it crystal-lises in sphero-crystals (Jaquelain, A. Ch. [2] butyric acid. Fusion with P_2O_2 forms $C_{12}H_{24}O_3$, 173). According to Brown a. Morris, this is $[54^{\circ}-60^{\circ}]$. Nitric acid forms sebacic, glutarically $[54^{\circ}-60^{\circ}]$. STORAX. 5)8

succinic, and other acids. Br and water at 140° form bromo- and dibromo-stearic acids (Oudemans, J. pr. 89, 193). Distillation in vacuo with NaOMe yields C₁₇H₃₆ (Mai, B. 22, 2133).

Salts.—KA'. Hygroscopic crystals. in hot water. Partially decomposed by a large quantity of water into an insoluble acid salt and free potash.—KHA'2. Silvery scales (from alcohol). S. (alcohol) 36 in the cold; 27 at 78°. Boiling water converts it into a more acid salt. NaA'.—NaHA'2: insol. water.—BaA'2: minute laminæ, insol. water.—CaA'2.—SrA'2.—MgA'2: minute laminæ (from alcohol).—CuA'2: lightblue amorphous powder. - PbA'2: amorphous powder, insol. ether. —Pb₂OA'₂. —AgA': white pp. Methyl ether MeA'. [38°]. Crystalline, insol. water (Hanhart, C. R. 47, 230).

Ethyl ether EtA'. [33°]. (224°). Formed by heating the acid with alcohol at 200° (Lassaigne, A. 13, 168; Berthelot, A. 88, 312) by passing HCl into an alcoholic solution of stearic acid (Redtenbacher, A. 35, 51) and by heating stearin with a little NaOEt (Duffy, C. J. 5, 197; Bouis, C. R. 45, 35). Crystalline mass, partially decomposed by distillation. V. sol. alcohol.

Ethylene ether C,H,A'₂. [76°]. Fro AgA' and C,H,Br, (Wurtz, A. Ch. [3] 55, 486).

Isoamyl ether C,H,A'. [25°].

Octyl ether C,H,A'. [45°].

Cetyl ether C,H,sA'. [55°-60°]. Laminæ (from ether) (Berthelot, A. Ch. [3] 56, 70).

Glyceryl ethers v. vol. ii. p. 622. A mixture of stearic acid and glycerin saturated with HCl at 100° forms C₂H₃Cl(OH)(OC₁₈H₃₅O)

Phenyl ether PhA'. [52°]. (mm.) (Krafft a. Bürger, B. 17, 1380). (267° at 15

p-Tolyl ether C,H,A'. [54°]. (276° at 15 mm.).

Chloride C18H38OCl. [23°]. (215° at 15

mm.). Crystalline mass.

Amide C₁₈H₃₅O.NH₂. [109°]. Formed by distilling ammonium stearate at 230° under pressure; the yield being 50 p.c. (Hofmann, B. 15, 984; cf. Carlet, Bl. 1859, i. 76). Formed also by the action of NH, on the ether or on the chloride. Converted by the action of Br and NaOHAq into stearyl-heptadecyl-urea (Turpin, B. 21, 2486).

Anilide C18H35ONHPh. [94°]. Formed by distilling aniline over stearic acid at 230°

(Pebal, A. 91, 152). White needles. Phenyl hydraside C_1, H_{ab} .CO. N_2H_2 Ph. [107°]. Formed by heating stearic acid with phenyl-hydrazine. White unctuous plates (from alcohol), sl. sol. cold alcohol, benzene, and ether

(Strache a. Iritzer, M. 14, 37).

Nitrile C₁₇H₁₈CN. [42°]. (275° at 100 mm.). S.G. 45 ·815; 140 ·779. Formed by dis-(275° at 100 tilling stearamide with P.O. (Krafft a. Stauffer, B. 15, 516, 1780) or by heating cyanostearic acid at

250° (Hell a. Sadomsky, B. 24, 2779).

DI - BROMO - IODO-, References. — Bromo-, CHLORO-, IODO-, NITRO-, and OXY-STEARIC ACID.

Isostearic acid (C.H.,),CH.CO.H. [88.5°]. 8° at 101 mm.). Formed from di-octyl-(278° at 101 mm.). acetoacetic ether or di-octyl-malonic acid (Conrad a. Guthzeit, A. 204, 11, 165). Colourless leaflets (from alcohol).—NaA': needles (from alcohol). AgA': thick white pp.

Ethyl ether EtA^{\prime} . (275°-280° at 100 mm.).

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STEARIC ALDEHYDE C, H, O2. TR4º1. (213° uncor. at 22 mm.). Formed by distilling calcium stearate with calcium formate (Kratt, B. 13, 1417). Plates with bluish glitter, sl. sol. ether.

STEARIN v. Stearyl derivative of GLYCERIN. STEAROLIC AJID C18H12O2. Mol. w. 280. [48°]. (260°). Formed by heating bromo-oleic acid or the dibromide of clerc acid with alcoholic potash at 100° (Overbeck, J. pr. 97, 159; A. 140, 49). Needles (from alcohol), insol. water. Br forms $C_1, H_{12}Br_1O_2$ and C_1, H_{11}, Br_1O_2 [70°]. I and FeI, in CS₂ form $C_1, H_{12}I_1O_2$ [51°] (Liebermann a. Sachsse, B. 24, 4116). Potash-fusion yields myristic acid $C_{14}H_{18}O_{2}$ and an acid $C_{16}H_{19}O_{2}$ [21°] (Marasse, Z. [2] 5, 571; B. 2, 359). Alkaline KMnO, oxidises it to suberic and stearoxylic acids (Hazuraa. Grüssner, M.9,952). HNO, yielda azelaic, stearoxylic, pelargonic, and nitrosopelargonic acids (Limpach, A. 190, 294). Phenylhydrazine at 140° gives $C_{11}H_{11}CO.N_{1}H_{1}Ph$ [82°] (Holt, B. 25, 2670).—BaA .—CaA', aq.—AgA'. SFEARONE ($C_{11}H_{20}$), CO. Di-heptadecyl

ketone. [88°]. S.G. (liquid) 89 .7979 (Krafft, B. 15, 1715). Formed by heating stearic acid (9.5 g.) with P₂O₃ (5g.) at 210° (Kipping, C. J. 57,537; cf. Bussy, A. 9, 269; Redtenbacher, A. 35, 57; Varrentrapp, A. 35, 80; Rowney, C. J. 6, 97; Heintz, P. 94, 272; 96, 65). Got also by distilling the stearyl derivative of heptadecyl-urea with lime (Turpin, B. 21, 2486). Plates, sl. sol. hot alcohol. Br forms C₃,H₃,Br₂O [72°].

Oxim (C₁,H₃,)₂C:NOH. [63°]. White powder,

m. sol. hot benzene and alcohol, insol. water and alkalis (Spiegler, B. 17, 1575; Kipping, C. J.

57, 540).

STEAROXYLIC ACID C, H,2O,. 312. [86°]. Formed by the oxidation of stearolic acid (Överbeck, A. 140, 63; Hazura a. Grüssner, M. 9, 952). Plates or needles, sl. sol. cold al-cohol. BaA'₂. - AgA': crystalline powder. STEARYL GLYCERIN v. GLYCERIN.

STEEL v. Iron, vol. iii. p. 53, and DICTIONARY OF APPLIED CHEMISTRY, vol. ii. p. 360.

SIIBINE. Synonym of antimony hydride, v.

vol. i. p. 288.

STILBENE v. DI-PHENYL-ETHYLENE. STILBENE ALCOHOL v. HYDROBENZOÏN. STILBENE DIBROMIDE v. DI-BROMO-PHENYL-

ETHANE STILBENE DICARBOXYLIC ACID v. Dr.

PHENYL-MALEÏC ACID. DICHLORIDE v. DI-CHLORO-STILBENE

PHENYL-ETHANE STOICH!OMETRY. The laws of chemical

combination, and their application to chemicat calculations; v. Combination, Chemical, Laws of, vol. ii. p. 235.

STORAX. A balsam produced by Styrax officinalis, a shrub growing in the Levant. It occurs in two varieties, liquid storax and reed storax, the latter containing a large quantity of bark. Liquid storax is a brownish-yellow sticky mass containing styrene, styracin, cinnamic acid. phenylpropyl cinnamate, a little ethyl cinnamate, a substance [65°] smelling like vanilla (possibly ethyl-vanillin), and (a) and (β)-storesinol (W. von Miller, N. R. P. 24, 1; B. 9, 274; A. 188, 184 ; 189, 338).

(α)-Storesinol C_{so}H_{se}O_s i.e. C24H25(OH)20 [160°-168°]. Amorphous, v. sol. dilute KOHAq,

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but conc. KOHAq ppts. C₁₆H₃₆O₄K. Yields a muno- and tri-acetyl derivative.

(β)-Storesinol. [140°-145°]. Amorphous. Forms amorphous $C_{s_0}H_{s_0}O_4K$, which is more solwater than its (α)- isomeride.

STRONTIA. Oxide of strontium (q.v. p. 516). STRONTIUM. Sr. At. w. 87·3. Mol. w. not known. Very little known about properties. Doubtful if approximately pure Sr has yet been isolated. S.G. c. 2·4 to 2·58 (Franz, J. pr. 107, 253; Matthiessen, J. 8, 324). For emission-spectrum v. B. A. 1884. 444. H.C. [Sr,O] = 128, 440 (Th. 3, 258).

Occurrence.—Compounds of Sr are widely distributed, but not in very large quantities. SrCO₂ occurs in small quantities in all specimens or arragonite; very small quantities are found in many calc-spars, marbles, and dolomites. SrSO₄ is an ingredient of many heavy spars. SrCO₂ is found as strontianite, and SrSO₄ as celestine in a few localities; brewsterite contains Sr silicate, with silicates of Ba or Ca. Traces of SrSO₄ and SrCl₂ are found in many mineral springs, in some hard river-waters, in sea-water, and in the ashes of certain plants, especially Fucus vesiculosus.

Historical.—A mineral found at Strontian, a village in Argyleshire, in 1787, and supposed to be barium carbonate, was observed to colour fame reddish by Crawford and Cruikshank in 1790 (Mem. Manchester Soc.). The supposition made by C. a. C. that the mineral contained a new element was confirmed by Hope (T. E. 4, 3), Klaproth (Crell's Ann. 1793 [ii.] 189; 1794 (i.) 99), and Kirwan-Higgins (Crell's Ann. 1795 (ii.) 119, 205). The metal was isolated by Davy in 1808 (T. 1808. 345).

Formation.—1. By electrolysing moist SrO₂H₂ or SrCl₂ in contact with Hg and a little naphtha, and heating the amalgam so formed (Davy, T. 1808. 345).—2. By heating saturated SrCl₂Aq with Na amalgam to 90°, quickly washing the Sr amalgam so formed, drying it by filter paper, and distilling off the Hg in a stream of H (Franz, J. pr. 107, 253).—3. By heating SrO or SrO₂H₂ intimately mixed with Mg powder, a mixture of Sr with MgO is obtained (Winkler, B. 23, 125, 2647).

Preparation. - A porous clay cylinder is placed in a crucible, and SrCl,, mixed with a little NH₄Cl, is placed in the crucible and in the cell, so that when the mixture is fused the surface of it is at a higher level in the cell than in the crucible. A cylinder of sheet iron surrounding the cell serves as the positive electrode, and an iron wire passing through a tobacco pipe, the bowl of which dips under the molten mixture in the cell, serves as the negative electrode. The crucible is heated till the mixture of SrCl, and NH Cl melts, temperature being kept so that there is always a solid crust on the surface of the mixture in the porous cell. A current from 5 or 6 Bunsen cells is passed through the molten mass; Sr is separated and runs into small pieces, which are protected from the air by the solid crust of SrCl. and NH,Cl; the pieces of Sr are removed by an iron spoon and kept under petroleum.

For the preparation of pure Sr salts from strontianite v. Barthe a. Falières (Bl. [3] 7, 104).

Properties.—A vellowish-white metal, some-

Properties.—A yellowish-white metal, somewas harder than Ca or Pb; can be beaten into

thin leaves; melts at full red heat. According to Mallet (A. 190, 62), Sr is slightly volatilised at a very high temperature. Easily oxidised by exposure to air; decomposes cold water rapidly; dissolves in dilute acids, not in HNOsAq, giving salts and H. Combines directly with S and the halogens. Sr is a strongly positive metal, less positive than the alkali metals and Ca, but more positive than Mg; it is closely related to Ca and Ba, less closely to Mg; it also shows resemblances to Zn, Cd, and Hg (v. Alkaline earths, metals of the, vol. i. p. 112; and Magnesium group of elements, vol. iii. p. 163). The atomic weight of Sr has been determined (1) by determining CO₂ in SrCO₂ (Stromeyer, S. 19, 228 [1816]; Salvétat, C. R. 17, 818 [1843]); (2) by ppg. Cl from SrCl, by Ag (Rose, S. 19, 228 [1816]; Pelouze, C. R. 20, 1047 [1845]; Marignac, A. 106, 168 [1858]; Dumas, A. Ch. [3] 55, 191 [1859]); (3) by determining water in SrCl₂.6H₂O (Marignac, A. 106, 168 [1858]); (4) by transforming SrCl2.6H2O into SrSO, (Marignac, l.c.). The S.H. of Sr has not been determined directly. The V.D. of no compound of Sr has been determined.

Reactions and Combinations (v. Bunsen, A. 94, 111).—1. Exposed to air or oxygen rapidly forms SrO, or SrO, H, if moisture is present.—2. Burns brilliantly when heated in oxygen, sulphur vapour, chlorine, bromine, or iodine vapour; also in dry carbon dioxide.—3. Decomposes cold water rapidly, giving off H and forming SrO, H, —4. Dissolves rapidly in dilute hydrochloric or sulphuric acid; reacts slowly with conc. sulphuric acid; scarcely acted on by nitric acid, even when hot and conc.—5. Reduces silica and silicates when heated therewith to full redness.

Strontium, antimonate of; v. vol. i. p. 286.

Strontium, arsenates of; v. vol. i. p. 309.

Strontium, arsenite of; v. vol. i. p. 307. Strontium, borates of; v. vol. i. p. 530.

Strontium, bromide of, SrBr₂. Mol. w. not determined. By heating Sr in Br. By dissolving SrCO₂ in HBrAq and evaporating, long white needles of the hydrate SrBr₂. 6aq are obtained (Löwig, Mag. Pharm. 33, 7); S.G. 2°358 (Favre a. Valson, C. R. 77, 579); these crystals do not effloresce over H₂SO₄ (Rammelsberg, P. 55, 238), but on heating give off their water, leaving SrBr₂ as a white solid, S.G. 3°985 (F. a. V., l.c.), that melts at red heat without decomposition (R., l.c.). [Sr,Br²] = 157,700 (Th. 3, 258). Carnelley (C. J. 33, 279) gives melting-point as c. 630°. Combines with ammonia to form 2SrBr₂.NH₃ (Rammelsberg, P. 55, 288).

Strontium, chloride of, SrCl₂. Mol. w. not determined.

Formation.—1. By burning Sr in Cl.—2. By passing Cl over hot SrO (Weber, P. 112, 619).

3. By heating SrO in a stream of HCl (Chevreul, A. Ch. 84, 285).—4. By decomposing SrCO, by conc. CaCl₂Aq or MgCl₂ by heating together, then dissolving out SrCl₂ and crystallising (SrCO₃ + MgCl₂Aq = SrCl₂Aq + MgO + CO₃;

Wackenroder's Patent; v. B. 19, Ref. 633).—5. By heating a mixture of SrSO₄, CaCl₂, and charcoal with a little chalk, lixiviating, and crystallising (SrSO₄ + CaCl₂ + 4C = SrCl₂ + CaS + 4CO; Mactear, D. P. J. 262, 288).

Preparation.—Strontianite (SrCO3) is dissolved in HClAq, the solution is digested in absence of air with more SrCO, (to remove iron, &c.), poured off and evaporated to the crystallising point; the crystals of SrCl2. 6aq that separate are purified by re-crystallisation, then dried and heated to 100° until they cease to lose

weight.

Properties.—A white, crystalline powder, with a sharp, bitter taste. Melts at 825° (Carnelley, C. J. 33, 280) to a glass-like mass with an alkaline reaction. S.G. 2-96 at 0°; 2-77 at m.p. (Quincke, P. 138, 141). S. 44-2 at 0°, 48-3 at 10°, 53-9 at 20°, 60 at 30°, 66-7 at 40°, 74-4 at 50°, 83-1 at 60°, 87-5 at 65°, 88-8 at 10°, 88-8 at 10 66.5°, 89.6 at 70°, 92.4 at 80°, 96.2 at 90°, 101.9 at 100°, 109.1 at 110°, 116.4 at 118.8°; saturated SrCl₂Aq boils at 118.8° (Mulder). Gerlach (Fr. 8, 245) gives the following data:-

S.G. SrCl.Aq	P.c. SrCl.	8.G. SrCl.Aq	P.c. SrCl.
1.0453	5	1.2580	25
1.0929	10	1.3220	30
1.1439	15	1.3633	33
1.1989	20	1	

SrCl, is insol. absolute alcohol; it dissolves in aqueous alcohol in proportion to the amount of water present (Gerardin, A. Ch. [4] 5, 156). $[Sr,Cl^2] = 184,550$; $[Sr,Cl^2,Aq] = 195,690$ (Th. 3, 258). For connection between solubility in water of SrCl₂ and temperature v. Étard (Č. R. 113,

854).

Reactions and Combinations.—1. Heated to redness in water vapour, HCl is given off and SrO remains (Kraus, P. 43, 138; Kahnheim, J. 1861. 149).—2. Heated with *bromine* to 200° is partially decomposed to SrBr₂ (Potilitzin, B. 7, 733; 8, 766); the amount of decomposition varies with temperature and the relative masses of SrCl₂ and Br.—3. Combines with water. Solution of SrCO, in HClAq evaporated yields long, six-sided, hexagonal needles of the hexahydrate SrCl2. 6H2O; S.G. 1.933 at 17°. By keeping these crystals over H2SO, in vacuo for some months the dihydrate SrCl, 2H,O is formed.-4. Combines with ammonia to form SrCl₂.8NH₃ (H. Rose, P. 20, 155).

Strontium, cyanide of; v. vol. ii. p. 347.

Strontium, ferrocyanide of; v. vol. ii. p. 337. Strontium, fluoride of, SrF2. A white, crystalline powder; scarcely sol. water or HFAq. Prepared by the action of HFAq on SrO or SrCO₂ (Berzelius, P. 1, 20); also by fusing 2 parts SrCl₂ with 1 part NaF and 1 part NaCl, and lixiviating the product (Röder, Dissertation [Göttingen, 1863] 14; Feldmann, B. 21, Ref. 866 [Patent]). Poulenc (C. R. 116, 987) obtained SrF₂, as an amorphous powder, by the reaction of HFAq and SrCl₂Aq; S.G. 2.44; partly decomposed to SrO by heating in air to c. 1000°. By fusing with alkali chlorides, or with KHF2, P. obtained SrF2 in regular octahedral crystals.

Strontium, hydride of. By heating to redness a mixture of 103 parts SrO (made from SrCO₂) with 24 parts Mg powder, in an iron tube, in an atmosphere of H, Winkler (B. 22, 1975) obtained a greyish-brown powder that quickly oxidised in air to SrO₂H₂ with evolution of H, and was decomposed very rapidly by water or HClAq with violent evolution of H. Analyses

indicated that the substance might be a mixture of c. 66 p.c. SrH with c. 29 p.c. MgO, c. 4 p.c. SrO, and a very little Mg.

Strontium, hydrosulphide of, SrS₂H₂.xH₃O. By saturating SrOAq with H₂S and evaporating in vacuo over H₂SO₄; or by dissolving SrS in water, evaporating, filtering from SrO₂H₂, and evaporating over H2SO, in vacuo. Large, white crystals; melts when heated in water of crystallisation, and then gives SrS. Aqueous solution boiled gives off all H₂S and forms SrO₂H₂ (v. Rose, P. 55, 430; Berzelius, P. 6,

Strontium, hydroxide of, SrO₂H₂. Mol. w. Prepared by the action of not determined. water on SrO. Also by heating celestine (SrSO.) with charcoal, dissolving SrS so formed in water, heating with CuO or ZnO, filtering from CuS or ZnS, evaporating to dryness, and heating to

redness (cf. Barium Hydroxide, vol. i. p. 442).

A white solid; S.G. 3.625 (Filhol, A. Ch.
[3] 21, 415). Sol. water, forming a markedly alkaline solution reacting similarly to BaO, H2Aq and CaO₂H₂Aq. Scheibler (C. C. [3] 13, 33) gives the following data, showing solubility in water, expressed in terms of SrO, and the hydrate SrO₂H₂.8H₂O, dissolved:-

m		s.	m		8.
Temp.	SrO	SrO.H.SH.C	Temp.	SrO	SrO, Hg.8H,O
0°	.35	.90	55°	2.54	6.52
5	•41	1.05	60	3.03	7.77
10	•48	1.23	65	3.62	9.29
15	•57	1.46	70	4.35	11.16
20	•68	1.74	75	5.30	13.60
25	.82	2.10	80	6.56	16.83
30	1.00	2.57	85	9.00	23.09
35	1.22	8.13	90	12.00	30·78
40	1.48	3.80	95	15.15	38·8 6
45	1.78	4.57	100	18.60	47.71
50	2.13	5.46	101.2	19.40	49.75

Dry SrO₂H₂ takes up only traces of CO₂ from the air (Heyer, B. 19, 2684); but the hydrate SrO₂H₂H₂O absorbs CO₂ till it is changed to SrCO_s (H., l.c.); according to Finkener (B. 19. 2958) a basic carbonate is formed.

The octohydrate, SrO₂H₂.8H₂O, crystallises in tetragonal crystals (a:c=1:6407) from SrOAq; S.G. 1:396 at 16°; in air falls to powder, giving the monohydrate SrO2H2.H2O, which loses H₂O at 100° (v. Finkener, l.c.; Müller-Erzbach, B. 19, 2874; 20, 1628). Weisberg (B. 11, 511) says that Cl reacts with the hydrates of SrO.H.

to give SrCl₂ and Sr(ClO₃)₂.
Strontium, iodide of, SrI₂. A white solid; by dissolving SrCO, in HIAq, evaporating, and heating the hexahydrate, SrI2.6H2O, which crystallises out. S.G. 4.415; melts at red heat; strongly heated in air gives SrO and I (Croft, J. pr. 68, 420). Thomsen gives [Sr,I²,Aq]

= 143,460 (Th. 3, 258).

Strontium, manganocyanide of; v. vol. ii. p. 342.

Strontium, nitride of. By heating Sr amalgam containing from 20 to 25 p.c. Sr (prepared by electrolysis and heating the product in vacuo) in a stream of N, Maquenne (Bl. [3] 7, 366) obtained a dark-coloured compound of Sr and N to which he gave the formula Sr. N

Strontium, oxides of. Two oxides have been isolated, SrO and SrO.

STRONTIUM OXIDE SrO. (Strontium monowide. Strontia.) Mol. w. not known. Prepared by strongly heating SrO_2H_2 , $Sr(NO_3)_2$, $SrCO_3$ (Brügelmann, Fr. 29, 127; Heyer, B. 19, 2684); also by strongly heating SrI_2 in air (Croft, J. pr. 68, 420). Best prepared similarly to BaO (v. BARIUM MONOXIDE, vol. i. p. 443). A grey-white, porous, infusible solid; prepared by heating SrCO, or SrO,H, forms a mass of regular cubes with S.G. 475 referred to water at 4° (Brügelmann, l.c.). According to Moissan (C. R. 115, 1034), SrO melts to a transparent liquid at c. 3000°, and this solidifies to a crystalline mass. Thomsen gives [Sr,O] = 128,440 (Th. 3, 258). Dissolves in water (for data v. Strontium hydroxide); from this solution the octohydrated hydroxide SrO₂H₂.8H₂O separates on evaporation. By passing moist air over SrO the monohydrated hydroxide, SrO₂H₂.H₂O, is formed (Heyer, B. 19, 2684). SrO is a strongly basic oxide, reacting with acids to form salts SrX, where $X = SO_4$, $2NO_{3}, \frac{2}{3}PO_{4}, &c.$

STRONTIUM DIOXIDE SrO₂. (Strontium peroxide.) Mol. w. not known. Addition of H₂O₂Aq to SrOAq ppts. the octohydrate SrO₂SH₂O (Thénard, A. Ch. [2] 8, 312; Schöne, B. 6, 1172; cf. Conroy, C. J. [2] 11, 812). By drying this pp. at 100° SrO₂ is obtained as a white powder, having similar properties to those of barium dioxide (q.v. vol. i. p. 443). SrO₂ is not obtained by methods similar to those generally used

for making BaO2.

Strontium, oxychloride of. According to André (A. Ch. [6] 3, 66) crystals of the compound SrCl₂SrO.9H₂O(=Sr₂OCl₂.9H₂O) are deposited from a mixture of saturated SrCl₂Aq and saturated SrOAq; the crystals are very unstable in

Strontium, oxysulphide of. Schöne (P. 117, 59) obtained SrS. SrO. 12aq(Sr. OS. 12aq) by

slow oxidation of SrS,Aq.

Strontium, platinocyanide of; v. vol. ii. p.

Strontium, salts of. Compounds obtained by replacing H of acids by strontium. These compounds belong to the type SrX, where X is SO,, CO,, 2NO,, $\frac{2}{3}$ PO, &c. The Sr salts are very definite compounds; not many basic salts are known. The chief salts of oxyacids are the following: bromate and hypobromite, carbonate, chlorate, chlorite and perchlorate, chromate and dichromate, iodate and periodates, molybdate, nitrate, nitrite, and hyponitrite, selenate and selenites, sulphate, sulphite, thiosulphate, and thionates (v. Carbonates, nitrates, &c.).
Strontium, selenide of. Obtained as a white

solid, rapidly decomposing in air with separation of Se, by heating a small quantity of SrSeO, in H to dull redness; [Sr,Se] = 87,160 (Fabre, C. R.

102, 1469).

Strontium, silicofluoride of, SrSiF.2H.O. Four-sided prisms; S.G. 2.999; loses 2H₂O when gently warmed; prepared by dissolving SrCO, in H.SiF.Aq and evaporating (Berzelius, Lehrbuch [5th edit.], 3, 385]. Fresenius (Fr. 29, 143) gives S. 3·2 at 15°; S. in alcohol (50 p.c. by vol.) .06 at 15°.

Strontium, sulphides of. Three sulphides

have been isolated: SrS, SrS,, and SrS,

STRONTIUM MONOSULPHIDE SrS. Formed by heating together Sr and S in the ratio 87:32; Needles, m. sol. boiling water and hot alcohol.

also by reducing SrSO, by charcoal; also by passing CS2 vapour mixed with H, H2S, or CO2 over red-hot SrCO₃ (cf. BARIUM MONOSULPHIDE, vol. i. p. 444). A white powder, which slowly becomes yellowish when exposed to air. Does not phosphoresce (Schöne, P. 117, 59). For phosphorescence of SrS mixed with small quantities of other salts v. Becquerel (C. R. 107, 892). Sabatier (A. Ch. [5] 22, 5) gives [Sr,S] = 49,600. SrS does not combine with S when heated therewith, but polysulphides are formed by boiling an aqueous solution of SrS with S (Schöne, P. 117, 59). SrS dissolves in water; on crystallising SrO₂H₂ separates and SrS₂H₂ remains in solution (H. Rose, P. 55, 430).

STRONTIUM TETRASULPHIDE SrS4.xH2O. The hexahydrate, SrS, 6H₂O, is obtained, as a reddish crystalline solid, by boiling 100 parts SrS in water, and evaporating the solution in vacuo at a temperature not higher than 16°. solution at 20°-25° the dihydrate SrS, 2H2O separates. An aqueous solution of SrS, oxidises in air to the oxysulphide SrO.SrS. 12aq, and then to SrS₂O₃, while S separates and some SrCO₃

is formed (Schöne, P. 117, 59).

STRONTIUM PENTASULPHIDE SrS, xH2O. An amorphous, yellowish, hygroscopic solid; formed by evaporating a solution of SrS, saturated with S in the cold. After drying at 100°, CS, dissolves out the S, leaving SrS, (Schöne,

l.c.).
Strontium, sulphydrate of; v. Strontium

HYDROSULPHIDE, p. 515.

Strontium, sulphocyanide of; v. vol. ii. p. M. M. P. M.

STROPHANTHIN $C_{s_1}H_{4s}O_{12}$ (A.); $C_{2o}H_{s_4}O_{1e}$ (F.). $[a]_{b}=30^{\circ}$. Extracted by alcohol at 70° from the seeds of Strophanthus hispidus (Gallois, C. R. 84, 261; Fraser, Ar. Ph. [3] 3, 229; Pr. E. 124, 370; Ph. [3] 18, 69; Elborne, Ph. [3] 17, 743; 18, 219; Gerrard, Ph. [3] 17, 923; Arnaud, C. R. 107, 179; Gley, C. R. 107, 348). White micaceous crystals, sol. water, v. sol. alcohol, insol. ether. Dextrorotatory. Resembles ouabain in toxic effects. Decomposed by boiling dilute acids (even H₂S) into glucose and crystalline strophanthidin, which yields a resin on further boiling with dilute H.SO. (F.). H₂SO₄ and a trace of FeCl₃ gives a reddish-brown pp., changing to emerald-green (Helbing, Ph. [3] 17, 924).

STRUTHIIN v. SAPONIN. STRYCHNIC ACID C20H22NO(NH).CO2H. Formed from strychnine, alcohol, and Na at 50°-55° (Tafel, A. 264, 50). When heated in a current of H at 190° it changes to strychnine. K2Cr2O, and H2SO4 give a brown-red colour or

Nitrosamine C20H22NO(N.NO).CO2H. Gelatinous pp. Exhibits Liebermann's reaction.

Methylo-iodide $C_{20}H_{22}N(MeI)O(NH).CO_2H.$ Formed from methyl-strychnine and HI (Tafel, A. 264, 55). Colourless needles (containing aq), v. sl. sol. cold alcohol and cold water.—NaA'aq: v. e. sol. water, m. sol. alcohol. Converted by MeI into crystalline C20H22N(MeI)(NMe).CO2Me.

Methyl-strychnic acid. Methylo-iodide. C,H,N(MeI)O(NMe).CO,H. Formed from dimethyl-strychnine and HI (Tafel, A. 264, 58).

Deoxystrychnic acid C₂₀H₂₆N(NH).CO₂H aq. Formed by heating deoxystrychnine with alcoholic NaOEt at 180° (Tafel, A. 268, 253). Needles (containing aq) (from dilute H.SO.) or anhydrous prisms (from alcohol), sl.sol. hot water. Quickly converted by acids into deoxystrychnine. Yields a nitrosamine.

 $C_{20}H_{22}NO(NH).CO_2H.$ acid Isostrychnic Formed from strychnine, Na, and alcohol at 70°. Got also by heating strychnine with barytawater at 140° in absence of air. needles. K₂Cr₂O₇ and dilute H₂SO₄ form a brown liquid on heating, and a brown pp. with excess of K2Cr.O7.-HA'HI aq: prisms, m. sol.

cold water and alcohol, insol. ether.

Nitroso- derivative

NO.C₂₀H₂₁NO(NH).CO₂H aq. Sol. alkalis, but ppd. by CO₂. Yields a nitrosamine, which gives Liebermann's reaction with phenol and H2SO4. -HA'HCl: needles, insol. water.

Methylo-iodide C₂₀H₂₂N(MeI)O(NH).CO₂H. Crystalline, m. sol. hot water.—NaA': minute needles. Converted by methyl iodide at 100° into crystalline

C₂₀H₂₂N(MeI)O(NMe).CO₂Me 2aq.

Methyl-isostrychnic acid C20H22NO(NMe).CO2H. From isostrychnic acid hydroiodide and McI at 100° (Tafel, A. 268, 240). Small colourless prisms (containing 2 aq). Its alkaline solution is not coloured by air (difference from isostrychnic acid).

Methylo-iodide

[271°-275°]. C₂₀H₂₂N(MeI)O(NMe).CO₂H aq Formed from methyl-strychnine methylo-iodide and HI (Tafel, A. 264, 77). Needles, m. sol. hot water and hot alcohol, insol. ether.

Nitroso-methyl-isostrychnic acid

NO.C., H21ON(NMe).CO2H. Formed from the preceding acid, alcohol, HCl, and amyl nitrite. Green needles (from benzene). When heated with benzoic aldehyde and ZnCl2 it gives a green mass, which dyes like malachite green. Diazo-benzene chloride forms a brownish-orange

STRYCHNINE C, H, N, O, Mol. w. 334. [221°] (Blyth); [284°] (Claus a. Glassner, B. 14, 773); [268°] (Loebisch a. Schoop, M. 6, 858); 778); [269°] (Stoehr). (270° at 5 mm.). S.G. ¥ 1·359 (Clarke, Am. 2, 174). S. 015 in the cold; 04 at 100°. S. '014 in the cold (Dragendorff, J. 1865, 739); '025 at 14'5° (Crespi, G. 13, 175). S. (alcohol) '3 at 8°; 1.8 at 78°. S. (95 p.c. alcohol) '936. S. (ether) '08. S. (benzene) '607. S. (isoamyl alcohol) '53 at 12°; 4.3 at 99°. Occurs in nux vomica, in St. Ignatius beans, in the wood of Strychnos Colubrina, in Strychnos Tieute, and in other species of Strychnos (Pelletier a. Caventou, A. Ch. [2] 10, 142; 26, 44; Pelletier a. Dumas, A. Ch. [2] 24, 176; Liebig, A. 47, 171; 49, 244; Regnault, A. Ch. [2] 68, 113; Gerhardt, Rev. Scient. 10, 192; Nicholson a. Abel, C. J. 2, 241; Henry, Ph. 8, 401; Corriol, Ph. 11, 492; Robiquet, Ph. 11, 580; Henry, jun., J. Ph. 16, 752; Berlekom, Z. [2] 2, 443; Shenstone, Ph. [3] 8, 445). Occurs also in the bark of Erythrophlaum guineense, from which the pigmies of Central Africa prepare their arrow-poison (Holmes, Ph. [3] 21, 921).

Preparation. - 1. The nuts (1 kilo) are softened by steam, dried, powdered, and digested with alcohol (5 kilos of S.G. 856) acidulated with

H₂SO₄ (45 g.). Lime is added to the filtrate: and, after standing, the decanted liquid evaporated; the residue dissolved in dilute acid; and the filtrate ppd. by ammonia. The strychnine is separated from brucine by crystallisation from alcohol (Henry) .- 2. The dilute alcoholic extract of nux vomica is evaporated to a small bulk and ppd. by lead acetate. The filtrate is mixed with magnesia and left for a few days. The pp. is dissolved in alcohol of 83 p.c. and evaporated to crystallisation. Strychnine is first deposited, and afterwards brucine. The bases are further purified by crystallisation of their nitrates, the nitrate of strychnine separating first.

Properties.—Trimetric prisms, permanent in the air and not altered by light. Lævorotatory, the rotation varying greatly with the nature of the solvent (Hoorweg, A. 166, 76; Tykociner, R. T. C. 1, 146). The dispersive power is constant (Grimbert, J. Ph. [5] 16, 295). At 169° it forms a sublimate of minute needles (Blyth). Very bitter. Alkaline in reaction. Strychnine is a violent poison, acting on the spinal cord and producing convulsions (Lovett, J. Physiol. 9, 99). ·03 g. may be fatal. Strychnine dissolves with difficulty in acids, the solubility being greater the more dilute the acid; when a conc. solution of a strychnine salt is slightly acidified a pp. is formed which dissolves in excess of acid, forming a solution which yields a pp. on dilution. Thus H2SO4 added to a conc. solution of strychnine sulphate ppts. the acid sulphate B'H₂SO₄, the mother-liquor retaining 113 p.c. of salts. HCl added to a solution of strychnine hydrochloride ppts. B'HCll aq in needles, the mother-liquor retaining 413 p.c. of salt (Hanriot a. Blarez, C. R. 96, 1504). Strychnine is readily soluble in dilute hypophosphorous acid (Jones, Ph. [3] 20, 256). KCy added to a solution of a salt of strychnine ppts. the pure base (Flückiger, N. J. P. 38, 138; Weith, B. 4, 527). A solution of ICl gives a yellow pp. v. sol. HClAq, from which it crystallises on cooling (Dittmar, B. 18, 1612).

Reactions.—1. Dry distillation yields carbazole, H, C₂H₂, and C₂H₄ (Loebisch a. Schoop, M. 7, 614).—2. Distillation with lime yields (\$\beta\$)-methylpyridine, methyl-indole (scatole), C2H4, NEtH2, and NH, (Stochr, B. 20, 810, 1108, 2729; J. pr. [2] 42, 405).-3. Distillation with soda-lime gives carbazole, scatole, and (β) -methyl-pyridine (Loebisch a. Malfatti, M. 9, 628).—4. Distillation with KOH gives indole (Goldschmidt, B. 15, 1977) and butyric acid (Loebisch a. Schoop, M. 7, 93).-5. Distillation with zinc-dust at 400° in vacuo yields solid C21H22N2O, which dissolves in alcohol with blue fluorescence, and gives no colour with K₂Cr₂O₇ and H₂SO₄. Zinc-dust at a red heat gives H, C₂H₄, C₂H₂, NH₂, and carbazole (Loebisch a. Schoop, M. 7, 609), and (γ) -lutidine (Scichilone a. Magnanini, G. 12, 444). 6. Alkaline KMnO, yields oxalic acid, CO, NH, and s crystalline acid [195°] (Hoogewerff a. Van Dorp, R. T. C. 2, 181). Half the nitrogen is given off as ammonia (Wanklyn a. Chapman, C. J. 21, 161).-7. KMnO, in dilute acid solutions yields an amorphous acid C₁₁H₁₁NO₂ aq (?) (Plugge, R. T. C. 2, 270), which yields amorphous C₁₁H₁₀AgNO₂ aq (Hanriot, C. R. 96, 1671).—8. By treatment with CrO₂ and H₂SO₄ it yields the same oxidation product C₁₆H₁₈N₂O₄ as that ob-

tained from brucine under the same treatment (Hanssen, B. 18, 1917).-9. PCl, (1 pt.) acting on strychnine hydrochloride (8 pts.) under boiling chloroform yields a compound crystallising from benzene in needles [224°], possibly penta-chloro-strychnine (Stoehr, J. pr. [2] 42, 414).—10. Boiling HNO, forms cacostrychnine and picric acid (Shenstone, C. J. 47, 142); nitric acid forms 20 p.c. of di-nitro-di-oxy-quinoline carboxylic acid C₁₀H₁N₂O₃, which yields KA', is reduced by SnCl₂ to C₁₀H₂N₃O₄ (a body having acid and basic properties), and when heated with water is split up into CO₂ and dinitro-di-oxy-quinoline (Tafel, B. 26, 353).—11. Baryta water in sealed tubes at 140° forms two bases, C, H, N,O, and C₂,H₂₈N₂O₃, both crystallising from water (Gal a. Etard, C. R. 87, 362).—12. Alloxan added to a solution of strychnine in a saturated solution of SO₂ forms small colourless prisms of B'C, H2N2O, H2SO, aq (Pellizzari, A. 248, 150). 13. Iodoform (5 g.) and strychnine (12 g.) dissolved in hot alcohol (500 c.c.) deposit on cooling crystals of B', CHI, sol. ether and chloroform, but decomposed by light, hot water, and dilute acids (Lextreit, C. R. 92, 1057).-14. Chloro-acetic acid at 180° forms C₂₂H₂₄N₂O₄, a base which crystallises in silky needles, v. sol. hot water and alcohol, insol. ether (Roemer, Z. 1871, 435). It yields $(C_{23}H_{24}N_2O_4)$ PtCl₄. — 15. AcCl forms B'AcCl, which gives (B'AcCl) PtCl₄ (Konrad, J. 1874, 876).—16. Chloro-acetone at 135° yields $B'C_3H_5ClO$, S. 6.5 at 15°, which yields (B'C,H,OlO),PtCl, 2aq and B'C,H,OSO,H 112aq (Konrad).—17. Aqueous SO₂ and alloxan give colourless prisms of B'SO₂HC₄H₂N₂O₄ aq (Pellizzari, G. 18, 329).

Detection.—1. H₂SO₄ forms a colourless solution which with K_Cr2O, gives a blue colour, changing through violet and red to yellow (Otto, A. 60, 278). A drop of nitric acid and PbO₂ or MnO₂ may be used instead of K₂Cr₂O₇ (Marchand, B. J. 24, 400; Davy, A. 88, 402). KMnO₄ and Ag₂O act in like manner (Lyman, Fr. 12, 126). The colour with H2SO, and K2Cr2O, is not shown in presence of brucine (Hanriot, C. R. 97, 267), and is interfered with by quinine, morphine, and sugar (Brieger, J. 1850, 617; Vogel, J. 1853, 686; Flückiger, Fr. 28, 102).—2. A minute quantity of KClOs added to a warm solution in dilute HNO, gives a scarlet colour, changed by ammonia to brown, and then, on evaporation to dryness, becoming green (C. L. Bloxam, C. N. 55, 155).-3. Ammonium vanadate in H2SO4 gives a blue colour, turned red by potash (Mandelin, Fr. 23, 240).-4. Chlorine water gives a pp.—5. Strychnine can be extracted from beer containing the poison by shaking with animal charcoal, the charcoal being subsequently extracted with alcohol (Graham a. Hofmann, A. 83, 39; cf. Wagner, Fr. 6, 387).—6. Obtained from acid liquids by adding ammonia and immediately shaking with chloroform (Cloetta, Fr. 5, 265; Gray, Fr. 12, 125).-7. A liquid containing strychnine sulphate may be shaken with benzene without loss of strychnine, but if the liquid be neutralised by magnesia the free strychnine can be extracted by benzene (Dragendorff, Z. [2] 2, 27).—8. A 1 p.c. solution of strychnine strongly acidified by HCl is ppd. by K.FeCy, while brucine remains in solution (Holst a. Beckurts). This may be used as a volumetric

method. Strychnine and brucine may also be separated by crystallisation of their picrates (Gerock, Ar. Ph. [3] 27, 158).

Salts.—B'HCl 12aq (Regnault, A. 26, 17). Trimetric crystals; a.b.c = 484:1:466 (Stoehr, J. pr. [2] 42, 399). Neutral in reaction. S. 2 at 22°. Leevorotatory. $[a]_r = -28^\circ$.—B'HAuCl. Orange crystals (from alcohol) (Nicholson a. Abel, C. J. 2, 241; A. 71, 84). Decomposed by boiling water (O. De Coninck, *Él.*[2] 45, 131).—B'₂H₂PtCl₆: golden scales (from alcohol). S.G. 185 1.779 (Clarke, Am. 2, 175).—B'2H2PtCle aq (Schmidt, A. 180, 295).—B'₂H₂ZnCl₄ aq: prisms (Grafinghoff, Bl. [2] 4, 391).—B'₂H₂CdCl₄ (Galletly, N. ed. P. J. 4, 94).—B'HHgCl₂. Crystals (from alcohol).—B'HgCl₂: insol. alcohol, water, and ether.—B'₂H₂SO₂2HgCl₂.—B'₂H₂PdCl₃: dark-insol. brown needles (from alcohol).—B'HCl2HgCy2: tables (Brandis, A. 66, 268).—B'HgCy2: small prisms, sl. sol. water and alcohol, insol. ether. -B'HBraq: needles, sol. water and alcohol.-B'HI aq: plates or needles, almost insol. cold water.—B'HI. S. 007 at 15°. Reddish-brown pp. got by adding a solution of I in HIAq to a salt of strychnine. Crystallises from alcohol in dark-brown needles with bluish metallic lustre. Dichroic (brown and yellow) in polarised light. M. sol. hot alcohol, sl. sol. chloroform, nearly insol. CS₂ (Tilden, C. J. 18, 99; Herapath, Pr. 8, 149; Bauer, Ar. Ph. [3] 5, 289; Jörgensen, o, 14v; Bauer, Ar. Fh. [5] 5, 289; Jörgensen, A. Ch. [4] 9, 115).—B'HHgI₃: yellow tables (from alcohol) (Groves, C. J. 2, 97).—B'H₂F₄ 2aq: prisms (Elderhorst, A. 74, 77).—B'HNO₃: needles.—B'HClO₄aq: prisms, sl. sol. cold water (Boedeker, A. 71, 62).—B'H₂SO₄ 2aq: needles.—R' H SO. water (Boetelet, A. 11, $\frac{1}{2}$, $\frac{1}{2}$ Rammelsberg, B. 14, 1231; cf. Baumhauer, C. J. 44,485).—B'2H2CrO4: lemon-yellow needles, sl. sol. water and alcohol.—B'2H2S2O34aq. S. 9 in the cold. Formed by atmospheric oxidation of a mixture of strychnine, alcohol, and ammonium sulphide (How, C. N. 18, 232).— B'2H2Sa. Formed from strychnine and alcoholic ammonium polysulphide (Hofmann, B. 1, 81; 10, 1087) and by atmospheric oxidation of a solution of strychnine in alcoholic H2S (Schmidt, B. 5, 1267; 10, 1288; A. 180, 288). Orange needles, insol. water, alcohol, and ether.—
B'H₃PO₄ 2aq. S. 19. Needles (Anderson, A. 66, 56).—B'₂H₃PO₄ 9aq: rectangular tables.— B'H, AsO, laq. S. 7 in the cold; 20 in hot water. —B'HAsO₂: efforescent cubes. S. 3 in the cold; 10 at 100°.—B',H,FeCy, 4aq. Crystalline powder (Holst a. Beckürts, Ar. Ph. [3] 25, 818; cf. Brandis, A. 66, 257). By atmospheric oxidation in presence of water it is converted into strychnine and the ferricyanide (Beckürts, B. 18, 1235).—B'H,FeCy,: white powder, sol. hot water. Acid in reaction.—B'sH,FeCy, 6aq: golden prisms.—B's(H,CoCy,) 4aq.—B's(H,NiCy,), 8aq (Lee, B. 4, 789).—B',H,PtCy, 2aq: needles (from (Hee, D, 4, 769).— D_2H_2 FtOy, $2\pi q$. Receives (1801) alcohol).—Oxalate B'_2 H_2C₂O₄: flat needles.— B'_2 H_2C₂O₄ $\frac{1}{2}$ aq. — B'H_2C₂O₄. — B'HSOy: long silky needles.—B'HSOyfr(SCy), $\frac{1}{2}$ NH₂. Red pp., v. sl. sol. hot water (Christensen, J. pr. [2] 45, 367).— B'_2 2H₂Pt(SCy)₆. Red crystals (Clarke and Owens, Am. **5**, **351**). — Cyanurates: B'₂H₁O₃C₄N₃ aq. [287°]. Prisms, sl. sol. water, m. sol. alcohol. — B'H₄O₅C₄N₃ aq. [295°]. Needles (Claus, J.pr. [2] 38, 228). — Nitroprusside. S. ·12 (Davy, Ph. [3] 11, 756). Succinate B'₂C₄H₆O₄ 6\frac{1}{2}aq (Crespi, C. 13, 176). — Tartrates: B'₂C₄H₆O₆. — B'₂C₄H₆O₆ 4aq; white efflorescent needles (Arppe, J.pr. 53, 331). — B'C₄H₆O₆ 7aq (Pasteur, A. Ch. [3] 38, 437). — B'C₄H₆O₆ 3aq. — B'C₄H₆(SbO)O₆. Needles (Stenhouse, A. 129, 25). — Malate B'₂C₄H₆O₅ 3\frac{1}{2}aq. — Pyrotartrate B'₂C₅H₆O₄ 3\frac{1}{2}aq. — Mucate B'₂C₄H₁₆O₈. Long needles (Ruhemann a. Dufton, C. J. 59, 754).

Di-hydrate B'2aq. Strychnol. Formed by warming strychnine with alcoholic NaOEt, adding water, and evaporating. The brown oily layer that forms is allowed to solidify, dissolved in water, and ppd. by CO2. The pp. is dissolved in NH, Aq and re-ppd. by expelling NH, by air (Loebisch a. Schoop, M. 7, 83). The same body is got by heating strychnine (15 g.) with BaO_2H_2 (30 g.) for 12 hours at 140° (Tafel, B. 23, 2737). Minute wedge-shaped needles, sl. sol. cold water, insol. alcohol and ether. Darkens at 150°. Sol. dilute alcohol and solutions of ammonium salts; sol. acids. Converted into strychnine on warming with dilute acids. Dilute HNO, and a little NaNO, give a crimson colour. Bromine water gives a pale-purple colour. Does not give the strychnine reaction with H2SO4 and K2Cr2O7. On heating at 100° with NaOMe and MeI it is converted into C21H22N2O2MeaI 2aq, crystallising in needles.

Tri-hydrate v. Reaction 10.

Pentahydrate B'5aq. [215°]. Got by heating strychnine with alcoholic NaOEt (Tafel). Needles, more soluble than the dihydrate. Becomes very hygroscopic when dried in vacuo.

Benzoyl derivative C₂₁H₂₁BzN₂O₂. V. sl. sol. water (Schützenberger, A. 108, 353; C. R.

47, 233).

Methylo-iodide B'MeI. S. 5 in the cold (Stahlschmidt, P. 108, 513). Poisonous (Crum Brown a. Fraser, Tr. E. 25). Formed by heating with MeI in sealed tubes. Pearly plates. Yields B'MeI, (Jörgensen, J. pr. [2] 3, 157) and the salts B'MeBr, B'MeCl 2aq, B'₂Me₂PtCl, B'MeAuCl, B'₂Me₂SO₄5aq, B'MeHSO₄, B'MeH_PO₄2aq, B'MeCl2½HgCl₂, B'MeNO₅, and B'₂Me₂FeCy₄.

Methylo-hydroxide B'MeOH or $C_{20}H_{22}O(NH) \ll_{CO}^{NMe} > 0$. Formed from B'MeI

by successive treatment with Ag, SO₄ and baryta (Tafel, B. 23, 2733; A. 264, 62). Colourless crystals (containing 4aq), v. sol. alcohol, but a product containing less water soon separates. Poisonous. Gives the same colour-reactions as strychnic acid. Alcohol and Na yield a compound [158°]. MeI forms B'MeIMeOH crystallising in plates, decomposed at 280°.

Isomeride of the methylo-hydroxide C₂₂H₂₆N₂O₂7aq. Formed from the methyloiodide of isostrychnic acid, Ag, and Ag₂O at 40° (Tafel, A. 264, 81). Colourless needles; v. e. sol. hot water and alcohol, insol. ether.

Methyl-strychnine methylo-hydroxide C₂₃H₂₈N₂O₂ i.e. C₂₀H₂₂O(NMe) ≪ CO O. drate C₂₁H₂₁ClN₂O₂ Saq or potash.—B'₂H₂SO₄ 7aq.

ment with Ag₂SO₄ and baryta. Got also by the action of Ag₂O on the methylo-iodide of methylstrychnic acid (Tafel, B. 23, 2735; A. 264, 66). Prisms (containing 6aq), v. sol. hot water and alcohol, nearly insol. ether. Gives a blood-red colour with HNO₂, CrO₃, or FeCl₃. Forms with acids salts which differ from those prepared by the action of silver salts on B'MeIMeOH. Forms a crystalline nitroso-derivative, v. sol. water and warm alcohol, forming a yellow solution changed by NaOH to brownish-red and olive-green. HI yields the methylo-iodide of methyl-strychnic acid. Diazobenzene sulphonic acid gives a brownish-red dye.—Hydroiodide. S. 7. Plates.—Methyloiodide C₂₄H₂₁N₂IO₃. S. 1. Needles.

Isomeride of methyl-strychnine methylo-hydroxide C₂₂H₂₂N₂O₂3aq. Formed from the methylo-hydroxide of methyl-isostrychnic ether (Tafel, A. 264, 82). Crystals, v. sol. hot water and alcohol, insol. ether. Yields a dye with diazo-benzene sulphonic acid.

Ethylo-iodide B'Etl. S. 6 at 15°; 2 at 100°. Formed by heating strychnine with alcohol and Etl at 100° (How, A. 92, 338). Foursided prisms. Not attacked by KOHAq. Gives the strychnine reaction with K₂Cr₂O, and H₂SO₄. Converted by moist silver oxide into the base B'EtOH 2aq, which yields the salts B'EtNO₃, B'2£tPtCl₆, B'EtHCO₃, B'EtHCrO₄ aq, B'EtCy [105°] (Claus a. Merck, B. 16, 2748), and B'Etl₄ (Jörgensen, A. Ch. [4] 11, 115).

Isoamylo-chloride B'C₅H₁₁Cl ½aq. Formed by heating strychnine with isoamyl chloride (How, Tr. E. 21, 1, 27). Prisms. Yields B'C₅H₁₁OH, B'C₅H₁₁I₅, B'C₅H₁₁I₆, crystallising in black prisms (Jörgensen, J. pr. [2] 3,

145), and B'EtHCrO,

Bromo-ethylo-bromide
B'C₂H₄Br₂.
Formed by heating strychnine with ethylene bromide and alcohol (Ménétriès, J. pr. 85, 230).
White crystals, sl. sol. cold water and alcohol. Converted by AgNO₃ into B'C₂H₄Br.NO₃ and by Ag₂SO₄ into B'C₂H₄Br.SO₄H, whence baryta produces alkaline B'C₂H₄Br.OH.—(B'C₂H₄Br)₂PtCl₃.
—B'C₂H₄Brl₂: shining brown lamins.—Moist Ag₂O converts B'C₂H₄Br₂ into B'C₂H₂OH, which is alkaline and is converted in aqueous solution by chlorine into C₂₃H₂₁Cl₂N₂O₂.—B'C₂H₄CrO₄H.
—(B'C₂H₃Cl)₂PtCl₄.

Chloro-ethylo-hydroxide B'ClC₂H₄.OH.
Got by heating strychnine with glycolic chlorhydrin at 150° (Messel, A. 157, 7). Silky needles
(containing aq). Yields (B'C₂H₄OH)₂SO₄2sq,
which is converted by baryts into crystalline
(B'C₂H₄OH)OH 2½ aq.—(B'C₂H₄OH)₂PtCl₈.

Benzylo-chlorideB'C₂H₄OI. [263°]. Prisms

Benzylo-chloride B'C,H,Cl. [263°]. Prisms (containing aq). Yields (B'C,H,Cl),PtCl, [216°], B'C,H,NO₃ [c. 264°], (B'C,H,),Cr,O₇, and B'C,H,SCy [237°] (Garzarolli, M. 10, 1).

Bensylo-hydroxide B'C,H,OH. Plates. Chlorostrychnine C₂₁H₂₁ClN₂O₂.[a]_D = -105°. Formed, together with di- and tri-chlorostrychnine, by passing Cl into a solution of strychnine hydrochloride (Laurent, A. Ch. [3] 24, 312; A. 69, 14; Richet a. Bouchardat, C. R. 91, 990). Crystalline, v. sol. alcohol, ether, and chloroform. Chlorostrychnine forms a hydrate C₂₁H₂₁ClN₂O₂ 8aq on heating with alcoholic potash.—B'₂H₂SO₄7aq.

Dichloro-strychnine C₂₁H₂₀Cl₂N₂O₃. Needles i (from alcohol).

(a)-Tri-chloro-strychnine C₁₁H₁₈Cl₁N₁O₂.
Formed by saturating a solution of strychnine hydrochloride with Cl. Minute crystals (from alcohol). Yields a trihydrate which is not very poisonous.

(B)-Tri-chloro-strychnine C₂₁H₁₈Cl₂N₂O₂. Formed by heating strychnine hydrochloride with PCl₃ and chloroform (Stoehr, J. pr. [2] 42, 412; B. 20, 813). Microcrystalline powder (from alcohol). Gives the strychnine reaction with K₂Cr₂O, and H₂SO₄.—B'HCl: plates (from dilute alcohol).—B'H₂SO₄. Plates, m. sol. water. Bromo-strychnine C₂₁H₂₁BrN₂O₂. [222°].

Bromo strychnine $C_{21}H_{21}BrN_2O_2$. [222°]. Formed by adding Br (2 at.) to a solution of strychnine hydrochloride (1 mol.), ppg. by alkali, and recrystallising from alcohol (Shenstone, C. J. 47, 189; Beckürts, B. 18, 1236; Loebisch a. Schoop, M. 6, 855). Trimetric crystals; a:b:c=1·459:1:1·195 (Miers, C. J. 47, 144).—B'HCl.—B'HBR.—B'HNO.—B'_HLSO,7aq.

B'HCl.—B'HBr.—B'HNO,—B'2H2SO,7aq.

Methylo-iodide B'Mel. Pearly plates,
sol. alcohol. Converted by moist Ag20 into

B'MeOH4aq.

Dibromide C₂₁H₂₁Br₂N₂O₂. Got by adding Br to a solution of strychnine at 40°. Bright

yellow crystalline powder.

Di-bromo-strychnine $C_{21}H_{21}Br_2N_2O_2$. Got by adding Br to strychnine in chloroform. Trimetric crystals, v. sol. chloroform and dilute

alcohol. -- B'HCl: needles.

Nitro-strychnine C₂₁H₂₁(NO₂)N₂O₂. [225°]. Formed by adding strychnine nitrate to H₂SO₄ (Loebisch a. Schoop, M. 6, 845; 7, 59). Golden plates (from alcohol). Gives no colour with H₂SO₄ and K₂Cr₂O₇. Dissolves in acids and alkalis. On heating with alcoholic potash it is converted into xanthostrychnol C₂₁H₂₁N₃O₄, which forms ruby-red crystals (containing 2aq) and gives salts with acids and bases.—B'HCl.—B'₂H₂PtCl₈.—B'₂KOH: red deliquescent needles.—B'₂BaO₂H₂ 2aq.—Ag₂C₂₁H₁₀N₃O₄: unstable pp. (a)-Di-nitro-strychnine C₂₁H₂₀(NO₂)₂N₂O₂.

[226°]. Formed by passing nitrous acid gas into an alcoholic solution of strychnine nitrate (Claus a. Glassner, B. 14, 774). Orange-yellow plates. Converted by dilute HNO₃ into cacostrychnine.—B'HNO₃: yellow nodules (from

acetone), v. sl. sol. hot water.

(β)-Ďi-nitro-strychnine. Formed by dis solving strychnine (60 g.) in fuming HNO, (800 g.) at -10° (Hanriot, C. R. 96, 585). Transparent amber-yellow prisms (from chloroform-alcohol), sol. hot water and alcohol, v. sol. chloroform. Decomposes at about 202°.—B'HCl.—Nitrate: plates (from hot water).

Amido-strychnine v. vol. i. p. 184. Di-amido-strychnine, ibid. Cacostrychnine v. vol. i. p. 654.

Deoxystrychnine C₂₁H₂₈N₂O i.e. C₂₀H₂₆N $\stackrel{\mathbf{N}}{\stackrel{\mathbf{CO}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}{\overset{\mathbf{O}}}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}}}}}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}}}}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}}}}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}}}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}}}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}}}}}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}}}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}}}}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}}}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}}}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}}}}}}{\overset{\mathbf{O}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}}}}{\overset{\mathbf{O}}}}$

[172°]. Formed by boiling strychnine with HI and P (Tafel, A. 268, 245). Crystalline mass (containing 3aq), almost insol. water, sl. sol. ether and benzene, v. sol. alcohol. Melts, when hydrated, at 75°. Poisons like strychnine. Very bitter. Very little produces vomiting. Leevo-rotatory, a 10 p.c. solution rotating 1.6° in a 100 mm. tube. Its solution in H₂SO, is coloured bluish-violet by K₂Cr₂O₇, quickly becoming brown.

Its salts are more soluble than those of strychnine.—B'HI aq. Groups of prisms.—B'₂H₂Or₂O₇. Thin yellow needles.—B'₂H₂PtCl₈.

Methylo-iodide B'MeI. Crystals. S. c. 10

at 100°.

Strychnine sulphonic acid $C_{s_1}H_{s_1}(SO_sH)N_2O_s$. Formed by dissolving dry strychnine sulphate in fuming H_sSO_s (containing 30 p.c. anhydride extra) in the cold and allowing to stand a fortnight (Loebisch a. Schoop, M. 6, 857). Deliquescent mass. Does not give the strychnine reaction with $K_2Cr_sO_s$ and H_sSO_s . Strychnine sulphonic acid. Formed by

Strychnine sulphonic acid. Formed by heating strychnine with H₂SO₄ (2 pts.) at 100° (Stoehr, B. 18, 3429; Guareschi, G. 17, 109). Amorphous powder, v. sl. sol. water and alcohol, sol. dilute acids and alkalis. Does not give a colour with K₂Or₂O₇ and H₂SO₄. Not poisonous.—NH₄A': v. sol. water, insol. alcohol.—KA': m. sol. water.—BaA'₂: amorphous pp.

Strychnine disulphonic acid

C₂₁H₂₀(SO₂H)₂N₂O₂. Got by heating strychnine with H₂SO₄ and SO₅ at 150°. Amorphous, v. sol. water, v. sl. sol. benzene, insol. alcohol and ether.—Na₂A"6aq.—BaA": plates, sol.

water.—BaA"7aq.—Ba(HA"),: yellow powder.

STRYPHNIC ACID C,H₂N₃O₂. Got by adding
HOAc to a mixture of uric acid and KNO₂ and
evaporating (Gibbs, B. 2, 341). Small paleyellow crystals, sol. hot water. Sodium-amalgam
forms crimson crystals.— KA' 1½aq: yellow
needles.—MgA'₂6aq.—CaA'₂2aq.—SrA'₂6aq.—
BaA'₂2aq.—PbA'₂.—Pb₂A'₂O 3aq.

STYCERIC ACID v. PHENYL-GLYCERIC ACID.

STYCERIC ACID v. PHENYL-GLYCERIC ACID.

STYCERIN CHPh(OH).CH(OH).CH₂OH.

Formed by heating CHPhBr.CH₂Br.CH₂OH (v.

DI-BROMO-PHENYL-PROPYL ALCOHOL) with 30 pts.

water and some AgOAc at 150°-165° (Grimaux,

J. 1873, 404). Yellow gummy mass with bitter
taste. V. sol. Aq and alcohol, nearly insol. ether.

STYPHNIC ACID v. TRI-NITRO-RESORCIN.

STYRACIN v. Cinnamyl ether of CINNAMIO

ACID.

STYRENE C_aH_a i.e. C_aH_a .CH:CH $_2$. Styrol. Styrolene. Cinnamene. Phenyl-ethylene. Mol. w. 104. (144°) (Schiff, A. 220, 93). S.G. $\frac{3}{2}$ ° 9074. $\mu_D=1$ -541 (Madan, C. J. Proc. 1, 107). S.V. 131 (Schiff). R_{∞} 58·1 (Nasini a. Bernheimer, G. 1, 93). Occurs in liquid storax (Bonastre, J. Ph. 17, 838; Simon, A. 31, 265; Blyth a. Hofmann, A. 53, 293, 325) and in coal tar (Berthelot, A. Suppl. 3, 368; Krämer a. Spilker, B. 23, 3282).

Formation.—1. By distilling cinnamic acid aloneor with lime or baryta (Gerhardt a. Cahours, A. Ch. [8] 1, 96; Kopp, C. R. 53, 634; Howard, C. J. 13, 135; Kraemer, Spilker, a. Eberhardt, B. 23, 3269).—2. By distilling cupric cinnamate (Hempel, A. 59, 316).—3. By distilling balsam of Peru with pumice (Scharling, A. 117, 184).—4. By distilling dragon's blood alone or with zinc-dust (Glénard a. Bondault, A. 53, 325; Bötsch, M. 1, 610).—5. By heating acetylene at a dull-red heat (Berthelot, C. R. 62, 905, 947; A. 141, 181).—6. By passing ethylene alone or mixed with benzene or diphenyl through a redhot tube (Berthelot, Z. [2] 4, 384; A. 142, 257; Barbier, C. R. 79, 660; Ferko, B. 20, 660).—7. By passing ethyl-benzene through a redhube (Berthelot, Z. [2] 4, 589).—8. By heating C.H.CH.Br alone or with alcoholic potash

at 180° (B.; Thorpe, Z. 1871, 130).—9. By heating O_aH_a .CHBr.CH₂ with alcoholic KCy (Radziszewski, B. 7, 140).—10. By distilling O_aH_a .CH(OAc).CH₃ (R.).—11. By heating O_aH_a .CH(OH).CH₃ (derived from accophenone) with ZnCl₂ (Emmerling a. Engler, B. 4, 147). 12. By the action of acetylene on benzene in presence of AlCl, (Varet a. Vienne, Bl. [2] 47, 918; C. R. 104, 1375).—13. From phenyl-acetylene, zinc, and HOAc (Aronstein a. Hollemann, B. 22, 1184).

Properties.—Oil, with aromatic odour. Inactive to light (Van't Hoff, Bl. [2] 25, 175). Miscible with alcohol and ether. Changes on keeping, or on heating in a sealed tube at 300°, to solid metastyrene. Combines with Cl2 and Br2. HBrforms C.H., CHBr.CH₃. HCl forms PhCHCl.CH₄ only (Schramm, B. 26, 1709). Potash has no action. Fuming HNO₂ forms nitro-styrene. Chromic acid mixture forms benzoic acid. Conc. H₂SO₄ changes it to metastyrene. On heating with aqueous NaHSO, at 120° styrene forms the compound C.H.,NaHSO. [306°] (W. von Miller, N. R. P. 24, 31). Condenses, in presence of H.SO., with toluene and xylene, forming diphenyl-propane and phenyl-tolyl-propane respectively (Kraemer a. Spilker, B. 23, 3169).

Metastyrene (C₈H₈)_x. S.G. 12 1.054. μ_D 1.593 (Madan, C. J. Proc. 1, 107). Occurs in liquid storax (Kovalevsky, A. 120, 66). Formed from styrene on keeping or on heating in a sealed tube at 200°. Formed also by heating styrene with aqueous NaHSO, at 110° (Miller, A. 189, 341). Transparent solid without taste or smell, becomes sticky when warmed. Highly refractive. Insol. water and alcohol, sl. sol. ether. Inactive to light (Van't Hoff, B. 9, 1339; cf. Berthelot, C. R. 85, 1191). Changes to styrene on distillation. Cl and Br act slowly, forming the same compounds as with styrene.

Di-styrene C₁₈H₁₈ i.e. Ph.CH:CH.CHMe.Ph(?). (c. 315° i.V.). S.G. 2 1.027; 18 1.016. V.D. 7.07 (calc. 7.2). Got by boiling cinnamic acid with H_2SO_1 (1 vol.) and water ($1\frac{1}{2}$ vols.) (Erlenmeyer, A. 135, 122; Fittig a. Erdmann, A. 216, 187). Liquid, with blue fluorescence, which slowly disappears. Inactive to light. If kept boiling for a long time it decomposes, forming toluene, styrene, high boiling products, and probably iso-propyl-benzene.

Reactions. - 1. Chromic mixture gives benzoic acid.—2. Bromine gives the di-bromide $C_{16}H_{16}Br_2$ [102°], crystallising from ether in needles. V. e. sol. CS2, ether or benzene, v. sol. hot alcohol.

Solid distyrene C₁₀H₁₀. [119°]. A product of the distillation of cinnamic acid or of its Ca salt (Engler a. Leist, B. 6, 256; Miller, A. 189, 340). Formed also by passing C.H., C.H., Br. over red-hot lime (Radziszewski, B. 6, 494). Tables. Yields a dibromide C₁₆H₁₆Br₂ [238°], crystallising from benzene in needles (Liebermann, B. 22, 2256).

References.—AMIDO-, BROMO-AMIDO-, Bromo-nitro-, Chloro-, Chloro-nitro-, Nitro-, and Nitro-amido-Styrene.

STYRENE DIBROMIDE v. DI-BROMO-ETHYL-

a. Michael, B. 10, 1558, 2200). Slender needles (from water). Changed on fusion into the lactone of oxy-carboxy-phenyl-propionic acid.—PbA".— Ag,A": sticky pp.

Isomeride v. p-Carboxy-cinnamic acid, vol. ii. p. 190.

STYRENE DICHLORIDE v. wa-DI-CHLORO-DI-ETHYL-BENZENE

STYRENE DI-IODIDE v. DI-IODO-ETHYL-BENZENE

STYRENE DISULPHOCYANIDE C,H,(SCN)2. [102°]. Formed by heating C,H,CHBr.CH,Br with potassium sulphocyanide in alcohol (Nagel, A. 216, 323). Pale-yellow needles, v. sol. ether, hot benzene, and toluene. Crystallises from benzene as $C_{10}H_{*}S_{2}N_{2}C_{e}H_{s}$ [62°]. Alcoholic ammonium sulphide at 100° forms $(C_{8}H_{*})_{2}S$ [151°]. Cold fuming HNO, gives rise to the compound [1:4] C_sH₄(NO₂).CH(SCy)CH₂(SCy) [112°] orystallising from alcohol in needles.

DI-STYRENIC ACID C, H, O, i.e.

Ph.CH:CH.CPhH.CH, CO,H or Ph.CH:CH.CH(CO₂H).CH₂Ph. Formed by boiling cinnamic acid with H₂SO₄ (1 vol.) diluced with water (1 vol.) (Fittig a. Erdmann, A. 216, 182). White amorphous powder. V. sl. sol. water, v. e. sol. ether, alcohol, or glacial acetic acid. It melts below 50°. May be distilled almost undecomposed.

Reactions.-1. Sodium amalgam has no action.-2. Br in CS2 forms no bromide.-3. Boiling H₂SO₄ (1 vol.) mixed with water (1) vols.) does not affect it. 4. Chromic mixture gives only benzoic acid.

Salts.— $Ca(C_{17}H_{18}O_2)_2$.— BaA'_2 .—AgA'. Ether.—EtA'. Liquid. STYROGALLOL C, H,O, i.e.

 $C_{e}H_{4} < C_{O} > C_{e}H(OH)^{3} > 0$ (3). Formed from

cinnamic acid, gallic acid, and conc. H2SO4 at 50° (Jacobsen a. Julius, B. 20, 2588; von Kostanecki, B. 20, 3137). Yellow needles, which may be sublimed; sl. sol. hot alcohol. Not melted at 350°. With mordants it dyes like nitro-alizarin.

STYROLENE ALCOHOL v. DI-OXY-ETHYL-BENZENE

STYRONE v. CINNAMYL ALCOHOL.

STYRONE DIBROMIDE v. (di)-Bromo-PHENYL-PROPYL ALCOHOL.

STYRYL-ACRYLIC ACID &. PHENYL-PEN-TINOIC ACID.

STYRYL AMYL KETONE CARBOXYLIC ETHER C1,7H22O3 i.e. C4H3.CH:CH.CO.CEt2.CO2Et. [102°]. Formed from di-ethyl-acetoacetic ether, benzoic aldehyde, and HCl in the cold (Claisen a. Matthews, A. 218, 184). Triclinic prisms (from ligroïn). Yields C₁₇H₂₂Br₂O₃ [55°] crystallising in small prisms.

STYRYL-ANGÈLIC ACID C₁₃H₁₄O₂ i.e. C₈H₃CH:CH:CH:CEt.CO₂H. [127°]. Formed by heating cinnamic aldehyde with butyric anhydride and sodium butyrate at 165° (Perkin, O.J. 81, 413). Crystals, v. sol. alcohol.—AgA'.

STYRYL BUTENYL KETONE C, H, O i.e. BENZENE.

STYRENE DICARBOXYLIC ACID C₁₀H_sO₄

i.e. [1:2] CO₂H_sCH:CH:CH:CO₂H_s o-Carboxycinnamic acid. [175°]. Formed by heating the salts
of oxy-carboxy-phenyl-propionic acid (Gabriel ing like strawberries. Yields C_{1s}H₁₄Br₄O [180°].

STYRYL-CROTONIC ACID $C_{12}H_{12}O_2$. [158°]. Formed by heating cinnamic aldehyde with propionic anhydride and sodium propionate (Perkin, C. J. 31, 413). Prisms (from ligroin).

-AgA': white pp., sl. sol. cold water.

STYRYL ETHYL KETONE CARBOXYLIC ACID C.H. CH:CH.CO.CH. CH. CO.H. Cinnamoyl-propionic acid. Benzylidene-levulic acid. [120°]. Formed from \(\beta\)-acetyl-propionic acid, benzoic aldehyde, and dilute KOHAq (Erdmann, A. 258, 130). Leaflets (from water). May be reduced to C.H., CH., CO.CH., CH., CO.H., ENGL. (See STYRYL ETHYL OXIDE C., H., 20 i.e.

C₆H₅.CH:CH:O.O.H₅. (217°). S.G. 981. Formed from ∞ -chloro-styrene and NaOEt at 180° (Erlenmeyer, B. 14, 1868). Aromatic liquid. Converted by heating with water into alcohol and phenyl-acetic aldehyde.

STYRYL-ETHYL-PYRIDINE C,,H,,N

 $\textbf{i.e.} \ \textbf{C}_{6}\textbf{H}_{\text{5}}.\textbf{C}\textbf{H}:\textbf{C}\textbf{H}.\textbf{C} \swarrow \begin{matrix} \textbf{N}.\textbf{C}\textbf{H} \\ \textbf{C}\textbf{H}:\textbf{C}\textbf{H} \end{matrix} \searrow \textbf{C}\textbf{E}\textbf{t}.$ Ethyl-stilbazole. [58°]. (356° cor.). Formed by heating methyl-ethyl-pyridine with benzoic aldehyde and ZnCl₂ at 220° (Plath, B. 21, 3087). Needles or plates, insol. water, v. sol. alcohol and ether, sl. sol. ligroïn. Bromine yields the dibromide Ogng, UHBr. CHBr. C, HgEtN. [128°]. — B'HCl. [193°]. — B'gH. PtCl. 2aq. [188°]. Needles.— [168°].—B'HSnCl, 3aq. [2 [196°].—B'C,H,N,O,. [203°]. L.GLYOXAL [246°].-B'HAuCl.

STYRYL-GLYOXAL. Oxim C₁₀H₀NO₂ i.e. CaHaCH:CH.CO.CH:N.OH. [144°]. Formed from nitroso-acetone, benzoic aldehyde, and alcoholic NaOEt (Claisen a. Manasse, B. 22, 529). Crystals (from water or benzene). Gradually decom-

poses, with evolution of HCy

STYRYL-GLY OXYLIC ACID C10H8Os i.e. C.H., CH:CH.CO.CO.H. Formed by saponification of its amide by conc. HClAq, and also by saturating a mixture of benzoic aldehyde and pyruvic acid with HCl (Claisen, B. 13, 2124; 14, 2472). Gummy mass, sol. water. Split up by alkalis, even in the cold, into benzoic aldehyde and pyruvic acid.-AgA': pp.

Amide CHPh:CH.CO.CO.NH, [130°]. Prepared by the action of conc. HClAq and HOAc on the nitrile of cinnamic acid. Flat prisms or

plates. Sol. ether and hot water.

Reference. - NITRO-STYRYL-GLYOXYLIC ACID. STYRYL-HYDANTOIC ACID C,1H12N2Os. [185°]. Formed by boiling styryl-hydantoin with baryta water (Pinner a. Spilker, B. 22, 692). White plates, sol. alcohol and hot water. Boiling HClAq converts it into styryl-hydantoin.—AgA'. STYRYL-HYDANTOIN v. DI-OXY-STYRYL-m-

PYRAZOLE. STYRYLIDENE-THIO-GLYCOLLIC ACID v.

CINNAMYLIDENE-THIO-GLYCOLLIC ACID.

DI-STYRYL-KETONE v. DI-BENZYLIDENE-ACETONE and NITRO- and OXY- DI-STYRYL-KETONE. STYRYL-DI-METHYL-GLYOXALINE

C₁₃H₁₄N₂ i.e. C₈H₅CH:CH.C≪NH.CMe N − CMe [202°]. Formed from di-methyl di-ketone, cinnamic aldehyde, and alcoholic ammonia at 100° (Wadsworth, C. J. 57, 11). Crystals, v. sol. alcohol.—B',H,PtCl,: minute yellow needles.
STYRYL METHYL KETONE v. BENZYLIDENE-

ACETONE and NITBO- and OXY- STYRYL METHYL

KETONE.

STYRYL METHYL KETONE CARBOXYLIC ETHER v. BENEYLIDENE-ACETOACETIC ETHER, vol. i. p. 24.

STYRYL-METHYL-OXAZOLE DIHYDRIDE C.H.CH:CH.C≪N.CH. [81°]. Formed from

β-bromo-propylamine hydrobromide, cinnamoyl chloride, and alcoholic potash (Elfeldt, B. 24, 3226). Transparent crystals. — B'2H2PtCl6. $-B'C_6H_3\hat{N}_3O_7$. [183°]. Needles. [198°].-

STYRYL-METHYL-PYRIDINE C14H18N i.e. C.H., CH.CH.C.H., MeN. Methyl-stilbazole. (c. 323°). S.G. § 1.0717. Formed by heating (c. 323°). S.G. ½ 1.0717. Formed by nearing benzoic aldehyde with (αγ)-di-methyl-pyridine and ZnCl₂ at 215° (Bachér, B. 21, 3072). Oil, v. sol. alcohol and ether. Gives C₁H₁₈Br₂N. [140°]. May be reduced to C₁4H₁₈N (c. 293°) and C₁4H₁₈N (c. 288°).—Hydroiodide. [211°]. B', H.PtCl6aq. [183°].—B'HAuCl₄. [142°]. B'HHgCl₃.—B'C₄H₃N₃O₇. [193°]. Yellow needles.

STYRYL-DI-METHYL-PYRIDINE DICARB-

OXYLIC ACID $C_{17}H_{18}NO_4$ i.e. $C_8H_5.CH:CH.C < \begin{array}{c} C(CO_2H).CMe \\ C(CO_2H):CMe \end{array} N. \quad [241°].$

Formed from its ether, which is got by the action of nitrous acid gas on an alcoholic solution of the dihydride C₂₁H₂₅NO₄ [149°], obtained from cinnamic aldehyde, acetoacetic ether, and alcoholic NH₃ (Epstein, A. 231, 1). Minute colourless tablets (containing 2aq), sl. sol. water, sol. alcohol. Melts at 219° when hydrated. Converted by hypochlorous acid into C_{1}, H_{1}, NO_{4} 2aq (Messinger, B. 19, 196).— $K_{2}A''$ 3aq. — $(H_{2}A'')_{2}H_{2}$ PtCl₆.— $H_{2}A''$ HCl. Decomposed by water.

Ethyl ether Et2A". [39°]. Yields the salt (Et,A"),H,PtCl, [195°].

STYRYL-METHYL-THIO-GLYCOLLIC ACID v. CINNAMYL-THIO-GLYCOLLIC ACID.

STYRYL-(a)-NAPHTHOQUINOLINE

C₂₁H₁₅N i.e. C₆H₄ CH:CH.C.CH:CH -C. N :C.CH:CHPh.

[104°]. Formed by distilling the carboxylic acid $C_{10}H_6 < C(CO_2H):CH$ $C_{10}CCH:CHPh$ [256°], which is

got by the action of (a)-naphthylamine on cinnamic aldehyde and pyruvic acid (Döbner a. Peters, B. 23, 1233). Concentric groups of yellowish needles, sl. sol. alcohol.—B'2H,PtCl, 2aq: orange-yellow pp. — B'2H2Cr2O7. — B'C4H3N3O7. [230°]. Golden needles.

Carboxylic acid C₂₂H₁₅NO₂. Yields BaA'₂2aq, CuA'₂ aq, and AgA'. Styryl-(\$)-naphthoquinoline [256°].

- Ç.CH:ÇH C.H.CH.C. N : O.CH:CHPh [175°].

Formed in like manner from (β) -naphthylamine. White silky needles or pearly plates.—Salts: B',H,PtCl, 2aq. — B',H,Cr,O, — B'C,H,N,O, [254°]. Golden needles, sl. sol. ether.

Carboxylic acid C22H15NO2. [305°]. Lemon-yellow needles, sl. sol. hot alcohol.

STYRYL-OXAZOLE DIHYDRIDE C,H,NO i.e. C_eH₅CH:CH.C≪N.CH₂. [53°]. Formed from CH,Br.CH,NH.CO.CH:CHPh H.NH.CO.CH:CHPh and alcoholic (Elfeldt, B. 24, 3225).—B'₂H₂PtCl₂. and potash [194°]. Orange-yellow powder.—B'CaHaNaO,. [189°]. Yellow needles.

STYRYL-PENTOXAZOLE DIHYDRIDE

 $\mathbf{C}_{e}\mathbf{H}_{s}\mathbf{CH}:\mathbf{CH}.\mathbf{C}\leqslant_{\mathbf{N},\mathbf{CH}}^{\mathbf{O}.\mathbf{CH}_{2}}\mathbf{CH}_{2}$. [56°]. by the action of alcoholic potash on the γ -bromopropyl-amide of cinnamic acid (Elfeldt, B. 24, 3227). Needles.—B'.H.PtCl_s. Decomposes at 193°.—B'C.H.N.O.. [196°].

STYRYL-PHENOL v. OXY-DI-PHENYL-ETHYL-Decomposes at

(a)-STYRYL-PYRIDINE C, H, N i.e.

C_eH_sCH:CH.C CH:CH CH. Stilbazole. [91°]. (825° cor.). Formed by heating benzoic aldehyde with methyl-pyridine and ZnCl2 at 220° (Baurath, B. 20, 2719; 21, 818). Crystals, m. sol. alcohol.

Reactions.—1. Br forms C₁₃H₁₁Br₂N [167°]. 2. HIAq reduces it to C₁₃H₁₃N [-3°].—3. Na and HOEt reduce it to C₁₃H₁₈N (288° cor.).

Salts.-B'HCl 4aq: needles. Melts at 177° when anhydrous.—B',H,PtCl, 2aq. [188°]. Red needles.—B'HAuCl,. [185°].—B'HHgCl, aq. [183°].—B'HI,. [159°]. Prisms, with blue reflex. Reference.—OXY-STYRYL-PYRIDINE.

(Py. 8)-STYRYL-QUINOLINE C17H12N i.e.

C.H. CH:CH:CHPh. [100°]. Formed by distilling its carboxylic acid (Doebner a. Peters, B. 22, 3008). Got also by heating (Py. 3) methyl-quinoline with benzoic aldehyde and ZnCl₂ (Wallach, B. 16, 2008; Jacobsen a. Reimer, B. 16, 2606). Crystals. C₁₇H₁₈Br₂N [174°].—Salts: B'₂H₂PtCl₄ 2aq.-B'H, Cr.O., 2 saq. References.—Nitro- and Oxy- Styryl-quin-

OLINE

STYRYL-QUINOLINE CARBOXYLIC ACID

 $C_8H_4 < N CC_2H): CH$ [295°]. Formed C.CH:CHPh' from cinnamic aldehyde, pyruvic acid, and aniline in alcohol (Doebner a. Peters, B. 22, 3006). Yellow needles, insol. water, sl. sol. ether.-MgA'2: concentric groups of needles.

STYRYL-QUINOLINE SULPHONIC ACID C,H,(SO,H)N.CH:CHPh. Formed by heating (Py. 1)-methyl-quinoline (B. 2)-sulphonic acid with benzoic aldehyde and ZnCl2 for six hours at 170° (Busch a. Koenigs, B. 23, 2682). Crystals (containing 2aq). V. sol. 50 p.c. acetic acid.

STYRYL THIENYL KETONE C, H, SO i.e. C₆H₅CH:CH.CO.C₄H₈S. [80°]. Formed by passing HCl into a mixture of thienyl methyl ketone and benzoic aldehyde (Brunswig, B. 19, 2895) Needles. Yields a dibromide C₁₃H₁₀Br₂SO [157°] crystallising from alcohol in plates

DI-STYRYL-DI-VINYL KETONE v. DI-

PHENYL-DI-BUTINYL KETONE.

METHYL STYRYL-VINYL KETONE v.

PHENYL-BUTINYL METHYL KETONE

SUBERCOLIC ACID C.Hs(CO2H)2. Formed in small quantity by the action of alcoholic potash on di-bromo-suberic acid (Hell a. Rempell, B. 18, 820). White powder, yielding at 225°-230° a sublimate of slender white needles.-

BaA".—CaA".—MgA" 2aq.—Ag,A": white pp. sUBERCONIC ACID. [165°-170°]. A crystalline product of the action of alcoholic potash on bromo-suberic acid (Ganttner a. Hell, B. 15,

SUBERENE CARBOXYLIC ACID C.H.,O

C₈H₁₃ClO₂ with NaOHAq (Spiegel, A. 211, 119). Pearly plates, volatile with steam. Reduced by sodium-amalgam to suberane carboxylic acid C₈H₁₄O₂.

SUBERIC ACID C.H.O. Mol. w. 174. [140°]. (c. 300°). (279° at 100 mm.) (Krafft, B. 22, 816). S. 142 at 15°. S. (ether) 8 at 15°. H.C.p. 992,409 (Louguinine, C. R. 107, 597); 985,600 (Stohmann, J. pr. [2] 40, 215). H.F. 249,400 (S.).

Formation.—1. By the action of nitric acid upon cork (Brugnatelli, Crell. Ann. 1787, i. 145; Bouillon-Lagrange, A. Ch. 23, 2; J. Ph. 8, 107; Chevreul, A. Ch. 62, 323; 96, 182; Brandes, S. 32, 393; 83, 83; 36, 263; A. 9, 295; Bussy, J. Ph. 8, 107; 19, 425; Boussingault, J. pr. 7, 211; Harff, N. Br. Arch. 5, 303).—2. By the action of nitric acid upon oleic and stearic acids (Laurent, A. Ch. [2] 66, 157; Bromeis, A. 35, 89), upon castor oil (Tilley, A. 39, 166; Dale, A. 132, 244; 199, 145; Grote, A. 130, 208), upon linseed oil (Sacc, A. 51, 222), upon cocoa-nut oil (Wirtz, A. 104, 261), upon almond oil, upon spermaceti (Arppe, A. 120, 292; 124, 89), upon palmitolic acid (Schröder, A. 143, 33), upon paraffin (Pouchet, C. R. 79, 320), upon palm oil (Ganttner a. Hell, B. 13, 1165; 14, 1549), and upon myristic acid (Noerdlinger, B. 19, 1896) .-3. Occurs among the products of the distillation of crude fatty acids in superheated steam (Cahours, C. R. 94, 610).—4. From di-oxy-stearic acid and alkaline KMnO, (Spiridonoff, J. pr. [2] 40, 250).—5. Suberio ether is formed by the electrolysis of CO₂Et.CH₂.CH₂.CH₂.CO₂K, the yield being 28 p.c. of the theoretical amount (Crum Brown a. Walker, A. 261, 120).

Properties.—Long needles (from water) or tables. May be sublimed in needles. Not volatile with steam.

Reactions.-1. Yields hexane when distilled with baryta.—2. Forms suberone on distilling with lime.-3. Bromine at 160° forms bromoand di-bromo-suberic acids, which when boiled with potash yield oxy- and di-oxy- suberic acids-(Gal a. Gay-Lussac, C. R. 70, 1175).

Salts.-K2A". S. 85 at 14°.-Na,A" \frac{1}{2}aq. S. (of Na₂A") 50 at 14°.—NaIIA".—(NH₄)₂A". S. 38 at 25°.—BaA": crystalline powder, less sol. hot water than cold. S. 2.2 at 7.5°; 1.8 at 100°.—SrA". S. 2.9 at 14°; 1.9 at 100°.—CaA" aq. S. 62 at 14°; 42 at 100°.—MgA" 2aq. S. 13.5 at 20°.—ZnA". S. 041 at 14°.—CuA" aq: blue pp. S. (of CuA") '024 at 16°.--CuA" 2aq. CdA" aq. S. (of CdA") ·08 at 17°.—HgA". S. ·012 at 7·5°.—Al₂OA"₂. S. ·0094 at 6·5°.—PbA". S. ·008 at 16°.—Pb₂A"O₂.—MnA" 3aq. S. 1·08 at 13°.—Fe₄A"₄O. S. ·0015 at 9°.—CoA" 4aq.— CoA" 2aq. S. (of CoA") 1·16 at 14°; ·85 at 100°.
—NiA" 4aq. S. (of NiA") ·79 at 7·5°.—Ag₂A". S. .0075 at 8°.

Methyl ether Me.A". S.G. 18 1.014. Ethyl ether Et.A". (282°). S.G. Ethyl ether Et₂A". (282°). S.G. $\frac{15}{15}$. 9852; $\frac{25}{25}$. 9783 (Perkin, C. J. 45, 517). M.M. 12.461 at 14°

Amide C.H. N.O. Crystalline. Amic acid C.H. (CONH.).CO.H. [c. 170°]. Formed by distilling ammonium suberate. Crystalline, v. sol. hot water.

Anilide C₆H₁₂(CO.NHPh)₂. [183°]. Formed [54°]. Formed by boiling chloro-suberonic acid by heating suberic acid (1 pt.) with aniline (1 pt.) (Gerhardt a. Laurent, A. Ch. [3] 24, 185). Pearly plates (from alcohol), v. sol. ether.

Phenylamic acid C.H. (CO.NHPh).CO.H. Formed at the same time as the anilide. Minute laminæ, sl. sol. hot water.—AgA': white pp.

References .- Bromo-, Chlobo-, and Oxy-

SUBERIC ACID.

Isosuberic acid is DI-ETHYL-SUCCINIC ACID

(Hell, B. 22, 67)

SUBERIC ALDEHYDE C.H.O. (202°). A product of the action of fuming HNO, on palmitolic acid (Schröder, A. 143, 34). Oil, partially decomposed by distillation. Oxidised by bromine water to suberic acid.

SUBEROCARBOXYLIC ACID v. HEXANE

TRICARBOXYLIC ACID.

SUBEROMALIC ACID v. OXY-SUBERIC ACID. SUBERONE C,H,2O. Mol. w. 112. (180° i.V.). V.D. 3.73 (calc. 3.89). Formed, together with hexane, by distilling suberic acid with lime (Boussingault, A. 19, 808; Tilley, A. 39, 167; Dale a. Schorlemmer, C. J. 27, 935; A. 199, 147; Wislicenus, A. 275, 356). Colourless liquid, smelling like peppermint. Oxidised by HNO₃ (S.G. 14) to n-pimelic acid [102°]. Does not reduce AgNO₃ or Fehling's solutions. Combines with bromine, forming a product which, when distilled over KOH, gives $C,H_{12}O$ (Ladenburg, B. 14, 2406). Combines with HCy yielding $C,H_{12}(OH)CN$, which is converted by HCl into suberyl-glycollic acid C,H₁₂(OH).CO₂H [80°], from which conc. HClAq at 130 produces chloro-suberane carboxylic acid C,H₁₂Cl.CO₂H, a thick oil, converted by KOH into suberene carboxylic acid C,H,1.CO,H [54°], which may be reduced by sodium-amalgam to suberane carboxylic acid C,H1s.CO2H, and this may be exidised by HNO, to a dibasic acid $C_8H_{14}O_4$ or $C_8H_{12}O_4$ (Dale a. Schorlemmer, C. J. 20, Spice of A. 211, 117). Subseque 1 39, 539; Spiegel, A. 211, 117). Suberone is reduced in alcoholic solution by sodium to $C_1H_1O_1$, a colourless liquid, with mouldy smell (185°), S.G. $\frac{15}{15}$ 9595, which unites with phenyl cyanate forming C,H,,O.CO.NHPh [85°], and also with HI forming a compound which is converted by alcoholic potash into suberonylene C,H₁₂ [114.5°] and a little C,H₁₃OEt. The compound C,H₁₄O is reduced by conc. HIAq at 250° to C,H14. Suberonylene combines with bromine (Markownikoff, C. R. 110, 466).

Oxim C,H12:NOH. Liquid, smelling like peppermint, sol. alcohol, ether, alkalis, and acids (Nägeli, B. 16, 497). Reduced by sodiumamalgam to C,H13.NH

SUBERO-TARTARIC ACID v. DI-OXY-SUBERIC

SUBERYL-GLYCOLIC ACID v. OXY-SUBERANIO

SUBLIMATION. The passage of a solid body, when heated, to the state of vapour without melting. The temperature of sublimation is definite for every solid, and is dependent on the pre sure. There is a definite pressure for each definite solid below which the substance cannot exist as a stable liquid, so that if heated below this pressure it passes from the solid directly to the gaseous state; but if heated under pressures greater than this pressure the substance first liquefies and then vaporises. As the definite pressure is considerably below that of the atmosphere for most substances, but few bodies undergo sublimation, properly so called, when heated under ordinary conditions. Sublimation is treated fairly fully in Thorpe's DICTIONARY OF AP-PLIED CHEMISTRY (vol. iii. p. 609), and instances are given where the process is made use of in M. M. P. M. manufacturing operations.

SUBSTITUTION. The replacement of one element in a compound by another element is the primary meaning given to the term 'substitution.' As our ways of looking at composition, and changes of composition, are saturated with the conceptions of the molecular and atomic theory, substitution is better described as the replacement of an atom in a molecule by another atom. But the term must be widened to include cases wherein an atom is replaced by a group of atoms, which group is regarded in that reaction as if it were a single atom. The moment attention is paid to the number of atoms that can be substituted for some specified atom in a molecule the conception of chemical equivalency is found necessary to give a common foundation to the facts that are observed (v. Equivalency, vol. ii., especially pp. 449-451). The properties of a molecule are changed by the substitution of an atom or atomic group in the place of one, or more, of the atoms in the original molecule; the prosecution of the inquiry into the connections between the properties of molecules derived, by substitution, from some parent molecule leads to the hypothesis of types, which is afterwards merged in the wider subject of chemical classification (v. Classification, Chemical, vol. ii. p. 196; Equivalency, vol. ii. pp. 450-451; Types, in this vol.).

Experiments show that the substitution of a more positive by a less positive atom, or atomic group, is accompanied by a change of properties in the direction that the substitution-product is more acidic than the parent substance. This field of inquiry has been cultivated systematically in recent years, and measurements have been made of the effects of substitution-changes of the kind indicated on the affinities of many acids. The results obtained have led, and are leading, to new conceptions of the nature of affinity, and of the constitutions of molecules (v. Physical methods, section Electrical methods, pp. 197-207 in this vol.).

M. M. P. M.

SUCCINAMIDINE C4H, N, i.e.

CH₂·C(NH₂)NH. Formed by the action of ammonia on succinic-di-imido-di-ethyl ether C.H.(C(OEt):NH)₂ (Pinner, B. 16, 362, 924, 1643, 1655). The salt B"2HCl forms minute needles, dissolving in water with formation of NH4Cl and

succinimidine C.H.(NH)...
SUCCINAMIDOXIM C₂H₄(C(NH₂):NOH)₂. [188°]. Formed by Succinene diamidoxim. hydroxylamine hydrochloride Na₂CO₂Aq to an alcoholic solution of the nitrile of succinic acid (Sembritzki, B. 22, 2958). Crystals, sol. hot water, sl. sol. hot alcohol, insol, ether. Coloured reddish-brown by FeCla. Fehling's solution gives a dirty green pp. Ac₂O forms C₂H₁(C(NH₂):NOAc)₂ [168°]. NaOEt and EtI form C₂H₁(C(NH₂):NOEt)₂ [119°]. BzCl and NaOH produce the benzoyl derivative C₂H₁(C(NH₂):NOBz)₂ [192°], crystallising in needles, converted by heating with water at 159° into $C_2H_4(O \swarrow_N^{N,O} \geqslant CPh)_2$ [159°]. Aquepotassium cyanate and HCl form

Succinimidoxim CH₂.C(NOH) NH. Formed CH₂.C(NOH) by heating the nitrile of succinic acid with hydroxylamine at 65° (Sembritzki, B. 22, 2964). Prisms (containing 2aq). Sol. hot water, insol. alcohol. Alcoholic NaOH forms a blue solution turning green. FeCl, gives a dark-violet colour.—
Ag₂C₄H₅N₃O₂: pearly plates.

Di-acetyl derivative C₄H₅Ao₂N₂O₂.

Di-acety l C,H,Ac,N,O,.

[171°]. Crystalline powder, sol. water.

Di - benzoyl derivative C.H.Bz.N.O. Crystals, insol. water, sol. hot alcohol.

SUCCINIC ACID C,H,O, i.e. 2H.CH,CH,CO,H. Mol. w. 118. CO₂H.CH₂.CH₂.CO₂H. (Krafft a. Noerdlinger, B. 22, 816; Reissert, B. 23, 2245); [185°] (Davidoff, B. 19, 406). (261°). S.G. 1.55. S. 2.88 at 0°; 5.14 at 14.5°; 121 at 100° (Bourgoin, Bl. [2] 21, 110; 29, 243; Miczynski, M. 7, 263). S. (alcohol) 7.5 at 15°. S. (ether) 1.25 at 15°. H.C.v. 357,100. H.C.p. 356,800. H.F. 226,200 (Stohmann, J. pr. [2] 40, 207); 229,000 (von Rechenberg). S.H. 290 (from 0°-50°); 365 (0°-150°) (Hess, A. Ch. [2] 35,410). Heat of Neutralisation: Gal a. Werner, C. R. 103, 871.

Occurrence.—In amber (Agricola; Berzelius, A. Ch. 94, 187; Lecanu a. Serbat, J. Ph. 8, 541; 9, 89; Liebig a. Wöhler, P. 18, 162; D'Arcet, P. 36, 80), in turpentine from several species of pine, in Chelidonium majus (Walz, N. J. P. 15, 22; Zwenger, A. 114, 350), in lettuce (Köhnke, B. J. 25, 443), in Papaver somniferum, in unripe grapes (Brunner a. Brandenburg, B. 9, 982), in exudations from the bark of mulberry trees (Goldschmiedt, M. 3, 136), in beet juice (von Lippmann, B. 24, 3299), and in rhubarb (Brunner, B. 19, 595). Occurs also in the thymus gland of the calf, the spleen of the ox (Gorup-Besanez, 1. 98, 28), in certain pathological exudations (Heintz, A. 76, 369; Brieger, H. 5, 368), in the urine of rabbits fed on carrots (Meissner a. Jolly, J. 1865, 675; cf. Salkowski, Pf. 4, 91), and in human urine after eating asparagus (Hilger, A. 171, 208).

Formation.—1. By the action of HNO, on many organic substances, including fats, fatty acids from butyric acid upwards, wax, spermaceti, sebacic acid, and azelaïc acid (Bromeis, A. 35, 90; 37, 292; Sthamer, A. 43, 346; Ronalds, A. 43, 356; Radcliff, A. 43, 351; Arppe, A. 95, 242; Dessaignes, A. 70, 102; 74, 361; Erlenmeyer, B. 7, 696; Noerdlinger, B. 19, 1895). -2. In the alcoholic fermentation of sugar (Pasteur, Bl. 1852, 52; A. 105, 264).—3. In the fermentation of asparagine (Piria, A. 68, 343), of calcium malate (Dessaignes; Liebig, A. 104, 363), of ammonium tartrate (König, B. 14, 211), and of flesh (Salkowski, B. 12, 649).-4. By reducing fumaric and maleic acids with sodium-amalgam (Kekulé, A. Suppl. 1, 133).—5. By reducing malic and tartaric acids with HI (R. Schmitt, A. 114, 106; Dessaignes, A. 115, 120; 117, 134).—6. By heating its nitrile (ethylene cyanide) with alcoholic potash at 100° (Maxwell Simpson, A. 118, 878), or by treating it with dilute nitric acid (Jungfleisch, Bl. [2] 19, 197).—7. From bromo-acetic acid and reduced silver at 180° (Steiner, B. 6,

184). - 8. From \$ chloro-propionic ether by successive treatment with KCy and potash (Wichelhaus, Z. [2] 3, 247).—9. From sodium acetoacetic ether by treatment with chloroacetic ether and saponification of the product (Wippermann, B. 3, 337; Noeddecke, A. 149, 224).—10. By heating ethane tricarboxylic acid at 160° (Bischoff, B. 13, 2162).—11. By potashfusion from gum arabic, milk sugar, and carminic acid (Hlasiwetz, A. 138, 76; 141, 340).— 12. By reducing acetylene dicarboxylic acid with zinc and HOAc at 60° (Aronstein a. Hollemann, B. 22, 1183).-13. Its ether is formed by electrolysis of CO₂Et.CH₂.CO₂K in aqueous solution; the yield being 60 p.c. of the theoretical (Crum Brown a. Walker, A. 261, 115).

Preparation.-1. By distilling amber, heating the watery distillate, filtering, and allowing to crystallise. The product is freed from oils by treatment with nitric acid.-2. By neutralising tartaric acid (2 kilos) with NH, Aq, adding to the aqueous solution (40 litres) K2HPO, (20 g.), MgSO₄ (10 g.) and a little CaCO₃. Fermentation is started by diluting a little (5 c.c.) of the solution with water (25 c.c.) and exposing it to the air for a few days. When fermentation has set in. the liquid is returned to the main quantity, and the whole kept at 25°-30° for seven weeks. The liquid is evaporated somewhat, clarified by white of egg, and boiled with lime. After cooling, the calcium succinate is collected and decomposed by H₂SO₄. The yield is good (500 g.) (Koenig,

B. 15, 172).

Properties.-Monoclinic prisms, permanent in air, acid in taste. V. sol. water, m. sol. al-cohol, sl. sol. ether. Resolved on boiling into water and anhydride; the lactone C,H,O, being also formed. BaCl₂ completely ppts. hot neutral solutions (Schmitt a. Hiepe, Fr. 21, 536). BaCl, followed by NH₃Aq and alcohol ppts. succinic (but not benzoic) acid. FeCl₃ gives in neutral solutions a reddish-brown pp. PbAc, gives a

pp. sol. excess.

Reactions .- 1. Not attacked by HNO, CrO, or chlorine water. MnO2 and H2SO, yield acetic acid. KMnO, in neutral solution oxidises it to oxalic acid and CO₂; and, in acid solution, to CO₂ (Berthelot, Bl. [2] 8, 390; Sorokin, J. R. 11, 383).- 2. Yields ethylene and CO, when decomposed in alkaline solution by an electric current (Kekulé, A. 131, 79; Bourgoin, A. Ch. [5] 20, 80; cf. Kolbe, A. 113, 244). In perfectly neutral solution only CO2, CO, and O are given off at the positive pole.—3. Potash-fusion yields oxalic acid.—4. PCl, forms successively: succinic anhydride C₂H₁(CO)₂O, succinyl chloride C₂H₁(COCl)₂, fumaryl chloride C₂H₂(COCl)₂, chloro-fumaryl chloride C₂HCl(COCl)₂, dichloro-maleyl chloride C₂Cl₂(COCl)₂, two tetrachlorinated di-chloro-maleyl chlorides, viz.: C,Cl,(CCl,)(COCl) and C,Cl,(CCl,),O(v.1)1-GHLORO-MALEIC ACID), and finally per-chloro-ethane, 2C.Cl. (Kander, J. pr. [2] 81, 1).—5. On long boiling it yields the dilactone of diethyl ketone s-di- ω -carboxylic acid CO .0 O.CU [75°] CH, CH, CH, CH, CH alohol in (c. 203° at 15 mm.), crystallising from alcohol in plates, converted by HClAq into hydrochelidonic acid CO(CH, CH, CO, H), [143°], which yields an oxim [129°] (Volhard, A. 253, 206).—6. Conver-

ted into succinic anhydride by treatment with

P₂O₂, AcCl, BzCl, Ac₂O, or succinyl chloride (Anschütz, B. 10, 325, 1881).—7. A solution containing 5 p.c. succinic acid and 1 p.c. uranium succinate when exposed to sunlight turns green and gradually gives off CO2, leaving propionic acid in solution (Seekamp, A. 133, 253). Barium succinate (1 mol.) heated with NaOMe (1 mol.) for 3 hours at 300° yields propionic acid (Mai, B. 22, 2133).—8. Heated with a large excess of lime it yields ethane as chief product (Hanriot, Bl. [2] 45, 79).—9. The salts on distillation yield a liquid product (160°-250°) which yields benzene when distilled over zinc-dust (Von Richer, J. pr. [2] 20, 206; cf. Fumaro, G. 11, 273). Hydroquinone is also a product of the distillation of succinates.—10. The sodium salt heated with P2S, yields thiophene (Volhard a. Erdmann, B. 18, 454).—11. The Na salt heated with aldehydes and NaOAc forms oxyacids of the form R.CH(OH).CH(CO₂H).CH₂.CO₂H (Fittig, B. 18, 2523).—12. By heating with phthalic anhydride and NaOAc at 250°, exhausting the product with water and alcohol, and boiling the residue with aniline, there is got 'diphthalsuccinanilide' C₃₀H₂₄N₂O₄ [267°] converted by boiling with HOAc and HCl into 'diphthalsuccindehydranilide ' C_{so}H₂₀N₂O₂, crystallising from HOAc in small yellow prisms, not liquid at 280° (Roser, B. 18, 3122).—13. By heating with glycerin at 200° it yields 'succinin' C3H3(OH)A", a gummy mass, insol. cold water, alcohol, and ether (Fumaro a. Danesi, G. 10, 58). Alkalis and acids split it up into glycerin and succinic acid.—14. a-Chloropropionic ether and alcoholic potash give C₂H₄(CO.O.CHMe.CO.Et)₂ (c. 302° at 730 mm.) (Wurtz a. Friedel, A. Ch. [3] 63, 101; Wislicenus, A. 133, 262).

Salts.—(NH₄)₂A". Hexagonal prisms, v. sol. water and alcohol. Yields succinamide when heated.—(NH₄)HA". Triclinic crystals, v. sol. water and alcohol (Brooke, Ann. Phil. 22, 286). Formed by evaporating a solution of the neutral salt.—K,A" 2aq. Deliquescent crystals, sol. alco-Salzer, B. 16, 3025).—KHA" 2aq. Trimetric crystals (Salzer, B. 16, 3025).—KHA" 2aq. Efflorescent six-sided prisms.—KH₃A"₂.—Na₂A" 6aq: monoclinic prisms.—NaHA". Triclinic prisms.—NaHA". Sl. sol. water, insol. NH, Aq and alcohol. S. 42 at 3°; 43 at 12°; 28 at 67° (Miczynski).— CaA" 8aq: small needles, deposited gradually on mixing cold conc. solutions of sodium succinate and CaCl₂. On mixing the hot solutions CaA" aq is ppd. S. 1.1 at 0°; 1.8 at 24°; .8 at 68° (Miczynski, M. 7, 266).—CaH2A"22aq.—SrA": (Miczynski, M. 7, 266).—CaH₂A"₂2aq.—SrA": monoclinic prisms.—BeA" 2aq (Atterberg, Bl. [2] 21, 162).—Be₂(OH)₂A"₂2aq.—MgA" 6aq. Prisms (Fehling, A. 49, 154).—MgK₂A"₂.—Mg₃A"₂O₄ aq.—CrA" aq: scarlet pp.—CuA" (dried at 200°).—CuA"2NH₂.—CuA"4NH₃ (Schiff, A. 123, 45).—Fe(OH)A": brownish-red pp.—NiA" 4aq.—MnA" 4aq. Triclinic prisms (Handl, Sitz. W. 32, 254).—PbA" (dried at 100°). White powder, sl. sol. water, v. sol. KOHAq and HNO₃.—Pb.OA"₂: sticky pp., got by adding lead sub-Pb_sOA"₂: sticky pp., got by adding lead subacetate to sodium succinate.—Pb,0,2A", (dried at 200°): white powder.—ZnA" (dried at 200°). Crystalline powder.—ZnA"3NH, (Lutschak, B. 5,30).—CdA".—(UrO),A"aq.—K(UrO)A" aq.—Na(UrO)A" aq.—Ag.A". White amorphous pp. -Quinine salt (C₂₀H₂₄N₂O₂)₂H₂A" aq. Prisms.

S. ·11 at 10° (Hesse, A. 135, 331).—Cinchonine salt $(O_{22}H_{24}N_2O)_2H_2A''$ aq. Thick crystals (Hesse, A. 122, 226).—Cinchonidine salt $(O_{22}H_{24}N_2O)_2H_2A''$ 6aq. Silky prisms. S. ·4 at 10° (Hesse, A. 135, 342).—Urea salt $(CON_2H_4)_2H_2A''$. Six-sided monoclinic prisms; a:b:c=1·483:1:1·365; $\beta=83^\circ$ 28' (Lischmidt, Sitz. W. 52, ii. 238).—Et hylene-diamine salt $C_2H_4(NH_2)_2H_2A''$. [182°]. Thick white prisms, v. sol. water, insol. ether (Mason, C. J. 55, 10).—Benzylamine salts $(C,H,NH_2)_2H_2A''$. [145°]. Thin plates, sol. aloohol (Werner, C. J. 55, 628).—(C,H,NH_2)H_2A''. [117°]. Rectangular prisms (containing aq), v. sol. water.

Methyl ether Me₂A". [19°]. (195° cor.).
S.G. ²⁰ 1·1209 (Emery, B. 22, 3185); ¹⁰ 1·1261; ¹⁰ 1·383 (Perkin, C. J. 45, 516). M.M. 6·232 at 18·2°. S.V. 159·7 (Lossen, A. 254, 64). H.C. (solid) 703,600. H.C. (liquid) 708,500. H.F. (solid) 205,400 (Stohmann, J. pr. [2] 40, 353). Formed from succinic acid, MeOH, and gaseous HCl (Fehling, A. 49, 195). Formed also from succinyl chloride and NaOMe in ether, and from Ag₂A" and MeI. Crystalline, sol. alcohol.

Methylethylether MeEtA". (208° cor.). C.G. 2 1.093. S.V. 184.6. Formed by the action of EtAgA" on MeI (Köhler, A. 221, 88). On saponification by baryta it yields H.A", HMeA", and HEtA" (Lossen a. Köhler, A. 262, 200).

Mono-ethyl ether HEtA". Formed by boiling succinic anhydride with alcohol (Heintz, J. 1859, 280). Syrup, miscible with water, alcohol and ether.—AgEtA": amorphous, sl. sol. water.—The salt NaEtA" is converted by POCl, into CO₂Et.C₂H₄.COCl (144° at 90 mm.), which is split up by distillation into EtCl and succinic anhydride, and is converted by phenyl hydrazine into CO₂Et.C₂H₄.CO.NH.NHPh [107°], crystallising in white needles, itself converted into O(CO.C₂H₄.CO.N₂H₂Ph)₂ [137°] by successive treatment with HOH and HCl (Michaelis a. Hermans, B. 25, 2748). The salt NaEtA" is converted by a-chloro-propionic ether into CO₂Et.C₂H₄.CO.CHMe.CO₂Et (280°) S.G. 2 1·119, which is split up by boiling with baryta water into succinic and lactic acids (Wurtz a. Friedel, J. 1861, 378).

Ethylether Et.A". (216.5° cor.). S.G 15 1.0465; $\frac{25}{95}$ 1.0383. M.M. 8.380 at 17.8^6 (Perkin). S.V. 209 4. Formed by boiling succinic acid (20 pts.) with alcohol (8 pts.) and H₂SO₄ (1 pt.) (Eghis, B. 6, 1178; cf. D'Arcet, A. Ch. [2] 58, 291). It is also produced by heating C₂H₄Br₂ in alcohol with potassium succinate in sealed tubes (Davidoff, Bl. [2] 46, 818; 19, 406). Oil. Converted by Na into the dihydride of dioxyterephthalic ether. Not attacked by hydroxylamine (Jeaurenaud, B. 22, 1273). Succinic ether (40 g.) treated with cyanamide (20 g.) and an alcoholic solution of K (18 g.) for 5 hours with inverted condenser yields C₂H₄(CO.NKCy)₂ and C₂H₄(CO₂K)(CO.NKCy). These bodies may be converted by AgNO, into Ag salts, which may be separated by HNO, in which silver 'succincyanamate' dissolves, while silver 'succinyl-di-cyan-di-amide' is insoluble. The Ag salts may then be suspended in alcohol and decomposed by H₂S (Möller, J. pr. [2] 22, 214). Succinyl-di-cyan-di-amide C₂H₄(CO.NHCy)₂ [105°] crystallises in monoclinic pyramids (containing 2aq), decomposed by warm water into succinic

acid and cyanamide. It may also be prepared acid and cyanamide. It may also be prepared by the action of NaNH.Cy on succinyl chloride, and by warming C₂H₄(CO)₂NCy with cyanamide. The salt C₂H₄(CO.NAgCy)₂ and crystallises in plates, sol. NH₂Aq.—Et₂A"5TiCl₄ (Demarcay, C. R. 70, 1414).—Et₂A"2TiCl₄.—Et₂A"TiCl₄.

Ethyl propyl ether EtPrA". (231°). S.G. § 1.0387. S.V. 230.2 (Wiens, A. 253, 300). Yields, on saponification by baryta, salts of EtHA", PrHA", and H₂A" (Lossen a. Köhler, A. 262, 2011.

A. 262, 201).

Ethyl butyl ether Et(C,H_o)A" (247°). S.G. § 1.0218. S.V. 255.9 (W.).

Ethyl heptyl ether Et(C,H₁₈)A". (291.4°). S.G. § 9850. S.V. 832.9 (W.). n. Propyl ether Pr₂A" (247.1°). (Wiens, A. 253, 300); (250.8°) (Perkin, C. J. 53, 561). S.G. $\frac{9}{6}$ 1.0189 (W.); $\frac{4}{5}$ 1.0157; $\frac{15}{16}$ 1.0062; $\frac{25}{25}$ 9986 (P.). S.V. 257.8. Formed by passing HCl into succinic acid in propyl alcohol.

Isopropyl ether Pr.A". (228°). S.G. 2

1.009 (Silva, A. 154, 255).

Propyl butyl ether Pr(C,H,)A". (258.7°). S.G. § 1 0106. S.V. 277.8 (Wiens, A. 253, 300).

Isobutyl ether (PrCH.), A". (265° cor.).

S.G. $\frac{15}{15}$ 9737; $\frac{25}{25}$ 9667. M.M. 12·707 at 14·5° (Perkin, C. J. 45, 519).

Isoamyl ether (C₂H₁₁)₂A". (290° cor. at 728 mm.). S.G. ¹² ·961. Formed from Ag₂A" and isoamyl bromide (Del Zanna a. Guareschi, Atti Real. Instit. Veneto [5] 6; B. 12, 1699).

yl ether (C,H₁₆),A" (350°). S.V. 459.6 (Wiens). Heptyl

2 ·9519.

Cetyl ether (O16H31)2A". [58°]. Formed by neating the acid with cetyl alcohol (Tütscheff, Rep. Chim. pure, 2, 463). Plates, sl. sol. alcohol, v. sol. ether.

Ethylene ether C2H4A". [c. 90°]. By heating succinic acid with glycol at 170° there is formed crystalline (CO₂H.C₂H₄,CO₂)₂C₂H₄ melting below 100°, which at 800° yields C₂H₄A" (Lourenco. A. 115, 858). Crystalline, insol. water and ether,

sol. hot alcohol. Decomposed by distillation.

Di-phenyl ether C₂H₄(CO₂Ph)₂. [119°]. (830°). Formed from succinic acid, phenol and POCI, (Rasiński, J. pr. [2] 26, 63). Got also from succinyl chloride and phenol (Weselsky, B. 2, 519). Completely decomposed by slowly heating (Anschütz, C. J. 47, 898). NaSEt gives di-thiosuccinic ether. Pearly plates (from alcohol).

Di-benzyl ether (CH2Ph)2A". [42°]. Formed from Ag,A" and benzyl bromide (Zanna a.

Guareschi, G. 11, 255). Lamine.

Chloride C₂H₄ (CO.Cl)₂ or C₂H₄ < COl₂>0. Mol. w. 155. (190°-200°). S.G. 1.39. Formed from succinic anhydride and PCl, (Gerhardt a.

Chiozza, A. 87, 293).

Preparation.—Succinic acid is heated with PCl, for two days with an inverted condenser in an oil bath. The POCl, is distilled off (below 120°) and the residue shaken with benzoline that has been dried over lime. The benzoline dissolves the rest of the POCl, and the succinyl The operachloride separates as a lower layer. tion is repeated until the oil no longer forms phosphoric acid when mixed with water (H. Möller, J. pr. [2] 22, 208).

Properties.—Solidifies at 0°.
Reactions.—1. Reduced to butyrolactone by sodium amalgam (8 p.c. Na) acting upon its

ethereal solution mixed with glacial acetic acid. -2. Benzene and aluminium chloride give $C_2H_4 < CO^{CPh_2} > O$ [90°], $C_2H_4(CO, C_4H_5)_2$ [134°], and a little β -benzoyl-propionic acid (Auger, A. Ch. [6] 22, 812; Bl. [2] 49, 845).-3. ZnEt, gives C₂H₄ C_C O.-4. Na₂S gives succinyl sulphide.—5. PCl, at 280° gives off HCl and the product, after treatment with water, contains C₄Cl₆O (199°-215°) which solidifies below 0°. Hot conc. H₂SO₄ dissolves C₄Cl₆O and the product, diluted with water, yields on evaporation hygroscopic crystals, which on sublimation give non-hygroscopic plates of C,Cl2O, [120°] (Kauder, J. pr. [2] 28, 191). Di-chloro-maleïc chloride is also a product of the action of PCl, on succinyl chloride.-6. Chlorine passed into boiling succinyl chloride forms chlorides of fumaric, chlorocinyl chloride forms chlorides of tumaric, chlorides fumaric, and di-chloro-maletic acids (Kauder, J. pr. [2] 31, 24).—7. C_aH.,SO₂NBzAg forms C₂H.,(CO.NBz.SO₂.C_aH.)₂ [146°] (G. a. C.).—8. Phenyl-hydrazine forms the compound C₂H.,(CO.NH.NHPh)₂ [218°] (Freund, B. 21, 2462; Fischer, B. 22, 2728), whence phosgene product [21, 128] (CO.NH.NB.) (CO.NH.NB duces C.H. (CO.N.Ph:CO), [225°]. Sodium phenylhydrazine in benzene forms C₂H₄ CO.NPh [199°], which gives C₂H₄ CO.NPh CO.NAc (Michaelis a. Hermans, B. 25, 2751).-9. Cyanamide in ethereal solution forms, on warming, succin-cyanimide C2H4(CO)2NCy [138°] orystal.

lising from alcohol in plates (Möller, J. pr [2] 22, 207).-10. Cyano-acetic ether in Et,O at 100° forms $O_2H_4: C_2O_2: CCy.CO_2$ Et [126°] (Muller, C.R.112, 1140).—11. Urea at 65° forms the compound C₂H₄(CO.NH.CO.NH₂)₂, a powder, v. sl. sol. hot water (Conrad, J. pr. [2] 9, 301).

Anhydride C₂H₄<CO>O. Mol. w. 100. [119°]. (261°) (Krafft a. Noerdlinger, B. 22, 816). Formation.—I. By rapidly boiling succinic acid.

2. By distilling the acid with P₂O₄ (D'Arcet,

A. Ch. [2] 58, 282), with PCl₅ (Gerhardt a.
Chiozza, C. R. 36, 1050), or with BzCl (Kraut, A. 137, 254).-3. By warming succinyl chloride with dry oxalic acid (Anschütz, A. 226, 16). 4. By heating succinic acid (100 g.) with POCl. (65 g.) at 120° (Volhard, A. 242, 150).-5. By distilling succinyl chloride (180 g.) with succinic acid (137 g.), and crystallising from alcohol (H. Möller, J. pr. [2] 22, 194). -6. By the action of dry Pb(NOs)2 (also nitrates of other heavy metals) upon succinyl chloride; PbCl2 is produced, and N₂O₄ and oxygen evolved; yield c. 60 p.c. of theoretical (Lachowicz, B. 18, 2990).

Properties .- Long trimetric needles (from alcohol); a:b:c = .595:1:.462 (Bodewig, B. 14, 2788). Sl. sol. water, v. sl. sol. ether. Converted by dry NH, into succinimide. Slowly converted by boiling alcohol into succinic ether. Phenylhydrazine reacts in alcoholic solution, forming CO2H.C2H,.CO.NH.NHPh [120°], converted by heat into C₂H₄<CO>N.NHPh [155°] (Auger,

A. Ch. [6] 22, 339; cf. this vol. p. 45) or [158°] (Michaelis, B. 25, 2750). Potassium cyanamide forms 'succincyanamic' acid, the compound CO.H.C.H., CO.NHCy [128°], which is v. e. sol. water, v. sol. alcohol and other, is split up by

dilute acids into succinic acid and cyanamide, and yields the following six salts Na,A"5aq, K,A"aq, CaA"4aq, BaA"2aq, AgHA", and Ag,A" (Möller, J. pr. [2] 22, 193). Urea heated with succinic anhydride at 125° forms succinuric acid CO2H.C2H4.CO.NH.CO.NH2 [205°] (Pike, B. 6, 1104).

Nitrile C₂H₄.(CN)₂. Ethylene cyanide. Mol. w. 80. [55°]. (147° at 10 mm.) (F.); (185° at 60 mm.) (Biltz, B. 25, 2541). H.C.p. 546,100. H.F. (from diamond) - 32,000 (Berthelot a. Petit, C. R. 108, 1217). Formed by boiling ethylene bromide (300 g.) with alcohol (500 g.) and KCy (200 g.) for two hours with inverted condenser (Maxwell Simpson, Pr. 10, 574; Geuther, A. 120, 268; Fauconnier, Bl. [2] 50, 214; Nevolé a. Tscherniak, C. R. 86, 1411). Amorphous, sometimes crystalline, solid, v. sol. water, alcohol, and chloroform; sl. sol. ether. Decomposed by distillation under atmospheric pressure. Converted into succinic acid by boiling alcoholic potash. Aniline hydrochloride at 200° reacts, forming di-phenyl-succinimidine $C_2H_4 < C(NH) > NPh$ (Blochmann, B. 20, 1856). $-C_2H_4Cy_24AgNO_3$: tables, sol. water and alcohol,

insol. ether. Amide-nitrile CN.C2H4.CO.NH2. Formed by heating the nitrile with alcoholic potash at 110° (Drouin, C. R. 108, 675). Crystals, sl. sol. Slowly decomposes at alcohol, insol. ether. 210°-220°.

Amide C₂H₄(CO.NH₂)₂. Succinamide. Mol. w. 116. [243°]. S. 625° at 9° (Henry, C. R. 100, 943); 11 at 100° (F.). Formed by the action of aqueous NH, on the ether and on the chloride (D'Arcet, A. 16, 215; Fehling, A. 49, 196). Formed also by warming succinimide with alcoholic NH, (Menschutkin, A. 162, 165, 187). Needles, insol. ether and alcohol. Decomposed at 200° into NH, and succinimide or by saturating at -15° with gaseous NH₃, and heating to 150° in sealed tubes (Roubtzoff, Bl. [2] 45, 250) .-HgC,H₈N₂O₂ 1½aq. White powder, deposited on cooling from a solution of HgO in hot aqueous succinamide.

Iso-amide $C_2H_4 < CO \longrightarrow 0$. [c. 95°]. Formed to the extent of 8 p.c. in the preparation of the amide from succinyl chloride and NH,Aq (Auger, A. Ch. [6] 22, 312). Hygroscopic mass. Its aqueous solution gives with AgNO, a pp. of C₂H₄(CO)₂NAg.

Methylamide C.H. (CO.NHMe)2. [175°]. Plates (Wallach a. Kamenski, B. 14, 170).

Dimethylamide O.H.(CO.NMe.)2. [81°]. Formed from succinyl chloride and NHMe2 in ether (Franchimont, R. T. C. 4, 202). Crystals (from ether), v. sol. water.
Benzyl-di-amide

C₂H₄(CONH₂)(CONHCH,Ph). [189°]. Formed from the benzylimide and ammonia at 100° (Werner, C. J. 55, 633). Minute prisms, sl. sol. ether and hot benzene. Yields the benzylimide when heated.

Di-benzyl-diamide $C_2H_1(CO.NHCH_2Ph)_2$. [206°]. Formed from succinic ether and benzylamine in alcohol. Thin plates, not decomposed by boiling NaOHAq.

Anilide O.H. (CO.NHPh). [227°]. Formed water (Menschutkin, by boiling succinic acid with aniline (Laurent a. crystals, v. sol. water.

Gerhardt, A. Ch. [8] 24, 179; A. 68, 27; Menschutkin, A. 162, 187). Needles (from alcohol), insol. water. Not affected by boiling alcoholic potash or by nitrous acid. Gives the phenylimide on distillation. Cold fuming HNO. converts it into C.H. (CO.NHC, H. (NO.)[1:4]), [260°] (Hübner, A. 209, 377). Benzoic aldehyde at 180° forms succinic phenylimide and benzylidene-aniline (Schiff, A. 148, 338).

Amide-anilide CO(NH2).C2H4.CONHPh. [181°]. Formed by heating the phenylimide with alcoholic NH₃ at 100° (Menschutkin, A. 162, 182). Broad needles (from water), sl. sol. hot alcohol. Alkaline KOBr followed by heating with HOAc forms C2H4:C2O2:(NHBr)(NHPh) crystallising in minute needles, converted by boiling alcohol into C.H., C.O. (NH.) (NHC.H.,Br.) [215°], whence potash forms the compound CO.H.C.H.,CO.NHC.H.,Br[1:4] [187°] (Hoogewerff a. Van Dorp, R. T. C. 9, 41).

Di-(a)-naphthyl-di-amide

C₂H₄(CO.NHC₁₀H₂)₂. [285°]. Needles (from HOAo). Converted by fuming nitric acid into C₂H₄(CO.NH.C₁₀H₅(NO₂)₂)₂, [225°], and C₂H₄(CO.NH.C₁₀H₂(NO₂)₄)₂ [256°].

Tetra-phenyl-di-amide

7. (CO.NDb.) [284°] (Pintti, G. 14, 467).

[234°] (Piutti, G. 14, 467). $C_2H_4(CO.NPh_2)_2$.

Needles (from alcohol).

o-Tolyl-diamide C₂H₄(CONH₂).CO.NHC,H₇. [160°]. Formed by heating the o-tolylimide with alcoholic NH, at 100°. Plates.

Di-o-tolyl-diamide C2H4(CO.NHC7H7)2. [100°]. White needles (Bechi, B. 12, 25, 321). V. sl. sol. water, m. sol. alcohol.

p-Tolyl-di-amide C₂H₄(CONH₂).CONHC₇H₇. [148°]. Formed by heating the p-tolylimide with alcoholic NH.

Di-p-tolyl-diamide C2H4(CO.NHC,H7)2. [256°]. Plates (from alcohol), sl. sol. hot water (Bechi, B. 12, 323; Hübner, A. 209, 380). Yields on nitration C,H,(CO.NHC,H,NO,)2 [217°] and $C_2H_4(CO.NH.C_7H_5(NO_2)_2)_2$

(β)-Naphthalide C2H4(CO.NHC,H7)2. $[266^\circ]$ (Bischoff a. Reebe, **B**. 25, 3267). Got by heating succinic acid (10 g.) with (β) -naphthylamine (24 g.). Insol. ordinary solvents, sol.

H₂SO₄.

Amic acid CO2H.C2H2.CO.NH2. Succinamic acid. [157°]. Formed by warming succinimide with an equivalent quantity of baryta water (Teuchert, A. 134, 136) or milk of lime (Menschutkin, A. 162, 175). Formed also from nitrosoglutaric acid by heating alone or with Ac₂O (Serda a. Wiedemann, B. 23, 8284; Wolff, A. 260, 114). Needles, m. sol. water, insol. alcohol. Decomposed at 200° into water and succinimide. Boiling water forms acid ammonium succinate.

Salts.—KA': very hygroscopic mass (Landsberg, A. 215, 201).—BaA'₂: needles, v. sol. water.—CaA'₂.—MgA'₂ Saq: trimetric crystals.—MgA'₂ 6aq.—OdA'₂aq: prisms.—CuA'₂.—PbA'₂. Concentric needles, v. sol. water. - MnA', 5aq. ZnA' .- AgA': monoclinic crystals, v. sol. NH,Aq.

Ethyl-amic acid CO,H.C,H,.CONHEt. Formed by warming the ethylimide with baryta water (Menschutkin, A. 182, 92). — BaA's:

Ethylene-di-amic acid C10H16N2O6 i.e. C.H. (NH.CO.C.H., CO.H.). Ethylene-disuccin-amic acid. [185]. Formed by boiling the diimide with baryta (Mason, C. J. 55, 13). Large plates, v. sol. hot water. Does not yield an oxim or aphenyl-hydrazide.—CaA" 3aq: prisms. -Ag₂A": white amorphous powder.

Bensylamic acid CO.H.C.H..CO.NHC,H,. [139°]. Formed by boiling the benzyl-imide (2 mols.) with aqueous barium hydroxide (1 mol.) (Werner, C. J. 55, 631). Flat prisms, m. sol. hot water, v. sl. sol.

ether.

Phenylamic acid CO, H.C, H, CO.NHPh. Succinanilic acid. [149°]. Formed by boiling the phenylimide with NH3Aq (Laurent a. Gerhardt, A. Ch. [8] 24, 179) or baryta (Menschutkin, A. 162, 176). Flat needles, v. sl. sol. cold water. Decomposed by heat into water and the phenylimide. Boiling alcoholic HCl forms succinic acid. — CaA', 4aq. — BaA', 3aq. — AgA': crystalline pp.

o-Tolyl-amic acid

CO₂H.C₂H₄.CO.NHC₇H₇. [97°]. Got by boiling the o-tolylimide with baryta water (Bechi, B. 12, White needles .- BaA', aq.

p-Tolyl-amic acid [157].-BaA', aq.

(a)-Naphthyl-amic acid

 $CO_2H.C_2H.CO.NHC_{10}H_7$. [171°]. Formed by heating the (a)-naphthylimide with KOHAq (Pellizzari, A. 248, 158; G. 18, 323). Needles or plates (from alcohol), v. e. sol. HOAc and

(β)-Naphthyl-amic acid [192°]. Crystals.

Di-phenyl-amic acid

CO₂H.C₂H₄.CONPh₂. [119°]. Plates (Piutti, G. 14, 468).—AgA': pp.

Phenylene-di-amic acid C₅H₄(NH.CO.C₂H₄.CO₂H)₂. Formed by heating phenylene-diamine with succinic acid at 200° (Biedermann, B. 9, 1668). Small crystals (from Melts above 360°. HOAc). Converted by KOHAq and MeI into C.H. NMe.O.CO crystallising in plates or tables (containing 1 aq)

(Griess, B. 18, 2410).

Imide CHrCO NH. [126°] (Erlenmeyer, Z. [2] 5, 175). (288°). H.C. 439,000. H.F. 110.500 (Berthelot a. André, Bl. [8] 4, 229). Formed from succinic anhydride and dry NH, (D'Arcet, A. Ch. [2] 58, 294), and also by heating succinamide or ammonium succinate (Fehling, A. 49, 198; Laurent a. Gerhardt, Compt. Chim. 1849, 108; Menschutkin, A. 162, 165, 187; 182, 93). Large efflorescent crystals (containing aq) or octahedra (from acetone) (Bunge, A. Suppl. 7, 118), v. sol. water and alcohol, m. sol. ether. Not attacked by pure HNO. (S.G. 1.53) (Franchimont, R. T. C. 6, 228). Converted by hot baryta water into succinamic acid, and finally into succinic acid. Alcoholic NH, at 100° forms succinamide. Yields pyrrole on distillation with ainc-dust. KOH (6 mols.) and KBrO (1 mol.), acting for two hours at 60°, form (\$\beta\$)-amido-propionic acid [196°] (Hoogewerff a. Van Dorp, R. T. C. 10, 4). Bromine forms bromo-succinimide [225°] and bromo-maleimide [152°] (Kisielinski, Sits. W. [2] 74, 561; cf. Kusseroff, A. 252, 158). Chlorine at 160° forms chloro- and di-chloro-VOL. IV.

at 145°-148° (Bernthsen, B. 13, 1047). A solution of bleaching-powder added to a solution of succinimide containing excess of HOAc forms C₂H₄ CO NCl [148°], which separates from benzene in large crystals, and is reconverted by acids and alkalis into succinimide (Bender, B. 19, 2273). Hydroxylamine at 65° NH(CO.C₂H₄.CO.NH.OH)₂ [171°], crystallising in needles, v. e. sol. water, and yielding a picrate B'C₂H₂N₂O, [266°] (Garny, B. 24, 3434). Methyl cyanate forms C₂H₄CO.NH₂CO [149°], while EtNCO forms corresponding C, H₁₀N₂O₃ [95°]

(Menschutkin, A. 178, 204).

Salts.-KC,H,NO, laq. Ppd. by adding alcoholic potash and ether to an alcoholic solution of succinimide (Landsberg, A. 215, 200).-Hold of succession and elochol.—

BaA', 22aq. — BaA', 2½aq. — MgA', 3aq. —

HgA', — NH₂.HgA'. — ClHgA'. — CyHgA'. —

Cu,A', (OH), 9aq. — Cu₂A', (OH), 2aq.—AgA' ½aq.

Needles. Converted by iodine into IC,H,NO₂, crystallising in dimetric prisms; a:c=1: 873, melting, with evolution of I, at 135°. Silver succinimide is converted by succinyl chloride in ethereal solution into 'trisuccinamide' (C,H,O2)2N2 [83°], crystallising from ether (Gerhardt a. Chiozza, A. 90, 108).—AgA'NH.: crystalline.

Methylimide C₂H₄:C₂O₂:NMe. (234°). Formed by distilling methylamine succinate. Plates. Got also by the action of conc. H2SO, on the oxim of \$-acetyl-propionic acid (Rischbieth; Bredt a. Boeddinghous, A. 251,

Ethylimide C₂H₄:C₂O₂:NEt. [26°]. (234°). Formed by distilling acid ethylamine succinate (M.), and by adding alcoholic EtI to sodium succinimide (Landsberg, A. 215, 212). Long lancet-shaped crystals, v. sol. water, alcohol, and ether. Yields ethyl-pyrrole on distillation over zinc-dust.

Allylimide C.H.: C.O.: NC.H. (245°). S.G. § 1·1543; 1 1·1432. Formed by heating succinic acid with allyl thiocarbimide. Liquid,

v. sol. water and alcohol.

Ethylene-imide $(C_1H_4:C_2O_2:N)_2C_2H_4.$ [251°]. (395°). Formed by heating succinio acid with the hydrate of ethylene-diamine at 180° (Mason, C. J. 55, 10). Prisms, sol. hot water, v. sl. sol. hot alcohol, insol. ether

Propylene-imide (C.H.:CO.:N)₂C.H... [100°]. Formed from the anhydride and propylene-diamine (Strache, B. 21, 2360). Crystals

(from benzene).

Phenylimide C₂H₄:C₂O₂:NPh. [156°] (L. a. G.; Kauder, J. pr. [2] cinanil. 31, 17). [150°] (Hübner, A. 209, 373; Bischoff a. Nastvogel, B. 22, 1807) (above 300°). Formed by heating succinic acid with aniline, and crystallising the product from water and alcohol successively (Laurent a. Gerhardt, A. Ch. [3] 84, 179). Formed also by the action of PCl, on C,H,(CO.NHPh).CO,H in chloroform (Anschütz, B. 21, 957). Needles, sl. sol. hot water. PCl. in presence of boiling POCl, forms the phenylimide of di-chloro-maleïc acid C,Cl,:C,O,:NPh. Alcoholic NH₂ forms C₂H₄(CONH₂).CONHPh. Boiling lime-water forms C₂H₄(CO₂H).CONHPh. malesc imide. PCl, forms a compound melting | Fuming HNO, forms an o-nitro-derivative [156°] and the p-nitro derivative [205°-208°]. The salt C₂H₄:C₂O₂:NC₄H₄:SO₂Na is got by heating succinic acid with sodium amido-benzene p-sulphonate (Pellizzari, G. 18, 317).

Bensylimide C2H4:C2O2:NCH2Ph. Formed from benzyl chloride, succinimide, and alcoholic NaOH (Werner, C. J. 55, 629). Six-

sided prisms, sol. alcohol.

o-Tolylimide C2H4:C2O2:NC6H4Me. [75°]. (345° uncor.). Formed by heating o-toluidine with succinic scid (Michael, B. 10, 579; Bechi, B. 12, 25, 321). Needles, v. sol. water.

p-Tolylimide. [150°]. (845°). Needles (from water), sl. sol. cold water. Yields on nitration C_2H_4 : C_2O_2 : $NC_2H_2Me(NO_2)[1:4:2]$

(Hübner, A. 209, 878).

Tri-methyl-phenyl-imide C₂H₄:C₂O₂:NC₆H₂Me. Succinmesidil. Formed by heating succinic acid with mesidine (Eisenberg, B. 15, 1018). Pearly plates, sol.

alcohol and ether, sl. sol. water.

(a) - Naphthylimide O.H.: C.O.: NC, BH,. [153°]. Formed by heating succinic acid with (a)-naphthylamine (Hahnemann, B. 10, 1713; Hübner, A. 209, 381; Pellizzari, A. 248, 158). Needles, m. sol. alcohol. Yields a di-nitro-[250°]. derivative The sulphonic C₂H₄:C₂O₂:NC₁₀H₆SO₂H, prepared from (a)-naphthylamine p-sulphonic acid, yields KA' 2aq (Pellizzari, G. 18, 321).

(8) Naphthylimide. [180°]. Colourless needles, v. sol. alcohol. Converted by KOHAq into CO2H.C2H4.CO.NHC10H, [192°] (Pellizzari,

A. 248, 159).

Benzimidide C.H.; C.O.; NCPh:NH. [212°]. Formed from acetyl-succinic ether C10H16O6, benzamidine hydrochloride, and NaOHAq (Pinner, B. 22, 2620). Needles, sl. sol. cold acetone and ether, sol. alcohol.

References.—Amido-, Bromo-, Bromo-Amido-, Chloro-, Nitro-, and Oxy-succinic acid and

DI-IODO-BUCCINAMIC ACID.

Isosuccinic acid v. METHYL-MALONIC ACID.

BUCCINIC ALDEHYDE. Reduction of succinic anhydride, which might be expected to yield the aldehyde, forms instead the isomeric

y-oxy-butyric lactone.

Owim C2H4(CH:NOH), [178°]. Formed by heating pyrrole with hydroxylamine hydrochloride at 100° and, together with ethylamine, by the action of hydroxylamine on v-ethylpyrrole (Ciamician, B. 17, 533; 22, 1968; 23, 1788). Small white crystals (from alcohol). May be reduced to tetramethylene-diamine

C₂H₄(CH₂.NH₂)₂ [28°].

Phenyl hydraside C,H,(CH:N,HPh), [125°]. Formed by heating a solution of the oxim (1 pt.) in water (50 pts.) with phenyl hydrazine (5 pts.) dissolved in dilute HOAc (Ciamician, B. 22, 1974; 28, 1784). Small silky plates, v. sol. alcohol. Converted by cold conc. HClAq into a base C₂₀H₂₀N₄ [185°], which crystallises from EtOAc in needles, v. sl. sol. alcohol, and gives a deep-blue colour with K,Cr,O, and H,SO,

Succinic semi-aldehyde. Nitrile CN.CH. CH. CHO. \$\beta\$-Cyanopropionic aldehyde (77°). S.G. 15 89. Formed by heating (77°). S.G. 12 ·89. Formed by heating sugars, before exhibiting this property, require CH_I.CH_CHO (92 g.) with alcohol (180 g.) and to be digested with dilute mineral acids. Enagcy (67 g) for ten hours (Chautard, A. Ch. symes also convert some of the non-reducing members into reducing ones. With phenyl-

with alcohol and ether. Reduces Fehling's solution. Does not combine with NaHSO, or phenyl-hydrazine. Decomposed by hot alkalis and acids. Aniline at 850° forms the compound CH2Cy.CH(NHPh)2.

Succinimidine C.H.N. i.e.

 $C_2H_4 < C(NH) > NH$. Formed, as hydrochloride, together with NH Cl, by the action of water on succinamidine (Pinner, B. 16, 362, 1657; 18, 2845).—B"HCl: long plates, v. sol. water, sl. sol. alcohol. Converted by acetoacetic ether into crystalline $C_0H_{11}N_3O_2$.— $B''(HNO_3)_2\frac{1}{2}aq$: leaflets (Grabowski, A. 265, 168).— $AgC_4H_6N_3$: ppd. by adding ammoniacal AgNO, to a solution of the hydrochloride.

SUCCINIMIDO-ACETIC ETHER C.H., NO. i.e. C₂H₄:C₂O₂:N.OH₂:CO₂Et. [67°]. Formed by heating succinimide with alcohol, NaOEt, and CH₂Cl.CO₂Et (Haller a. Arth, C. R. 105, 280). Needles, v. sol. water, alcohol, and ether. holic NaOEt forms gelatinous C,H10NaNO4.

SUCCINIMIDO-ETHYL ETHER C.H. N.O. i.e. C.H. (C(NH).OEt). The hydrochloride B"2HCl is prepared by passing gaseous HCl into an ethereal solution of ethylene cyanide (Pinner, B. 16, 359). It is v. sl. sol. alcohol and ether, and decomposed by water into succinic ether and NH₄Ol. With ammonia it gives $C_2H_1(C(NH).NH_2)_2$.

 $C_2H_4 < C(NOH) > NH.$ SUCCINIMIDOXIM [197°]. Formed by heating ethylene cyanide with alcoholic hydroxylamine at 65° (Garny, B. 24, 3427). White crystals, sol. water.—B'HOl. [98°]. Needles.—B'C₆H₂N₃O₇. [212°].

Benzoyl derivative

 $C_2H_4 < \frac{C(NOBz)}{CO} > NH.$ [184°]. Formed from the oxim, NaOH, and BzCl. White powder, sol. alcohol, sl. sol. ether.

SUCCINOPHENONE v. DI-PHENYL ETHYLENE

DIKETONE.

SUCCINOXYL-AMIDO-BENZOIC ACID CARBOXY-PHENYL-SUCCINAMIC ACID.

SUCCINYL-AMIDO-BENZOIC ACID AMIDO-BENZOIC ACID.

SUCCINYL - SUCCINIC ACID v. DI-OXY-TEREPHTHALIC ACID DIHYDRIDE, vol. iii. p. 777.

SUCCISTERENE C. A. [160°]. 300°). Occurs in the product of distillation of amber (Pelletier a. Walter, A. Ch. [3] 9, 96). Flat needles, nearly insol. cold alcohol, v. sl. sol. ether.

SUGAR, a term applied to the members of a group of carbohydrates in consequence of the property they possess of being sweet to the taste and of their relationship to the substance to which the term was originally given. The known natural members of this group are not very numerous. They are all easily soluble in water, crystallisable, and diffusible. When solutions of them are heated with alkaline solutions of the more easily reducible metals, reduction takes place with precipitation of the lower oxides in some cases, and of the metals in others; some

hydrasine they yield phenylhydrasides and phenylosasones. Oxidising agents easily convert them into soids, and reducing agents into alcohols. Some of them undergo fermentation when submitted to the action of beer yeast, i.e. they are decomposed into alcohol, CO₂, &c. They yield solutions which are optically active, those of some rotating the plane of polarisation to the right, of others to the left. They give colour reactions with aromatic acids and phenols.

The group may be divided into two classes: (1) those which, when digested with dilute acids, do not yield any other sugar or sugars—the sugars of this class are distinguished by the termination -ose; (2) those which, when digested with dilute acids, do yield some other sugar or sugars—these sugars are distinguished by the termination -on. They are designated according to the number of carbon atoms they contain: thus, pentose containing C, hexose containing C_6 , &c., and di-penton containing C_{16} , di-hexon containing C12, &c.

The members of the -ose class are:

Triose C₂H₆O₃, glycerose

Tetroses C,H,O,, erythrose C,H,(C,H,)O,, phenyltetrose

Pentoses C.H.O., arabinose

ribose

99

xylose

CaHa(CHa)Oa, rhamnose (methyl

pentose) fucose (ditto)

Hexoses C₈H₁₂O₈, d- l- and i-1 glucose (dglucose = dextrose)

d- l- and i-mannose

d- l- and i-gulose

d- l- and i-fructose (d-

fructose = lævulose) d- l- and i-galactose

sorbose (sorbinose)

formose

lokaose

B-acrose C,H, (CH,)O, rhamnohexose (methyl hexose).

Eleptoses C,H,O, mannoheptose

a- and \$-glucoheptose

galaheptose

fructoheptose

digitalose

C,H,a(CH,)O, rhamnoheptose (methyl heptose).

Octoses C₈H₁₆O₈, mannoctose a- and \$-glucoctose

Nonoses C, H1, O, mannononose

glucononose

The members of the -on group are:

Di-penton $C_{10}H_{10}O_{9}$, arabinon Di-hexon $C_{12}H_{21}O_{11}$, cane-sugar (saccharon)

milk-sugar, lactose (lacton)

(di-glucon, maltose

amylon)

iso-maltose

melibiose

trehalose

cyclamose para- saccharose

tewfikose (tewfikon)

" d , L and i. Pischer's notation, see p. 583.

Tri-hexon C15H22O16, raffinose melezitose Hex-hexon C_{se}H_{ex}O_{s1}, gentianose stachyose

Occurrence.-Some of these sugars have been obtained by synthesis, others are products of the action of dilute acids or ferments on more complicated bodies, and others occur naturally. The most important source of the latter is the vegetable kingdom. Little, if anything, is known concerning the formation of the sugars in nature; it is very probable that they are intermediate products of assimilation between CO, and H,O, and starch, inulin, cellulose, and such bodies. For an account of the various theories, see textbooks of vegetable and animal physiology. sugars found in animals are most probably of vegetable origin.

Determination. - Owing to the fact that the sugars possess many properties in common, the presence of a particular sugar can only be defi-nitely proved by its isolation in the pure state, and a determination of its optical activity, reducing power, and other definite distinguishing

properties (v. end of art.)

Molecular Weight.—Until quite recently none of the accepted methods for determining molecular weights were applicable to the sugars. At the present time there are several, chief among which are the physical methods of Raoult (A. Ch. [5] 28, 133; [6] 2, 66-124; [6] 4, 401; [6] 8, 289 a. 817), and De Vries, which are of general application, and the various chemical ones which are of special application. By Raoult's method Brown and Morris (C. J. 1888, 610; 1889, 462), Tollens and Mayer (B.21, 1566), and Tollens, Mayer, and Wheeler (B. 21, 3508), Ekstrand and Mauzelius (Vetensk. Akad. für Handl. 1889. 157), and O'Sullivan (C. J. 1890. 62) have determined the molecular weight of xylose and arabinose to be 150, corresponding to a formula C, H, O,; of dextrose, lævulose, galactose to be 180, corresponding to a formula C₆H₁₂O₆; of arabinon to be 282, corresponding to a formula CieHisOe; of cane sugar, maltose, lactose to be 342, corresponding to a formula C₁₂H₂₂O₁₁; and of rashinose to be 504, corresponding to a formula CieHarOis. These numbers have been confirmed in the case of the simpler sugars by various chemical methods (see succeeding paragraphs). In the case of arabinose it was first shown to be CaH10Oa by Kiliani (B. 20, 339) by the analysis of some of its compounds, its formula previously having been con-

sidered to be C_eH₁₂O_e.

Synthesis.—The first step towards the synthesis of the sugars was made by Butlerow (A. 120, 295; C. R. 53, 145). He obtained a sweet syrup, having the common characteristics of the sugars, by adding lime-water to a hot solution of di-oxymethylene; the product he called methylenitan, and considered its formula to be C, H,4O. The next step was made by Loew (J. pr. 33, 321), who, having discovered a method for the preparation of formaldehyde (CH,O) in quantity, investigated its condensation by lime-water. He obtained a sweet syrup, which he called formoss, and considered it to have a formula $C_eH_{12}O_e$. Neither methylenitan nor formose fermented with yeast (see also Tollens, B. 19, 2188). Fischer (B. 21, 989) finds these products to be a mixture of various saccharine bodies,

the chief which is a sugar, formose $C_0H_{12}O_4$, yielding an osazone $C_{10}H_{22}N_4O_4$, m.p. 144°. Another sugar is present in smaller quantities in the condensation products of formaldehyde. It yields an osazone greatly resembling glucosazone, which has been identified with a-acrosazone. Later, Loew (B. 22, 475), by the condensation of formaldehyde with lead oxide and magnesia, obtained a sugar which is fermentable. Loew called this methose, but Fischer has shown it to be identical with a-acrose.

Acrose was obtained by Fischer (B. 20, 1093 and 2566) by acting on acrolein bromide with bases $2C_2H_4OBr_2 + 2Ba(OH)_2 = C_6H_{12}O_6 + 2BaBr_2$. Two isomeric sugars, α -acrose and β -acrose, are

thus produced.

Glycerose yields these two sugars by the action of alkalis. (Glycerose is a sugar C.H.O., obtained by the action of bromine and soda on glycerol and other methods by Van Deen, J. 1863. 501; Grimaux, C. R. 104, 1276; and Fischer a. Tafel, B. 20, 1088 and 3385; 22, 106. It is probably a mixture of the aldehyde and ketone of glycerol.) The formation of α-acrose from glycerose may be represented CH_{*}(OH).CH(OH).COH+CH_{*}OH.CO.CH₂OH= CH_OH.CH(OH).CH(OH).CH(OH).CO.CH_OH. The sugars can only be separated from the products thus obtained as osazones; a-acrosazone is identical with glucosazone in every way except in its action on polarised light.

a-acrosazone CH_OH(CH.OH)_C-CH.N.NHC_H_

N.NH.C.H.

acted on by fuming hydrochloric acid, is converted into a-acrosone CH, OH. (CH.OH), CO.COH, which, when reduced by zinc and acetic acid, yields a sugar CH, OH(CH.OH), CO.CH, OH, in the form of a sweet syrup; it ferments with yeast, yields lævulinic acid when heated with hydrochloric acid, and is reduced by sodiumamalgam to a hexahydric alcohol a-acritol, which resembles mannitol in every way except that it is optically inactive. By these various reactions so much is lost that from 1 kilo. glycerol only 0.2 g. acritol is obtained. Further, all the bodies obtained in the processes were optically inactive. More knowledge was necessary before the natural sugars could be synthesised. At this stage a new light was thrown on the subject by an observation of Fischer (B. 23, 370) that arabinose carboxylic acid and mannonic acid were identical in every way except that their optical activities, although equal, were opposite in sign, and that they combined to form an optically inactive acid. These three acids were reduced by sodium-amalgam to three sugars resembling one another in all respects except that one of them had a right-handed optical activity, the other an equal but left-handed one, while the third was optically inactive. They are named l-mannose, i-mannose, and d-mannose; by further reduction they yielded the corresponding hexahydric alcohols I-mannitol, i-mannitol, and d-mannitol.

The sugar obtained from a-acrosazone as

above described is identical with lavulose except that it is inactive. If, then, we can decompose this lævulose into d-lævulose and l-lævulose, the synthesis of the natural sugar will be accomplished. This Fischer has done (B. 23, 370). If i-lævulose be submitted to the action of yeast, the lævo- constituent ferments and the residue is dextrorotatory. This is l-levulose; it is not the natural sugar: that belongs to the d-group, and was destroyed by the ferment.

To obtain the natural sugar from the inactive synthetic one, Fischer proceeded as follows:—i-mannitol, i.e. a-acritol, is oxidised by nitric acid to i-mannose, and this further by bromine-water to i-mannonic acid. By fractional crystallisation of the strychnine or morphine salts it can be mesotomised, the result being dand l- mannonic salts, which, on being freed from the base, yield by reduction the corresponding mannoses and mannitols. These, by means of the osazones, can be converted into the corresponding lævuloses (see Dextrosazone).

Dextrose and mannose both yield the same osazone; the difference in their constitution must therefore be caused by the position of the group marked with a * in the formula

ОН. (ОН).СН(ОН).СН(ОН).СН(ОН).ОН(ОН).ООН. Fischer (B.23,799 a. 2611) found that by heating gluconic acid with quinoline, part was converted into mannonic soid and part unaltered; and also that mannonic acid, treated in the same way, yielded some gluconic acid. Dextrose may then be obtained by reducing the gluconic acid thus obtained. The corresponding reaction with I-mannonic acid does not take place easily, if at all, but l-gluconic acid is produced simultaneously with arabinose carboxylic acid (l-mannonic acid) by Kiliani's mode of preparation (see Arabinose). From I-gluconic acid, by reduction, I-glucose, the optical isomer of dextrose, is obtained. The table on next page gives a concise view of the synthesis of dextrose, lævulose, and mannose.

Another sugar, gulose, stereoisomeric with glucose, has been obtained by Fischer (B. 23, 93; 24, 521) by the reduction of saccharic acid, the d-, l- and i- modifications being obtained by the reduction of d-, l- and i- saccharic acids. As these are the products of the oxidation of d-land i- glucose, the synthesis of gulose is com-

plete.

Galactose has not been synthesised up to the present (June 1893), but by the reduction of mucic acid Fischer and Hertz (B. 25, 1247) have obtained i-galactonic acid and i-galactose; i-galactonic acid, by the fractional crystallisation of its strychnine salt, may be mesotomised into d- and l-galactonic acids from which d- and l-galactose are obtained; d-galactose is identical with ordinary galactose.

Arabinose has not been directly synthesised, but its relationship to the synthetical sugars is shown by the fact that the two acids obtained from arabinose by the nitrile reaction are I-mannonic acid and l-gluconic acid (Fischer, B. 23,

2611; 24, 539).

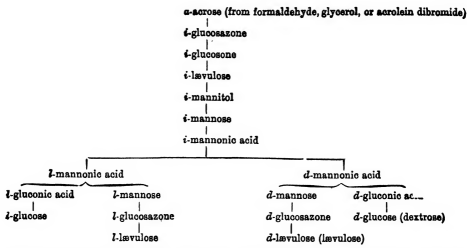
Of xylose the same must be said; but by the nitrile reaction it yields l-gulonic acid (Fischer

a. Stahel, B. 24, 528).

Arabonic acid, when heated with quinoline, is partly changed into its stereoisomeride ribonic

^{17, 4,} and d are very unsatisfactory, and somewhat mis'eading; but as Fischer has introduced them we must let them stand. A little study will make their meaning

SUGAR. \$33



acid, just as gluconic acid yields gulonic acid; ribonic acid, by reduction, yields the stereoisomeride of arabinose, which it is proposed to call

The synthetical passage from an -ose sugar to an -on one is said to be accomplished for lactose (lacton) and cane sugar (saccharon) as well as for maltose (amylon).

Lacton. A mixture of dextrose and galactose, is acted on by acetic anhydride, and the resulting products, carefully saponified with alkali, yield lacton (Demole, C. R. 89, 481). This has been contradicted by Berthelot (Bl. [2] 84, 82) and by Herzfeld (A. 220, 219).

Sucron. A body having a dextrorotatory power, and by inversion becoming lævorotatory, was obtained by acting on aceto-chlorhydrose (a derivative of dextrose) with an alcoholic solution of lævulose in the presence of barium carbonate. Another body which was lævorotatory, and became less lævorotatory on inversion, was obtained by acting on aceto-chlorhydrose by sodium lævulosate (Colley a. Vakovitch, Bl. [2] 84, 826).

Isomaltose has been prepared by Fischer (B. 23, 8687) by the action of HCl on dextrose. Dilute acids again hydrolyse it to dextrose, the same product as is yielded by maltose.

Phenyl-tetrose has been prepared synthetically from cinnamaldehyde cyanhydrin; this, by the action of bromine, yielding phenyl-dibromovy-butyronitrile CH.Ph.Br.CH.Br.CH.OH.CN, which, when heated with hydrochloric acid, yields phenyl-bromo-di-oxy-butyrolactone, from which the corresponding acid OH.CH.Ph.CHBr.CH.OH.COOH is easily obtained, and which on reduction yields the sugar OH.CH.Ph.CH(OH).CH(OH).COH.

It will not be out of place here to indicate what has been done in the way of converting a sugar into one containing a carbon atom more. To do this, Fischer (B. 22, 2204; 23, 370, 799, 930, 2226, and 3102; Ann. 270, 64) made use of the well-known reaction for proceeding from one alcohol to its next higher homologue by means of the nitrile. Now, in this reaction, where an asymmetric carbon atom is added to the molecule.

Van't Hoff has predicted that two isomeric bodies will be formed which will differ by the arrangement of the groups round the new carbon atoms. Fischer (Ann. 270, 64) has shown that this is so in the case of the glucose homologues, the two acids obtained from glucose being a- and B-glucoheptonic acids; a-glucoheptonic acid is always the principal product of the reaction, and is identical with the dextrose carboxylic acid described by Kiliani (see Dextrose). When the reaction takes place at 20°-25° hardly a trace of the \$- acid is formed; but at 40° about 18 p.c. of the latter is obtained. By oxidation these acids yield two penta-oxy-pimelic acids, the a- acid being optically inactive, and by reduction the two sugars a- and β-glucoheptose are obtained.

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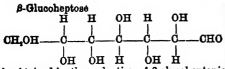
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a-Glucoheptose

separates from its aqueous solution in rhombic prisms, m.p. 180°-190°; they have a faintly sweet taste, dissolve in 10.5 pts. of water at 14°, and are easily soluble in hot water and sparingly in hot alcohol. Its solution is optically active. Freshly-prepared solutions show a slight bi-rotation; for a 10 p.c. solution, $[\alpha]_{D=0} = -19.7^{\circ}$. It does not ferment with yeast, but reduces Fehling slightly less than dextrose. It is oxidised by bromine to a-glucoheptonic acid. It yields a hydrazone C,H,,O,N,HC,H,, very soluble in water, m.p. 170°, and also an osazone C,H₁₂O₃(N₂HC₃H₃)₂, forming yellow needles, m.p. 195°, almost insoluble in water and difficultly in hot alcohol. Treated with acetic anhydride and zinc chloride it yields a hexacetate, m.p. 156°, and with sodium acetate and acetic anhydride it yields dec-acetyl-diglucoheptose C14H16(C2H2O), O121 m.p. 181°-182°.

By reduction with sodium-amalgam, a-glucoheptose yields a-glucoheptitol, which crystallises in delicate prisms, m.p. 127°–128°, is optically inactive, easily soluble in water and sparingly in alcohol. It yields a heptscetyl derivative CH.(C.H.(C.).



is obtained by the reduction of β -glucoheptonic acid, it has not yet been obtained in a crystalline state, it forms a phenyl-hydrazide $C_1H_{14}O_4N_2H.C_6H_5$, crystallising from alcohol in slender needles, m.p. 192°; the osazone is identi-

cal with a-glucoheptosazone.

Just as dextrose yields two isomeric gluco-heptonic acids, so a-glucoheptose yields two isomeric glucoctonic acids. The a-acid is the chief product, and the amount of β -acid formed varies with the temperature. a-Glucoctonic acid yields a lactone $C_0H_{14}O_9$, sparingly soluble in alcohol and readily in water, m.p. $145^\circ-147^\circ$ [a]_{D20}° = $-45^\circ9$; by reduction with sodium-amalgamit yields a-glucoctose C.H.O. 2H.O. which crystallises in colourless needles, m.p. 93°. Its aqueous solutions are optically active and show bi-rotation, [a] mass = -50.5° for the anhydrous sugar. It yields a phenyl hydrazone C,H₁,O, (N,HC,H₃), m.p. 190°, and an osazone C,H₁,O, (N,HC,H₃), m.p. 210°–212°, almost insoluble in water. By reduction with sodium-amalgam it yields a-glucoctitol C.H. O., which is easily soluble in water; it forms slender white needles, m.p. 141°. From a-glucoctose two glucononic acids may be prepared. Only one has been examined; this when reduced yields glucononose C,H18O, its phenylhydrazide C₀H₁₈O₈N₂HC₆H₅ is sparingly soluble in cold water and alcohol, m.p. 195°-200°, its phenyl osazone C₀H₁₈O₂(N₂HC₆H₅)₂ is sparingly soluble in hot water and alcohol, m.p. 220°-223°. Unlike mannononose, glucononose does not ferment with yeast. By reduction glucononitol is obtained.

Mannoheptose $C_1H_{14}O_7$ is obtained by the reduction of mannoheptonic acid. It crystallises from alcohol in fine needles, it has a sweet taste, m.p. $134^\circ-135^\circ$. It is easily soluble in water, and does not ferment with yeast; its solution is dextrorotatory and exhibits bi-rotation, the constant value is $[a]_D = +68^\circ6^\circ$. It yields a difficultly soluble hydrazide $C_1H_1, O_a, N_2H.C_aH_3$, m.p. $197^\circ-200^\circ$, and an osazone $C_1H_{12}O_3(N_1H.C_aH_3)$, m.p. 200° . By reduction with sodium-amalgam it yields a heptahydric alcohol which is identical with the naturally produced body, perseit (Maquenne, A. Ch. [6] 19, 1; C. R. 107, 583). It combines with hydrocyanic acid, and the compound decomposed by hydrochloric acid yields mannoctonic acid.

Mannoctose $C_sH_{1s}O_s$ is obtained by the reduction of mannoctonic acid. It is easily soluble in water, has a sweet taste, and does not ferment with yeast. Its aqueous solutions are laworotatory, $[a]_D = -3.3^{\circ}$. It yields a hydrazone $C_sH_{1s}O_s(N_sH.C_sH_s)$, difficultly soluble in water, m.p. 212°; and an osazone $C_sH_{1s}O_s(N_sH.C_sH_s)$, m.p. 223°. By reduction it yields the octohydric alcohol mannoctitol. It combines with hydrocyanic acid, and the compound decomposed by hydrochloric acid yields mannononnic acid.

Mannononose $C_pH_{1s}O_p$ is obtained by the reduction of mannonononic acid. It is easily soluble in water and crystallises from alcohol, m.p. about 180°, its solution is dextrorotatory, $[\alpha]_p$

= +50° approx. It ferments readily and completely with yeast, and resembles dextrose in many ways. It yields a difficultly soluble hydrazide C₂H₁₀O₈.N₂H.C₆H₈, m.p. 228°, and an osazone, m.p. 217°.

Galactoheptose C,H,O, is obtained by the reduction of galacto-heptoic acid. Its hydrazide (m.p. 199°) is sparingly soluble; so is its osazone

(m.p. 220°).

Fructoheptose is obtained from lævulose by

the same set of reactions.

Rhamnohexose CH3.(CH.OH)5COH is obtained by the reduction of rhamnohexolactone (Fischer a. Piloty, B. 23, 3102); it does not ferment with yeast, but exhibits all the characters of a sugar; the osazone is obtained in stellate groups of needles, m.p. 200°. On reyields rhamnohexitol the sugar CH,(CH.OH),CH2OH; by the action of hydrocyanic acid on rhamnohexose, the lactone of rhamnoheptonic acid is obtained, which on reduction yields rhamnoheptose. The osazone forms yellow needles, m.p. 200°; by the action of hydrocyanic acid, rhamnooctonic acid is obtained. These compounds have the following optical activities :-

> Rhamnose, $[a]_D = +8^\circ$. Rhamnohexose, $[a]_D = -61\cdot1^\circ$. Rhamnohexose, $[a]_D = +8\cdot4^\circ$. Rhamnitol, $[a]_D = +10\cdot7^\circ$. Rhamnohexitol, $[a]_D = +11\cdot6^\circ$. Rhamnohexonolactone, $[a]_D = +83\cdot8^\circ$. Rhamnohexonolactone, $[a]_D = +55\cdot6^\circ$. Rhamnoctonolactone, $[a]_D = -51\cdot2^\circ$.

The reverse of the method of proceeding from one sugar to its next higher homologue—i.e. the production of a sugar containing one less carbon atom than the original one—has been effected by Wohl (B. 26, 730). When dextrose oxim (wide below) is treated with acetyl chloride it yields an acetyl compound from which hydrocyanic acid can be eliminated, and on hydrolysis of the resulting compound a pentose is obtained, which is d-arabinose; l-arabinose (ordinary arabinose) treated in the same way as dextrose yields a tetrose.

As is well known, mucic and saccharic acids are products of the oxidation of the sugars with nitric acid. The action of reducing agents on these bodies indicates their relationship to the sugars and helps to a clearer idea of the synthetical process.

Reduction of mucic acid. Sulphurio acid

Reduction of mucic acid. Sulphuric acid and zino-dust reduce mucic acid to i-galactonic acid (Fischer a. Hertz, B. 25, 1247). The lactone crystallises in delicate prisms, m.p. 122°–125°. It is optically inactive. By further reduction with sodium-amalgam i-galactose is obtained. This can be prepared as a crystalline mass, m.p. 140°–142°; it yields a hydrazide, m.p. 158°–160° and an osazone, m.p. 195°, which is identical with that obtained from the oxidation product of dulcitol (Fischer a. Tafel, B. 20, 3884).

i-Galactonic acid may be separated into the d- and l- acids by fractional crystallisation of the strychnine salts. The two acids yield respectively d- and l-galactose; d-galactose is identical with ordinary galactose. l-Galactose may also be obtained by the fermentation of i-galactose syrup. The sugar crystallises from alcohol, m.p. 162°-163°, its specific rotatory power is [a]_b = -78.6°, the phenyl-hydrazide

melts at 158°-160° and the osazone at 192°-195°. From this, as well as from other matters connected with the synthetical portion of the work, it will be seen that d-, i-, and l- were intended to mean dextro-, inactive, and lævo-, as indicating the optical activity of the substances to which they are prefixed; but it is obvious the d-body is not always dextro-, nor the l-body lævo-. Hence the unsatisfactoriness already indicated.

Reduction of saccharic acid. Fischer (B. 23, 930; 24, 521) found that the lactone of saccharic acid was easily reduced by sodium-amalgam with formation of glycuronic acid. Thierfelder (H. 15, 71) has shown that this latter is further reduced to an acid C₈H₁₀O₂, which reaction Fischer has confirmed, and calls the acid gulonic acid and the corresponding sugar gulose. The relationship of these bodies to the others of the sugar group is shown here:

Dextrose . OH.CH., (CH.OH), COH. Gluconic acid . OH.CH., (CH.OH), COOH. Saccharic acid . COOH. (CH.OH), COOH. Glycuronic acid . COOH. (CH.OH) . COH. . COOH.(CH.OH), CH2OH. Gulonic acid . COH.(CH.OH), CH2OH. Gulose

Glycuronic acid obtained by the reduction of d-saccharic acid has $[\alpha]_D = +19.1^\circ$, m.p. 175°-178°; d-gulonic acid obtained by the reduction of glycuronic acid yields a lactone, m.p. 180°-181°, $[\alpha]_D = +55^{\circ}$; its phenyl-hydrazide is easily soluble in hot water and hot alcohol, m.p. 147°-149°; d-gulose obtained by the reduction of dgulonic lactone is a colourless syrup, easily soluble in water and sparingly in absolute alcohol, it is oxidised by nitric acid to saccharic acid, it does not ferment with yeast.

Constitution. Dextrose and galactose are

represented by the formula OH_(OH).OH(OH).OH(OH).OH(OH).OH

and lævulose by

CH_(OH) CH(OH).CH(OH).CH(OH).CO.CH_OH

and for the following reasons.

By oxidation dextrose and galactose yield acids containing the same number of carbon atoms as themselves, whilst lævulose yields acids containing less carbon atoms than itself; dextrose and galactose yielding gluconic and galactonic acids CH₂(OH){CH(OH)}₄.COOH and by further oxidation saccharic and mucic COOH{CH(OH)},COOH respectively; acids whilst lawulose gives rise to trioxybutyric CH. (OH) CH(OH) COOH, formic H. COOH, and glycollic CH.OH. COOH acids.

By reduction dextrose and galactose yield respectively mannitol and dulcitol; lævulose also yields mannitol. These yield hexacetyl derivatives, and are reduced by hydriodic acid to secondary hexyl iodide CH₃.(CH₂)₂CHI.CH₄ (Wanklyn a. Erlenmeyer, J. 1861. 731; 1862. 480); they are evidently hexahydric alcohols of

normal hexane.

All three sugars combine with hydrocyanic acid to form nitriles which when boiled with hydrochloric acid yield different acids C,H14Os, which are reduced by hydriodic acid to heptoic acids; dextrose and galactose yielding normal heptoic acid CH₄(CH₂)₃.COOH and lævulose yielding methyl-butyl-acetic CH, CH.COOH (Kiliani, B. 18, 3066; 19, 221, 767 a. 1128). Galactose, dextrose, and lævulose | E

form oxims or iso-nitroso- compounds by combination with hydroxylamine (Rischbieth, B. 20, 2673; Jacobi, B. 24, 696; Wohl, B. 24, 998). With phenyl-hydrazine, dextrose and galactose yield hydrazides

CH₂(OH){CH(OH)}, CH.N₂H.C₂H₃; lævulose does not; but all three yield osazones CH₂(OH){CH(OH)}, C.N₂H.C₂H₃.CH.N₂H.C₃H₃. All three yield pentacetyl derivatives (Erwig a. Koenigs, B. 22, 2207). Previous to Fischer's synthetic work, other formulæ had been suggested (Tollens, Kurzes Handbuch d. Kohlen-hydrate; B. 16, 921; Sorokin, J. pr. [2] 37, 312), but there is no doubt now that the above are correct.

From the synthetic work above described, it is evident that the aldehyde formula belongs also to mannose, gulose, arabinose, xylose, and ribose, and also their synthetic homologues. Rhamnose, as is evident from its reactions, has also an aldehyde constitution; it appears to be methyl-arabinose.

Sorbose appears to be isomeric with lavulose. By oxidation it yields tri-oxyglutaric acid COOH. (CH(OH)), COOH and other acids, and by reduction it yields mannitol. It combines By reduction with with hydrocyanic acid. hydriodic acid and phosphorus it yields hexyl iodide (Kiliani a. Scheibler, B. 21, 3276).

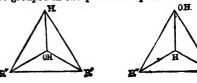
Little is known concerning the -on sugars. Maltose and lactose are probably anhydrides of equal molecules of dextrose and dextrose, and dextrose and galactose respectively; they yield acids (maltobionic and lactobionic) which on boiling with dilute mineral acids are decomposed into dextrose and gluconic acid and galactose and gluconic acid respectively. Both sugars thus appear to contain the dextrose-aldehyde group unaltered. Fischer (B. 21, 2633) has represented lactose as

CH (OH). CH(OH) CH OOH-{OH(OH)},OOH.

Cane sugar appears to be differently constituted from either maltose or lactose.

It thus appears that the formula CH₂(OH).{CH(OH)},COH must be assigned to eight known sugars—i.e. d- and l- glucose; d- and l- gulose, d- and l- mannose, d- and l-galactose (the i- modifications being analogous to racemic acid, i.s. compounds of the d- and lmodifications).

Now, as this formula contains four asymmetric carbon atoms, according to Van't Hoff's theory there will be possible 2'-16 isomers, which in the case of the symmetrical derivatives, as the hexahydric alcohols and dibasic acids, will be reduced to 10, of which $\frac{1}{2}2^4 = 8$ compounds are grouped in pairs (d. and l-) and $\frac{1}{2}2^{\frac{3}{2}}=2$ inactive, amesotomic (Fischer, B. 24, 1836) a. 2683; Van't Hoff, La Chimie dans l'Espace and Dix Années dans l'Histoire d'une Théorie). Each of the groups R"-C.H.OHbe grouped in two positions represented thus



(the carbon atom being considered to be at the centre of the tetrahedron). For the sake of brevity these two groupings will be written

The sixteen possible modifications of the glucose formula may then be expressed by the following symbols; in the case of the symmetrical derivatives (hexahydric alcohols and dibasic acids) Nos. 11-16 are identical with Nos. 5 to 10:—

Since saccharic acid may be obtained either from glucose or its stereoisomeride gulose, then the d- and l- saccharic acids must be represented by formulæ corresponding to two of those numbered δ to 10. Now, 7 and 8 may be eliminated, as they are optically inactive from internal compensation. Nos. 6 and 10 may be excluded for the following reasons: Glucose and mannose yield the same caszone, and consequently differ only in the arrangement of the groups round the asymmetric carbon atom next to the COH group. Other facts which confirm

these formulæ are (1) I-gluconic and I-mannonic acids are both produced by the nitrile reaction from arabinose; (2) fructose reduced by sodium-amalgam yields mannitol and sorbitol; (3) mannonic and gluconic acids can be converted, one into the other, by heating with quinoline; (4) all attempts to resolve gluconic and mannonic acids into two components have been unsuccessful.

If saccharic acid has the configuration 6 or 10, then manno-saccharic acid must have the configuration 7 or 8; but the latter are optically inactive, and therefore cannot be the configuration of manno-saccharic acid. Hence d-and l-saccharic acid must have a configuration corresponding to 5 and 9; for convenience we may assign 5 to the d- and 9 to the l- bodies.

Now the two sugars corresponding to d-saccharic acid—i.e. d-glucose and d-gulose—must consequently have the configuration 5 and 11; but in order to determine which to assign to one and which to the other, we must consider xylose and arabinose.

Arabinose by the nitrile reaction yields l-glucose and xylose yields l-gulose; now since l-glucose and l-gulose are

then xylose and arabinose must be

According to theory, there will be 2° = 8 isomeric pentoses, but when the formula becomes symmetrical, the asymmetry of the centre carbon is destroyed, and there will be only three isomeric pentahydric alcohols and three isomeric trioxy-glutaric acids, of which two are optically active, *.6.:

The same applies to the pentahydric alcohols. As arabitol and the tri-oxy-glutaric acid from arabinose are both laworotatory, whilst the corresponding compounds from xylose are optically inactive, the formula marked (a) is the formula for xylose, and (b) is the formula for arabinose, (9) and (15) being the formula for l-gulose and l-glucose respectively.

As lævulose yields the same osazone as man hose and glucose, d-fructose

The alcohols will be represented as follows: d-mannitol

and the dibasic acids: d-monosaccharic

The other di-basic acids known are mucic acid, which is optically inactive; allomucic acid, formed by the action of pyridine on mucic acid, which is also optically inactive; another isomeride of mucic acid, which is optically active, obtained by treating galactonic acid with quinoline and oxidising the product; and iso-saccharic acid. We have as yet not sufficient data to write the constitution of these with certainty.

So far we have considered the sugars in general; we must now turn to a description of the individual members of the group.

Class I. -Ose sugars.

Arabinose C.H.O. It does not occur in nature.

Formation.—By the action of dilute acids on metapectic acid, a constituent of beetroot, gum arabic, cherry-tree gum, gedda gum, gum tragacanth, and many other gums (Scheibler, B. 1, 58, 108; 6, 612; Claësson, B. 14, 1271; Martin, Sachsse's Phytochem. Unter. p. 69; Bauer, J. pr. [2] 34, 47; Kiliani, B. 19, 8030; v. Sandersleben, Sachsse, Lc., p. 90; O'Sullivan, C. J. 1884. 41, 1891, 1029; Stone a. Tollens, A. 249, 257; Stone, Am. 12, 435); wheat and rye bran (B. 23, 3110). Owing to the fact that most of the above gums yield galactose besides arabinose, the two were for some time confused (H. Kiliani, B. 13, 2304; 15, 34; Claësson, B. 14, 1270; B. C. 205; Scheibler, B. 17, 1729; E. O. v. Lippmann, B. 17, 2238). Arabinon yields on hydrolysis arabinose alone (O'Sullivan, C. J. 1890. 59).

Preparation.—A dextrorotatory gum, or one yielding little mucic acid when oxidised by nitric acid, is selected. A solution containing 30-40 g. per 100 c.c. is heated to 100° in a water bath and digested at that temperature with 2 p.c. sulphuric acid for 10-15 minutes. The solution is then cooled, neutralised with milk of lime, baryta water, or calcium or barium carbonate, and hot alcohol, S.G. 0.83, added as long as a precipitate is formed. The clear alcoholic solution is poured off the pp. which in a short time collects at the bottom, and evaporated in a vacuum to a syrup. After standing a few hours arabinose crystallises out in well-formed rhombic prisms with monoclinic terminations, and is purified by recrystallisation from water.

Formula and synthesis. See general part. Properties .- Arabinose crystallises in fine rhombic prisms out of aqueous solution. The termination and habitat of the crystals vary with the source. From alcoholic solutions it separates in sphenoids. The crystals melt at 160°. Its specific rotatory power is $[a]_{\rm p}$ + 104.5° for 10 p.c. solution at 20°C.; in more concentrated solutions it is higher, $[\alpha]_p = +110^\circ$ being observed for nearly saturated solutions. Freshly-prepared solutions possess bi-rotation, an angle of $[a]_p = 156.7^\circ$ having been observed (Scheibler, l.c.; O'Sullivan, l.c.; E. O. v. Lippmann, l.c.; Bauer, L. V. 36, 304; Tollens, A. 257, 160; Groth, B. 6, 615; Griess a. Harrow. B. 20, 3111). The specific gravity of aqueous solutions does not increase regularly with the strength, being less in proportion for high con-centrations than for low ones. The S.G. of a solution containing 10 g. in 100 c.c. at 15.5° is 1.0385-1.0384.

Action of acids. The action of boiling dilute mineral acids destroys much of the opticity and K. of arabinose in a short time (O'Sullivan, 1884, 55). The first products of the change have not been studied. When, however, the action is continued for a considerable time, no lævulinic acid is produced, but furfural is present in large quantities (Ganz, Stone, a. Tollens, B. 21, 2148; 23, 3791). Chalmont a. Tollens (B. 24, 694) obtained 52.7 p.c. furfural by precipitating the distillate from the products of action of acids with phenylhydrazine acetate. Formic and aceto-propionic acids and humus substances are also present (Conrad a. Guthzeit, B. 18, 2905).

Reduction.—Treated with sodium-amalgam

Reduction.—Treated with sodium-amalgam in acid solutions, arabinose yields arabitol, a body resembling sorbitol, m.p. 102°. This body

does not reduce Fehling's solution, and is optically inactive (Kiliani, B. 20, 1233)

Oxidation.-By oxidation with nitric acid arabinose yields arabonic acid (lævorotatory trioxyglutaric acid), and by violent oxidation at an elevated temperature oxalic acid (Kiliani, B.21, 3006). Bromine also oxidises it to arabonic acid C₅H₁₀O₆, m.p. 89° (Bauer, J. pr. [2] 34, 47; C. C. 1877, 732; Kiliani, B. 19, 3031; 20, 345). It is oxidised by alkaline solutions of metallic oxides, 100 c.c. of Fehling's solution being reduced by 0.4303 g. arabinose, and 100 c.c. of Sachsse's solution being reduced by 0.4375 g. arabinose (Bauer, L. V. 36, 304). Its K=108 110 (O'S., l.c.), i.e. 100 pts. of it reduce as much CuO as 108-110 pts. dextrose.

Fermentation.—Arabinose does not seem to be capable of fermentation with yeast (Stone a. Tollens, L.c.; Scheibler, L.c.; v. Lippmann, B. 17, 2238); with Bacillus ethaceticus it ferments with the production of ethyl alcohol, acetic, formic, and carbonic acids, and hydrogen (Frank-

land a. MacGregor, C. J. 1892. 737).

Compounds with cyanogen. By the nitrile reaction arabinose yields a mixture of l-gluconic and l-mannonic acids (Kiliani, B. 19, 3029; 20, 282, 339, 2710; Fischer, B. 23, 2611; 24, 539). The first product of the action of HCy on arabinose seems to be a compound of both bodies. HCl converts this into a crystalline amide C.H.,NO, which on treatment with baryta yields the acids just mentioned.

Hydrazine compounds. Arabinose yields with phenylhydrazine, just as dextrose (q.v.) does, an osazone, m.p. 157°-158° (Scheibler, B. 17, 1729; Kiliani, B. 20, 345).

Compounds with diamines. With o-diamidobenzene, arabinose yields arabinose-o-diamidobenzene C₆H₄.(NH)₂C₅H₈O₄. It is a neutral body, not reducing Fehling's solution, and is dextrorotatory, m.p. 235° with decomposition; it withstands boiling with dilute hydrochloric acid or potash, and yields a hydrochloride and hydrobromide. Arabinose-m-pdiamido - toluene C₆H₂(CH₂)(NH)₂:C₅H₆O₄, m.p. 288°, and arabino - γ - diamido - benzoic acid COOH.C.N.:(NH).:C.H.O. + 2H.O. m.p. 235°, have also been obtained (Griess a. Harrow, B. 20, 3111). Arabinose gives colour reactions with orcinol and phloroglucinol (Wheeler a. Tollens, A. 254, 314; 260, 304). Ribose. Fischer a. Piloty, B. 24, 4214.

When certain carboxylic acids of the sugar group are heated with quinoline or pyridine, they are converted with stereo-isomeric acids.

When arabonic acid OH.CH₂(CH.OH)₃.COOH is treated in this manner it yields a new acid COOH. (CH.OH) 3. CH OH, which the authors name ribonic acid; ribonic lactone crystallises from solution in ethyl acetate in long prisms, m.p. 72°-76°, does not reduce Fehling's solution, and is easily soluble in water, alcohol, and acetone. Its specific rotatory power is $[\alpha]_{D20} = -18$. Its phenylhydrazide forms colourless needles, m.p. 162°-164°. Ribonic acid may be converted into arabonic acid by heating with quinoline.

By reducing ribonic lactone with sodiumamalgam, it yields the new sugar ribose, which forms a phenylhydrazide crystallising from alcohol in colourless crystals, m.p. 154°-155°; ribose-osasone is identical with arabinoseosazone. By oxidation ribonic acid yields trioxy-glutaric acid, which differs from the tri-oxyglutaric acid obtained by the oxidation of xylose, but which is also optically inactive. By the further reduction of ribonic lactone, a penta-hydric alcohol was obtained which has been found to be identical with the naturally occurring adonitol, obtained from Adonis vernalis (Fischer, B. 26, 636).

It does not occur free in Xylose. $C_5H_{10}O_5$. nature.

Formation.—It is obtained by the action of dilute boiling acids on xylan or wood gum (Koch, B. 20, ref. 145; Wheeler a. Tollens, B. 22, 1046; A. 260, 289; Winterstein, H. 17, 381; Stone a. Test, Am. 15, 195), on the carbohydrate from the epidermis of Psyllium gallicum (Bauer, A. 248, 140), on vegetable amyloid (Winterstein, Z. P. C. 17, 353), on brewer's grain (Stone a. Tollens, A. 249, 227; 271, 55), on jute (Wheeler a. Tollens, l.c.), and other vegetable bodies (Voswinkel, C. C. 1891, 2, 655; Hebert, Ann. Ag. 16, 358; 18, 261; Stone a. Lotz, B. 24, 1657; Allen a. Tollens, B. 23, 137; Bertrand, Bl. [3] 5,

Preparation.—Xylose may be prepared by extracting straw or any other of the above-mentioned materials by 5 p.c. soda, precipitating the wood gum from the solution by alcohol, and hydrolysing this by boiling with dilute sulphuric acid. The digested solution is treated in the same way as described in the case of arabinose, and the xylose may be easily purified by

crystallisation from water.

The formula and synthesis have already been

discussed in the general part.

Properties. - Xylose is easily soluble in water; 100 pts. of water at 20.3° dissolve 117 pts. of xylose; it is insoluble in absolute alcohol, 90 p.c. alcohol will dissolve in 10 c.c. 0.4 g, sugar at 19° (Bertrand, Bl. [3] 7, 499). Its solutions are optically active, freshly-prepared solutions having about four times the constant optical activity (Parcus a. Tollens, A. 207, 160). For a 10 p.c. solution at 20° [a]_D = +19·3° (Parous a. Tollens, l.c.; Bertrand, l.c.). Schulze a. Tollens (A. 271, 40) give the formula [a]_D = +18·095 + 0·06986p at 15° -20°, where p = percentage of xylose in solution. Xylose frequently crystallises in prisms belonging to the ortho-rhombic system. Its solutions are not fermentable by yeast. It reduces Fehling's solution, K = 109.6, agreeing in this respect very closely with arabinose (Bertrand, l.c.). Boiled with acids it yields about 50 p.c. furfural (Günther a. Tollens, B. 23, 1751; Chalmont a. Tollens, B. 24, 694; Bertrand, l.c.).

It is reduced by sodium-amalgam to a pentahydric alcohol xylitol (Bertrand, Bl. [3] 5, 554; Fischer a. Stahel, B. 24, 528). It is oxidised by bromine to xylonic acid (Bertrand, l.c.; Allen a. Tollens, A. 260, 806); it is oxidised by nitric acid to inactive tri-oxy-glutaric acid C, H,O, (Fischer, B. 24, 1836). By the nitrile reaction xylose yields *l*-gulonic acid, the stereo-isomeride of *l*-gluconic acid (Fischer a. Stahel, B. 24, 528; Fischer, B. 23, 2625; Fischer a. Curtis, B. 25, 1025).

With phenyl hydrazine it yields an osazone, m.p. 160°. Warmed with phloroglucinol and hydrochloric acid, it gives a cherry-red coloura-

tion, just as arabinose does under the same conditions (Tollens a. Mayer, B. 21, 3508; Wheeler a. Tollens, B. 22, 1046).

Rhamnose. Isodulcite, rhamnodulcite

C₆H₁₂O₅.H₂O.

Preparation.—Rhamnose is obtained, together with other bodies, by the action of hot dilute acids on quercitrin, hesperidin, xanthorhamnin, naringin, and frangulin (Rayman a. Kruis, Bl. [2] 48, 632; C. C. 1888, 6; Hlasiwetz a. Pfaundler, A. 127, 362; Liebermann a. Hörmann, A. 196, 323; Kruis, A. 196, 333; Berend, A. 196, 328; Rigaund, A. 90, 292; Will, B. 18, 1316; 20, 297 and 1186; Dehn, Zeit. d. Ver. 15, 562; Rayman, Bl. [2] 47, 668; Tanret, Bl. 49, 20; Thorpe a. Miller, C. J. 1892. 8).

Properties.—Rhamnose forms fine crystals, which have a sweet taste, and a composition corresponding to the formula C₆H₁₂O₅.H₂O; by careful drying it loses water and then melts at 90°-110° (Websky, B. 18, 1318; Hirschwald, A. 196, 330). It is easily soluble water, and its solutions are dextrorotatory, $[a]_{D} = +9^{\circ}$. Freshlyprepared solutions show bi-rotation (Schnelle a. Tollens, A. 271, 61). In alcoholic solutions the optical activity decreases as the proportion of alcohol to water increases, until the solution becomes lavorotatory (Rayman a. Kruis, C. C. 1888. 6). It reduces Fehling's solution, 1 c.c. being reduced by 0.0052-0.0055 g. rhamnose (K = 91-96) (Rayman a. Kruis, l.c.). Its solution does not ferment with yeast.

Action of acids. Boiled with moderately strong acids furfural is evolved (Maquenne, C. R. 109, 603). Concentrated nitric acid converts it into a nitrate.

Oxidation .- Nitric acid oxidises rhamnose to tri-oxy-glutaric acid (Will a. Peters, B. 22, 1697). Malin (A. 145, 197) obtained an acid C_eH₁₀O₉. Bromine water oxidises it to rhamnonic acid C₆H₁₂O₆, which is obtained in the form of the lactone C₆H₁₀O₅, m.p. 148° (Will a. Peters, B. 21, 1813; Rayman, B. 21, 2046). With bromine and silver oxide it is oxidised to acetaldehyde and acetic acid (Herrig, Chem. Zeit. 1887, Rep. 145; M. 8, 227).

Reduction. - Sodium-amalgam reduces rhamnose to rhamnitol CH4(CH.OH)4.CH2OH, which crystallises from alcohol or acetone in triclinic prisms, m.p. 121°, and has a sweet taste (Fischer

a. Piloty, B. 23, 3102).

Compounds .- With phenylhydrazine rhamnose yields a hydrazide C₆H₁₂O₄:N₂HC₆H₅, m.p. 159°, and an osazone C₁₄H₂₂N₄O₃, m.p. 180° (Will, B. 20, 1186; Fischer a. Tatel, B. 20, 1089, 1091 a. With aniline it forms a compound C₈H₁₂O₄NC₈H₂, m.p. 118° (Rayman a. Kruis, l.c.). With ammonia and ethylaceto-acetate it forms rhamnodiasine C18H22O8N2, m.p. 186° (Rayman a. Chodoünský, B. 22, 304 and 3247).

Rhamnose combines with hydrocyanic acid to produce rhamnonamide. The lactone C, H12O, crystallises in needles, melting-point 168°; this is reduced by hydriodic acid and phosphorus to normal heptoic acid (Fischer a. Tatel, B. 21, 1657 a. 2173). Rhamnose hexoic acid lactone is reduced by sodium-amalgam to a sugar (Fischer, B. 22, 2204; Fischer a. Tafel, B. 23, 8102). Sodium rhamnose C.H., O.Na, is pre-cipitated by adding sodium ethylate to an alcoholic solution of rhamnose (Liebermann a.

Hamburger, B. 12, 1186). With benzoyl chloride tri- and tetra- rhamno-benzoates are formed; with acetic anhydride rhamno-tetracetate (Rayman, B. [2] 47, 668). From thermo-chemical considerations rhamnose appears to be methylxylose (Stohmann a. Langbein, J. pr. [2] 45, 805).

Fucose C₆H₁₂O₅ is obtained by the hydrolysis of fucus. The sugar forms microscopic needles. Its solution has a sweet taste and is levorotatory, $[\alpha]_D = -77^\circ$. With phenyl-hydrazine it forms a hydrazide, melting-point 170°, and an osazone, melting-point 159° (Bieler a. Tollens, A. 258, 110; Günther a. Tollens, B. 23, 2585). Fucose appears to be methyl-arabinose (Stohmann a. Langbein, l.c.).

Dextrose. d-glucose, glucose, starch-, corn-, grape-, honey-, diabetic-sugar. (Ger. Krummelsucker, Traubenzucker; Fr. Sucre de raisin.)

 $C_{4}H_{12}O_{4}$ and $C_{6}H_{12}O_{4} + H_{2}O_{5}$

Occurrence.—Dextrose is widely distributed throughout the vegetable kingdom, being usually accompanied by lævulose and cane sugar. The most important sources are honey, most fruits, the sap of plants, seeds. It occurs in the animal kingdom in the liver, blood, and other parts of the body, and in diabetic urine (S. Sachsse, Die Ch. und Phys. der Farbst. Kohlenhydrate u. Protein subst., Leipzig, 1877; F. Mack, J. 1869, 892; Girard, B. 17, 17c; Neubauer, Fr. 12, 39; Gorup-Besanez, B. 4, 906; Petit, B. 6, 1313; v. Schneider, A. 162, 235; Report on Sorghum and Cornstalks, Dep. of Agr., Report No. 35, Washington, 1881; Hermann a. Tollens, A. 230, 50; Vogel, B. 15, 2271; Fischer a. Bödecke, A. 117, 111; De Bary, Hoppe-Seyler's Medic. Ch. Unt. p. 72; Hoppe-Seyler, Handb. d. phys. Ch. Ana., 1883, 301; Klason, B. 19, 2541; Wehmer, Inaug. Diss., Göttingen, 1886, 40; O'Sullivan, C. J. 1886, 58; Wehmer a. Tollens, B. 19, 707; Seegen a. Kratzschmer, Pf. 22, 206; Kütz, Pf. 24, 52; Boussingault, C. R. 91, 639; J. Sieben, B. C. 1885. 134; Hermann a. Tollens, A. 230, 50; Müller, Ann. Agronom. 13, 88; Seegen, Pf. 37, 348; 89, 121; 40, 48; Cuisinier, C. C. 1886, 614; H. Thoms, B. 21, 1916a; Reali, G. 17, 325; Maquenne, A. Ch. [6] 17, 495).

Formation .- By the action of acids and ferments on the following glucosides: amygdalin (Schmidt, A. 119, 92; Hesse, l.c., 176, 112); populin (v. Lippmann, B. 12, 1648b); ruberythrio acid (Liebermann a. Bergami, B. 20, 2247); salicin (Schmidt, A. 119, 97; Hesse, Lc., 176, 112); lupinin (B. 11, 2200 b); hesperidin (Tanret, Bl. 49, 20); phlorizin (Rennie, C. J. 1887, 636). Other glucosides yield dextrorotatory, reducing, fermentable, and crystallisable sugars, but there is not sufficient evidence to prove with certainty that they are dextrose. These are: sesculin (Rochleder a. Schwarz, A. 87, 186; Zwenger, l.c., 90, 76); arbutin (Hlasiwetz a. Grabowski, Ar. Ph. 141, 329); coniferin (Tiemann a. Harmann, B. 7, 608).

By the action of dilute acids the following carbohydrates yield dextrose, alone or with other glucoses. Some of them are also hydrolysed to dextrose by ferments:

a. and B. amylan yield dextrose.

Cellulose yields dextrose (Flechsig, H. 7, 528; Lindsay a. Tollens, A. 267, 370).

Glycogen yields dextrose.

Lichenin yields dextrose.
Tunicin
Starch

Dextrin ,, Maltose ,,

Lactose yields dextrose and galactose.
Cane sugar yields dextrose and lævulose.
Raffinose yields dextrose, lævulose, and

Preparation.—Dextrose can be prepared from any of the substances above mentioned, but it is in many cases difficult to obtain it in any quantity in the pure state. Starch and sucrose are the bodies usually employed, and we find that lactose can also be used with advantage.

(a) A white saccharum (the commercial glucose obtained by the action of acids on starch) with an optical activity approaching that of dextrose is selected. This is scraped as fine as possible, and treated with boiling ethyl alcohol S.G. 820. On cooling, the clear saturated solution is separated from the undissolved syrup and a little previously prepared dextrose added to it. Crystallisation soon begins and continues for some time; a crop of fairly pure dextrose is thus obtained. Further purification can be effected by melting the crystals in 8-10 p.c. boiling water and adding dry boiling methyl alcohol until a slight cloud is produced. Anhydrous dextrose in a state of great purity separates abundantly from this solution. Or the saccharum is melted in 5 or 6 p.c. water, allowed to crystallise, the crystals pressed and further purified by treatment with methyl alcohol as described.

(b) A 25 p.c. aqueous solution of crystallised sucrose is taken and heated to 50°-55° and 2-8 p.c. pressed brewers' yeast stirred in (Tompson, E. P. 8686, 1884), the temperature is maintained for 2-3 hours, when the sucrose is completely inverted—that is, converted into dextrose and lævulose. The solution is then evaporated in a vacuum to a syrup, S.G. 1·5 or thereabouts. On standing for a short time this solidifies to a crystalline mass; it is washed with cold alcohol, S.G. 830, and the residue dissolved to saturation in alcohol of the same strength. Dextrose soon begins to crystallise out; the crystals can be further purified by dissolving in

methyl alcohol and recrystallisation.

Sucrose may also be inverted by digestion with sulphuric acid, the acid separated by baryta water, and the filtrate from the barium sulphate evaporated to a syrup and treated in the way mentioned. Inversion by yeast yields the best result. Soxhlet (J. pr. [2] 21, 242) recommends that inversion be effected by alcoholic hydrochloric acid. 480 c.c. of strong hydrochloric acid are mixed with 12 litres of alcohol, S.G. 810, in a glass vessel, the mixture heated 45°-50°, 4 kilos. powdered sucrose stirred in, and the whole maintained at that temperature for two hours. When cool a little previously prepared anhydrous dextrose is added, and in a few days there is an abundant crystallisation of dextrose. This can be purified by crystallisation from methyl alcohol. Should any colour be developed, the aqueous solution is treated with animal charcoal and again concentrated in a vacuum. Under such circumstances it is better to begin again, for if proper care be taken no colour need be produced.

An aqueous solution containing 25 p.c.

lactose and 8 p.c. sulphuric acid is digested in a water-bath for three hours, care being taken by continued agitation that no colour is developed, the sulphuric acid is separated by baryta water, and the filtrate concentrated to a syrup containing 76-80 p.c. solid matter. On standing this solidifies, and after a few days is treated with boiling methyl alcohol; this dissolves out the dextrose, leaving the galactose undissolved; the former soon crystallises from the saturated methyl alcohol. A recrystallisation yields the pure body in moderately large, well-defined crystals.

Hydrous dextrose $C_sH_{12}O_s$: H_2O can be obtained from any of the above preparations by dissolving in 12 to 20 p.c. water, and allowing to crystallise. If honey is used as a source of dextrose, any sucrose it may contain should be inverted by yeast or acids, and the residue obtained by evaporation crystallised from methyl alcohol as in the case of inverted sucrose. A pure product may be obtained in a few crystal-

lisations.

Formula and synthesis. See first part on the

sugars generally.

Properties.—Dextrose crystallises out of 93-95 p.c. ethyl or methyl alcohol in the anhydrous state. Anhydrous crystals may also be obtained by special treatment from concentrated aqueous solutions at 30°-35°. As a rule aqueous solutions deposit crystals containing one molecule of water of crystallisation. Anhydrous dextrose crystallises in rhombic, hemibedric forms; the axial relationship is a:b:c = 0.704:1: 335. Hydrated dextrose forms hemimorphous crystals; the elements are a:b:c = 1.7850:1:1.9080, $\beta = 97^{\circ} 59'$ (F. Becke, M. 10, 231). Well-defined crystals of both varieties are difficult to obtain; usually the first separates as a powder and the second in cauliflower-like masses; both dissolve easily in water, the first with evolution of heat. Hydrated dextrose dissolves easily in strong ethyl and methyl alcohol, but the anhydrous sugar is almost insoluble in them. It is insoluble in ether and hydrocarbons, slightly soluble in aniline, acetic acid, and acetic ether. Hydrated dextrose easily loses its molecule of water of crystallisation when exposed in a vacuum over sulphuric acid, or when heated gently. The anhydrous sugar melts at 144°-146° (Hesse, A. Ch. 176, 106); its specific gravity is, according to Guerin-Varry a. Heintz, 1.386; according to Bödecker, 1.5384 (v. Lippmann, Zuckerarten, p. 12). Dextrose is not so sweet as cane sugar. Behr (B. 15, 1106) finds it ? as sweet, and T. Schmidt (B. C. 1887, 405) that 1.53 pts. of dextrose are as sweet as 1 part of cane sugar. An aqueous solution of 10 g. dry dextrose in 100 c.c. of solution at 17.5° has a specific gravity of 1.0381 (water at 17.5 = 1) (Salomon, B. 14, 2710). This is fairly accurate. K = 100. Its solutions rotate the plane of polarisation of a ray of polarised light to the right. According to Hesse a. Tollens (B. 17, 2284), the amount of rotation for the D line is expressed by the formula $[a]_D = 52.5 + .018796P + .00051683P^2$ for anhydrous dextrose, and $[a]_D = 47.78 + .015534P + .00038888P^2$ for hydrated

[a]_D = 47.73 + 010534P + 0003888P for hydrated dextrose, in which P is the percentage of the two varieties respectively in solution. In solutions containing 10 p.c. dry sugar and thereabouts, the

factors $[a]_D = 52.7$ and $[a]_i = 58.3$ are sufficiently accurate, but probably a shade too low. These values are not materially affected by the temperature. The presence of some inactive bodies in the solution appears to have some effect (Pribram, M. 9, 395); bodies such as potash, soda, lime, &c., which act on dextrose reduce its optical activity; other bodies, such as urea, do not alter it (N. Wender, B.24, 2200). Freshlyprepared solutions have a greater optical activity than those which have stood a few hours. F. Urech (B. 17, 1547) has observed a rotation 2.19 times greater than the constant value. Parcus a. Tollens (A. 257, 160) observed an angle of $[a]_D = 105.16$ in a 10 p.c. solution $5\frac{1}{3}$ min. after commencing solution. This phenomenon is called bi-rotation; the optical activity quickly falls, and the constant value is reached in about six hours in the cold and immediately on boiling. bi-rotation is also immediately destroyed by potash (O'Sullivan a. Tompson, C. J. 1890, 920), or ammonia (Schulze a. Tollens, A. 271, 49). Dextrose when produced by the inversion of cane sugar at a low temperature is set free in the birotatory state (O'Sullivan a. Tompson, l.c.). Only crystallised dextrose exhibits this phenomenon; fused dextrose does not show it (Hesse, A. 176, 113). A solution of dextrose in strong alcohol retains its bi-rotation (?) The dispersive power of dextrose solution is practically the same as that of quartz (Hoppe-Seyler, Fr. 3, 1866, 412). The ratio between the specific rotatory power for the C and D lines respectively is 1:1-258 (Grimbert, J. Ph. [5] 16, 295 and 345). The heat of combustion of dextrose is +677.2 cal., and the heat of formation +300.8 cal. (Berthelot a. Recoura, C. R. 104, 1571).

Action of heat. When the dry sugar is heated to 170°, a molecule of water is eliminated and dextrosan C.H.O. is produced, which may be reconverted into dextrose by the action of water and dilute acids. If the temperature is raised to 200°, gas, water, and volatile acids are given off, and if the heating be continued without raising the temperature, a brownish black mass is finally left. This substance is soluble in water and constitutes the colouring matter (caramel) used in beer, wine, cooking, &c. Should the temperature be raised, carbon monoxide, dioxide, and methane are evolved, and aldehyde, furfural, acetone, metacetone, propionic, acetic, and formic acids distil over, and a black carbonaceous mass is left. If the hydrate be heated for a considerable time in a closed tube a fluid is obtained which absorbs oxygen and nitrogen (?) from the air (Thénard, C. R. 52, 795).

Action of acids. Dextrose is dissolved by strong sulphuric acid without blackening, dextrose-sulphuric acid being formed (Péligot, A. Ch. [2] 67, 170); from this solution alcohol precipitates a compound of diglucose with alcohol (Musculus a. Meyer, B. 14, 850). Similar compounds are produced by the action of chlorosulphonic acid (Claësson, J. pr. [2] 20, 1). A mixture of sulphuric and nitric acids converts dextrose into a nitro-compound (Carey Lea, Bl. [2] 10, 415). Dextrose phosphoric acid is said to exist (Amato, B. 4, 413). Several acetyl derivatives have been described. Dextrose discetate C₆H₁₆O₄(C₂H₁O₄)₂ and dextrose triacetate

 $C_eH_sO_s(C_sH_sO_s)_s$ are amorphous, soluble in water, alcohol and ether (Schützenberger a. Naudin, Bl. [2] 12, 107, 204; Liebermann a. Hörmann, B. 11, 1619; Herzfeld, B. 13, 265). Dextrose pentacetate C₂H₂O₂(C₂H₂O₂), is prepared by the action of acetic anhydride and zinc chloride on dextrose, m.p. 111-112 (Erwig a. Koenigs, B. 22, 1464, 2207). Didextrose octoacetate $C_{12}H_{14}O_{2}(C_{2}H_{2}O_{2})_{8}$ is prepared by the action of acetic anhydride and sodium acetate on dextrose, m.p. 184 (Herzfeld a. Erwig a. Koenigs, l.c.). Dextrose-monochlorhydrintetracetate is formed by the action of acetyl chloride on dextrose (Culley, C. R. 70, 401). Tri-, tetra-, and penta-benzyl (C_eH₁(C_rH₅O)₅O₆) dextroses are obtained by the action of benzoic chloride and potash on dextrose (Skraup, M. 10, 389; Kueny, H. 14, 330; Panormoff, J. R. 23, 375). Compounds of dextrose with other organic acids have been prepared (Berthelot, A. Ch. [3] 54, 74; Gm.-K. 7, 764; v. Lippmann, Zuckerarten, p. 40; Baumann, B. 19, 3218; Colley, C. R. 76, 436). If dextrose is boiled with dilute sulphuric acid the specific gravity of the solution gradually increases for some time, apparently from the fixation of the elements of water. On continued boiling the solutions become coloured, brown humus substances, acetopropionic and formic acids, are produced (Malaguti, A. Ch. [2] 59, 407; Tollens a. Grote, A. 175, 181; 206, 207; Wohl, B. 23, 2084). Dilute hydrochloric acid produces similar results, but more rapidly. Gummy bodies are also produced (Grimaux a. Lefèvre, C. R. 103, 146). See also-Conrad a. Guthzeit (B. 18, 439; 19, 2569) and Wehmer a. Tollens (B. 19, 707).

Action of alkalis. Alkalis give a brownish colour with dextrose solution, quickly on heating, slower in the cold, with production of acetal, acetone, and lactic, acetic, and formic acids. Other bodies of unknown composition are also formed, such as glucinic, saccharumic and japonic acids. When potash is used oxygen is absorbed. With calcium hydroxide, saccharin or calcium saccharinate is also produced; this is not one of the products of the action of potash (Emmerling a. Loge, B. 14, 1005; 16, 838; Henry, B. 14, 2272; Breuer a. Zincke, B. 13, 638; Rochleder a. Kawalier, J. pr. 94, 403; Hoppe Seyler, B. 4, 346; Kiliani, B. 15, 700; Nencki a. Sieber, J. pr. [2] 24, 498; Péligot, A. Ch. 30, 75; Mulder, A. Ch. 36, 260; Reichardt, Z. 1870. 404; v. Lippmann, Zuckerarten, p. 22; Péligot, B. 13, 196; Scheibler, B. 13, 212; Kiliani, B. 15, 701; Cuisinier, v. Lippmann, Zuckerarten, p. 25; Scheibler, B. 16, 2434). Thenard (C. R. 52, 444) obtained bitter bodies containing 10-19 p.c. nitrogen, by acting on dextrose with ammonia; they absorboxygen from the air, and resemble certain nitrogenous humic acids. Tanret (Bl. [2] 44, 102) obtained two volatile bases: a-glucosine $C_aH_aN_2$ b.p. 186° and β -glucosine $C_sH_{10}O_2$ b.p. 160°, by heating dextrose with strong ammonia. Heated with aniline, dextrose anilide is produced (Schiff, A. 140, 123; Sorokin, B. 19, 298).

Reduction.—Nascent hydrogen reduces dextrose in alkaline solutions to mannitol, but not in neutral or acid solutions (Linnemann, A. 128, 136; Dewar, Z. 1870. 418: Bouchardat, Bl. [2] 16,88; Krusemann, B. 9, 1465). Under the most

favourable conditions the yield does not exceed 10 p.c. If the reduction be carried on at a high temperature other fatty alcohols are formed, Meunier (C. R. 111, 49) obtained sorbitol by reducing dextrose with sodium-amalgam.

Oxidation .- Gluconic acid C. H12O, is produced by oxidising dextrose with mercuric oxide (Heffter, B. 22, 1049; Herzfeld, A. 245, 27) or with chlorine or bromine and subsequent treatment with silver oxide (Hlasiwetz a. Haber-mann, A. 155, 122; 156, 253; Kiliani, A. 205, 182; Kiliani a. Kleeman, B. 17, 1298; Herzfeld, A. 220, 335, 358), or by the action of Bacterium aceti (Brown, C. J. 1886. 179; Boutroux, C. R. 1880. 236). Oxygen has no action on dextrose in neutral or acid solutions, but it is absorbed by alkaline solutions. Ozone oxidises dextrose to saccharic and formic acids (Renard, A. C. 5, 17, 289). Platinum black acts similarly (Loew, B. 23, 865). Nitric acid oxidises dextrose to saccharic acid C.H.O., and finally to oxalic, formic, and carbonic acids (Liebig, A. 113, 1; Heintz, P. 51, 183; Hornemann, J. pr. 89, 304; Kiliani, A. 205, 172; Guérin-Varry, A. 8, 31; Sohst a. Tollens, Z. 11, 99). Iodine when acting on alkaline solutions of dextrose yields a little iodoform (Lieben, A. Suppl. 7, 228; Hermann a. Tollens, B. 18, 1335; Millon, C. R. 21, 828). Alkaline solutions of many metallic oxides oxidise dextrose, in some cases being reduced to the metal, in others to lower oxides. Potassium ferrocyanide, potassium permanganate, indigo, picric acid, litmus, &c., oxidise dextrose also. The chief products are formic, oxalic, glycollic, and carbonic acids. Some of these reactions are employed for the quantitative estimation of dextrose (Tollens, B. 16, 921; Herzfeld, Zeitschr. d. Ver. d. Rubenzucker-Ind. 87, 837; Habermann a. Hönig, B. 15, 2624; Worm-Müller a. Hagen, Pf. 22, 325; B. 13, 2096 a. 2442; Herzfeld a. Bruhns, Zeitschr. d. Ver. d. Rübenzucker-Ind. 86, 110; Reichardt, A. 127, 297; Felsko, A. 149, 856; Beyer, A. 131, 353; Claus, A. 147, 115; Smolka, B. 20, 167; M. 8, 1).

Fermentation.—Under the influence of various species of saccharomyces and other organisms, dextrose ferments, yielding alcohol and carbon dioxide as the chief products, glycerol, succinic acid, and small quantities of other bodies are formed at the same time. Bacteria decompose dextrose in other ways, one yielding chiefly lactic acid, another butyric, another cel-

lulose, &c. (v. FERMENTATION).

Compounds with acids (v. Action of acids).
Compounds with bases. Potassium and sodium dextrose C₈H₁₁KO₆ and C₈H₁₁NaO₆ are prepared by adding potassium or sodium ethylate to a solution of dextrose in strong alcohol. A white, bulky precipitate is obtained, which is very hygroscopic, and decomposed by water (Hönig a. Rosenfeld, B. 10, 871; Franchimont, B. 10, 994). The compounds with the alkaline earths are differently constituted, being apparently a combination of the sugar and base. They are formed by mixing alcoholic solutions of the base and sugar, or by precipitating their mixed aqueous solutions with alcohol. Lime forms C₈H₁₂O₆CaO.H₂O, C₈H₁₂O₆CaO.2H₂O, C₈H₁₂O₆SaCaO.2H₂O, 4C₈H₁₂O₆SaCaO.2H₂O, and similar compounds are

formed with baryta, strontia, and magnesia. Neither lead acotate nor subacetate precipitate aqueous dextrose solutions, but lead oxide yields compounds similar to those with lime &c. (Brendeche, Ar. Ph. [2] 29, 84; v. Lippmann, Zuckerarten, p. 45; Péligot, J. pr. 15, 105; Franchimont, B. 12, 1939). Alkaline dextrose solutions dissolve copper hydroxide, and from these solutions soda or potash in proper proportions precipitates compounds of copper oxide and dextrose of varying composition (Guignet, C. R. 109, 528; Worm-Müller a. Hagen, Pf. 22, 325; Fileti, B. 8, 441; Salkowski, B. 12, 704). The compound C₄H₁₂O_g.2ZnO.3H₁O is formed by adding an alcoholic solution of dextrose to an ammoniacal solution of zine oxide (A. C. Chapman, C. J. 1889. 576). Dextrose forms amorphous compounds with nickel, chromium, and iron oxides (Chapman, C. J. 1891. 323).

Compounds with metallic salts. Dextrose forms well-defined crystalline compounds with sodium chloride and bromide; their formulæ are 2C,H₁₂O_e,NaCl.H₂O, which crystallises from diabetic urine, C₆H₁₂O_e,NaCl, C₆H₁₂O_e,2NaCl, 2C₆H₁₂O_e,NaBr (Pasteur, A. Ch. [3] 31, 92; Erdmann a. Lehmann, J. pr. 13, 111; v. Lippmann, Zuckerarten, p. 45; Städeler, C. C. 1854. 930; Hönig a. Rosenfeld, B. 10, 872; Stenhouse,

A. 129, 286).

Compounds with cyanogen. By digesting an aqueous solution of dextrose with hydrocyanic acid solution, the following reaction takes place: $C_0H_{12}O_0+HCN+2H_2O=C_1H_{12}O_0NH_1$ (Kiliani, B. 19, 767). To prepare this compound Kiliani gives the following directions: 100 g. crystalline dextrose are dissolved in 30 g. water, and an equivalent quantity of 60 p.c. hydrocyanic acid is added and digested for six days at the ordinary temperature; after twenty-four hours' further digestion at 35°C. all smell of hydrocyanic acid had disappeared, and the liquid smelt of ammonia. The ammonium salt thus obtained is converted into a barium one by boiling with baryta, and the free acid obtained by removal of the barium by sulphuric acid. Fischer (A. 270, 64) dissolves 5 kilos. dextrose in 25 litres 3 p.c. hydrogen cyanide solution, adds 10 c.c. ammonia solution, and digests the solution at 25°C. for six days, boils with baryta (6.7 kilos.) dissolved in water (20 litres) until all the ammonia is expelled, the hot liquid acidified with sulphuric acid, and boiled to expel hydrogen cyanide, neutralised with baryta, and filtered. The filtrate is concentrated. This syrup contains the lactones of a-heptonic acid and \$-heptonic acid. After standing some weeks nearly the whole of the a-lactone will have crystallised out; the mass is triturated with alcohol of 80 p.c.; the filtrate yields a further crop of a-lactone, and the mother-liquor contains the β -lactone. This may be obtained by neutralising with brucine; the brucine β-heptonate crystallises out, and is purified by recrystallisation; the brucine may be separated by precipitation with baryta, and the barium salt decomposed by sulphuric acid.

 α -Glucohepto-lactone forms rhombic crystals readily soluble in water, and has $\{a\}_b = -55^{\circ}3$; boiled with concentrated hydriodic acid it yields a heptolactone, b.p. $216^{\circ}-230^{\circ}$ at 717 mm., with partial decomposition. About an equal quantity

of normal heptoic acid is also produced (Kiliani, B. 19,1128). By digestion with nitric acid S.G. 1.2 at 40° for 24 hours, a-glucoheptoic acid is oxidised to normal penta-oxy-pimelic acid (Kiliani, B. 19, 1916); this melts at 143°, is easily soluble in water, difficultly in alcohol, and soluble in ether. a-Gluco-heptonic acid yields a hydrazide C,H₁₂O,N₂H₂C₆H₅ m.p. 171°-179° (Fischer, B. 22, 2732).

B-Gluco-heptonic lactone crystallises in delicate needles, m.p. 151°-152°, reduces Fehling's solution, and is very soluble in water; its sp. rot. power is $[a]_{D20}^{\circ} = -67.7^{\circ}$. Its phenylhydrazide $C_1H_{10}O_1N_2H_1$ Ph crystallises from alcohol in yellowish leaflets, and melts at 150°-152°. Heated with pyridine (4 lactone, 4 pyridine, and 20 water) in a sealed tube for three hours it is

converted into the a- acid.

Hydrazine compounds. A concentrated solution of dextrose yields, with phenylhydrazine, dextrose phenylhydrazine $C_{12}H_{18}N_{1}O_{5}$; this forms fine crystals, melting at $144^{\circ}-145^{\circ}$. When heated with an aqueous solution of phenylhydrazine, hydrochloride, and sodium acetate on the waterbath, yellow crystals of phenylglucosazone C₁₈H₂N₄O₄ are formed (Fischer, B. 20, 824). Phenylglucosazone can be prepared by heating on the water bath 1 pt. dextrose, 2 pts. phenylhydrazine hydrochloride, 8 pts sodium acetate, and 20 pts. water; after 10-15 minutes' heating the separation of osazone commences, and after 12 hours the precipitate amounts to about 85-90 p.c. of the weight of dextrose taken. It is purified by recrystallisation from alcohol. Another method which gives satisfactory results is to heat a mixture containing 1 g. sugar in 100 c.c. water and 5 c.c. of a solution containing in 1 litre 400 g. phenylhydrazine and 400 g. glacial acetic acid. According to Maquenne's (C. R. 112, 799) numbers, the yield is low, but by increasing the strength of the sugar solution and keeping up the quantity of the phenylhydrazine acetate solution in proportion, it is materially increased. It is insoluble in water; not very soluble in cold, but fairly easily in boiling alcohol. It melts at 204°-205° with decomposition, and so in m.p. determinations it is important that the heating be done as quickly as possible. It reduces Fehling's solution, and is lesvorotatory (Fischer, B. 17, 579). It dissolves in cold fuming hydrochloric acid, with a dark-red colouration, the osazone group C_cH_s,HN_s:C:C:N_sHC_cH_s being thereby converted into the group .CO.CO. This reaction is common to all the osazones of the sugars, and Fischer (B. 21, 2681) has called the compounds thus obtained -osones. Dextrosone reduces Fehling's solution on boiling, and does not ferment with yeast; with phenylhydrazine acetate it again yields phenylglucosazone. The osazone is reduced by zinc and acetic acid to iso-glucosamine, which yields lævulose by treatment with nitrous acid (Fischer, B. 19, 1920; 20, 2569). The osone yields compounds with methyl phenylhydrazine and ortho-diamines (Fischer, B. 22, 87). By proceeding in the same way o-tolyldextrosazone C₂H₂N₂O₄ m.p. 201° and p-tolyl-dextrosazone C₂H₂N₂O₄ have been prepared (Baschen, A. 239, 229). Diphenyl-hydrazine sembines with the sugars in alcohol solution; sparingly soluble crystalline hydrazones are

formed, which are resolved into their constituents by concentrated hydrochloric acid. The dextrose compound C₂H₁₂O₂N₂(C₂H₃)₂ melts at 161° (Stahel, A. 258, 242). Roder (A. 286, 229) has prepared phenyldextrosazone carboxylic acid.

Compounds with aromatic amines. Dextrose - o - diamido - benzene C_eH₄(NH)₂·C_eH₁₀O₈ and anhydro - dextrose - o - diamido - benzene C_eH_4 : N_2 : $C_eH_{10}O_4 + 2H_2O$ are formed by the interaction of dextrose and o-phenylene-diamine in acid solution. Di-dextrose-o-diamido-benzene is obtained when the reaction takes place in neutral solution. They are all crystalline bodies, soluble in water, and having a bitter taste. The first one does not reduce Fehling's solution, the last two do (Griess a. Harrow, B. 20, 281 and 2205); o-tolylene-diamine gives a similar compound (Hinsberg, B. 20, 495). Dextrose- γ -diamido-benzoic acid is obtained by mixing boiling aqueous solutions of dextrose and γ -diamidobenzoic acid (Griess a. Harrow, l.c.). aniline dextrose forms a crystalline anilide m.p. 147° (Schiff, A. 154, 30; Sorokin, B. 19, 513; J. R. 1887, 877). Dextrose forms compounds with aldehyde, acetone, benzaldehyde, chloral, &c. (Schiff, A. 244, 19; Heffter, B. 22, 1050).

Dextrose-oxim C.H.,O.N is formed by the combination of dextrose and hydroxylamine. It forms colourless, microscopic crystals, m.p. 136°-137°; very soluble in water, sparingly soluble in alcohol, and insoluble in ether. reduces Fehling, and is laworotatory, having

[a]_{D200} = -2.2° (Jacobi, B. 24, 696). On warming an alkaline solution of dextrose with picric acid a blood-red colour is produced (Braun, Fr. 4, 185). If to crystallised diazobenzene sulphonic acid be added soda, dextrose solution, and a little sodium-amalgam, a red colourstion is produced in a short time (Penzold a. Fischer, B. 16, 657).

Many aromatic alcohols give a colouration with dextrose and other carbohydrates in the presence of sulphuric acid. If two drops of a 15-20 p.c. alcoholic solution of (α) -naphthol be added to a dextrose solution, and one part of concentrated sulphuric acid, a violet colouration will be produced, changing to bluish violet.

Optical isomerides of glucose, gluconic acid and saccharic acid (E. Fischer, B. 23, 2611).

l-Gluconic scid. l-Mannonic scid can be converted into l-gluconic acid by heating with quinoline, but it is most conveniently prepared from arabinose by treatment with hydrogen cyanide and baryta (Kiliani, B. 19, 3029); l-mannonic and l-gluconic acids are formed simultaneously. The greater portion of the I-mannonic acid may be separated as lactone, and the remainder may be separated from the l-gluconic acid by crystallising out the calcium l-gluconate.

i-Gluconic acid is obtained by mixing the I- and d- acids, or from i-mannonic acid by

quinoline.

L-Glucose is formed by reducing L-gluconic lactone with sodium-amalgam; it resembles dextrose closely, but does not ferment with yeast, $[a]_{D} = -51.4^{\circ}$; its derivatives also resemble the

corresponding ones of dextrose very closely.

i-Glucose is prepared by mixing i- and
d- glucose, or reducing i-gluconic acid.

l-Saccharic acid is prepared by oxidising l-gluconic acid with nitric acid.

i-Saccharic acid is prepared by oxidising i-gluconic acid or by mixing l- and d- saccharic acid.

Lævulose. d-Fructose, lævoglucose, fruit sugar; $C_sH_{12}O_s$.

Occurrence.—Lævulose is widely distributed in the vegetable kingdom, being found, together with dextrose, in many parts of plants, in honey, &c. (v. Dextrose).

&c. (v. Dextrose).

Formation.—The following carbohydrates yield lævulose, either alone or accompanied by other sugars, when acted on by dilute acids:

Cane sugar yields dextrose and lævulose. Raffinose yields lævulose, galactose, and dextrose.

Inulin yields lævulose.

Irisin (O. Wallach, A. 234, 364), lævulan (v. Lippmann, B. 14, 1509), triticin (Reidemeister, C. C. 1880. 808), scillin (Riche a. Remont, J. Ph. [5] 2, 291), sinistrin (Schmiedeberg, B. 12, 704), lævulin or synanthrose (Dieck a. Tollens, A. 198, 228; Ville a. Joulie, Bl. [2] 7, 262; Popp. A. 156, 185; Müntz, C. R. 287, 679), and inuloid (Popp, A. 156, 190) also yield lævulose. Lævulose is formed by the oxidation of mannitol (Cotton, Bl. [2] 33, 546; Gorup-Besanez, A. 118, 273; Bodenbender, Zeit. d. Ver. 14, 812; Renard, A. Ch. [5] 17, 316), by the fermentation of mannitol by B. aceti and B. xylinum (A. J. Brown, C. J. 1886, 184; 1887, 641); from phenylglucosazone by reduction to isoglucosamine, and thence by the action of sodium nitrite; from glucosone by reduction with zinc and acetic acid (Fischer, B. 22, 87, 2566). Preparation.—Inulin is heated with very

weak sulphuric acid for some hours, after which the sulphuric acid is carefully precipitated by baryta water and the filtrate decolourised by charcoal and evaporated on the water-bath to a syrup. This is treated several times with absolute alcohol, which removes the water, and the syrup which remains put on one side for some time in a closed flask, when fine needles make their appearance, and in time the whole solidifies (Jungfleisch a. Lefranc, C. R. 93, 547; also Crookewitt, A. 45, 184; Krusemann, A. 282, 26; Wohl, B. 23, 2084). A 10 p.c. solution of cane sugar is carefully inverted with hydrochloric acid (20 c.c. to a litre of solution) at 60°. The solution is cooled to -5° , 6 g. of slaked lime added for each 10 g. sugar, the temperature allowed to rise to 2°, and the lime lavulose carefully pressed from the rest of the solution and washed repeatedly with water. It is finally suspended in water, decomposed by oxalic acid, and the filtered solution evaporated in vacuo to a syrup which, when treated with absolute alcohol as above described, yields crystals of lævulose (Girard, Bl. [2] 33, 154; Dubrunfaut, C. R. 42, 901; 69, 438)

Formula and synthesis. See general part. Properties.—Lævulose forms colourless, fine, shining needles, belonging to the rhombic system, the axial ratio is a:b:c=0.80067:1:090674; they are slightly biaxial as regards their action on polarised light; they have a composition corresponding to the formula C₂H₁₂O₄, melt at 95°, and lose water at 100°. According to Hönig a. Jesser (M. 9, 562), the crystals con-

tain water of crystallisation and have a formula 2C,H12O,H2O. If free from alcohol they are not deliquescent in the air (Hönig a. Schubert, M. 8, 529; Jungfleisch a. Lefranc, l.c.). The sugar is almost insoluble in cold absolute alcohol, and soluble in a boiling mixture of alcohol and ether (Diech a. Tollens, A. 198, 228). With regard to the specific rotatory power of lavulose there are many and varied statements; all those based on the factors of unpurified solutions obtained by the inversion of inulin or cane sugar are worthless, as lævulose is extremely easily decomposed by acids (Jungfleisch a. Grimbert, C. R. 108, 144), and many apparent discrepancies may be explained by the fact that the optical activity varies very much with the temperature. Jungfleisch a. Grimbert (C. R. 107, 390) found for crystallised levulose $[a]_b$ = -101·38° -0·56t + 0·108 (p-10), where t is the temperature and p the weight of lævulose in 100 c.c. solution. This formula holds for temperature 0°-40° and for concentrations below 40 p.c. The $[a]_j = -108^\circ$ (c = 10 g. sugar in 100 c.c.), calculated from $[a]_j = -24^\circ$ 5 for invert sugars, and $[a]_j = +58^\circ$ 5 for dextrose agree very well with the numbers of Jungfleisch a. Grimbert. Hönig a. Jesser (M. 9, 562) give $[a]_1 = -111.38, c = 10, t = 20^{\circ}$ for a carefully crystallised preparation. Herzfeld a. Winter (B. 19, 390) give for 20 p.c. solution at 20° [a]_D = -71.4° ; but these determinations cannot be for the pure body. Freshly-prepared solutions of crystallised lævulose have a greater sp. rot. power than after standing a short time; Parcus a. Tollens (A. 257, 160) found for a 10 p.c. solution at 20° observed in a 200 mm. tube, [a]_p = -104.02° six minutes after commencing solution, which sank to a constant value $[a]_{D} = -92.25^{\circ}$ in 35 minutes. This cannot have been pure lævulose. K = 100; but Soxhlet makes it 92-93 (J. O'Sullivan, C. J. 1892, 408).

Action of heat and acids. Leevulose is exceedingly easily decomposed by heat and acids. Its aqueous solutions colour exceedingly quickly if heated for any length of time. Wohl (B. 23, 2084) found that by the action of minute quantities of acid on a hot concentrated solution of lævulose, a dextrin-like body was produced which was reconverted into lævulose by boiling with dilute acids. This instability accounts for the many different numbers which have been given for the sp. rot. power and for the fact that the opt. act. of invert sugar does not in all cases correspond with that of a mixture of pure lævulose and dextrose in equal quantities. On boiling with dilute sulphuric or hydrochloric acids, acetopropionic, formic, and other acids are produced (Jungfleisch a. Grimbert, C. R. 108, 144; Grote a. Tollens, A. 175, 181; Sieben, Zeitschr. d. Ver. 84, 237; Conrad a. Guthzeit, B. 19, 2569).

Action of alkalis. Alkalis quickly decompose lævulose, especially on warming; caustic soda and baryta yield lactic acid among other products (Sorokin, J. R. 1885, 368). Sodium lævulose C₂H₁₁NaO₂ is produced by acting on an alcoholic solution of lævulose with sodium ethylate (Hônig a. Rosenfeld, B. 12, 45). Lime yields a compound C₂H₁₂O₂.CaO.H₂O, which is soluble in 187 parts cold water (E. Péligot, C. R. 90, 153; Herzfeld a. Winter, B. 19, 890); and 2(C₂H₁₂O₂)SCaO, which forms needles solu-

ble in 838 parts cold water (Dubrunfaut, A. Ch. [3] 21, 169). Similar compounds are formed with potash, baryta, lead oxide, and bismuth oxide (Herzfeld a. Winter, B. 9, 390, and v. Lippmann, Zuckerarten, p. 68). If a solution of lævulose be boiled with lime, saccharin is formed (Scheibler, B. 13, 2212).

Reduction.—Lævulose in alkaline solution is reduced by nascent hydrogen to mannitol, and more easily than dextrose (Scheibler, B. 16, 3010; H. Krusemann, B. 9, 1465; Herzfeld, A. 244, 274). Fischer (B. 23, 3684) finds that equal parts of mannitol and sorbitol are pro-

duced.

Oxidation.—Mercuric oxide and baryta boiled with an aqueous solution of lævulose yield formic, glycollic and normal trioxybutyric acid (Börnstein a. Herzfeld, B. 18, 3353). Bromine water in the same way yields normal trioxybutyric acid (Hönig, B. 19, 171; Herzfeld a. Winter, B. 19, 390; C. C. 1886, 271). Lævulose by oxidation always yields an acid containing less carbon atoms than itself (Herzfeld a. Börnstein, C. C. 1886, 187; Herzfeld a. Winter, C. C. 1886, 271; B. 19, 890). Chlorine and bromine with silver oxide oxidise lævulose to glycollic acid (Hlasiwetz a. Habermann, A. 155, 130; Kiliani, A. 205, 175). Nitric acid yields oxalic acid, tartaric acid, glycollic acid, formic acid, a little saccharic acid, malic acid, acetic acid, and racemic acid (Dragendorf, Materialien su einer Monographie des Inulins; Kiliani, A. 205, 162; Hornemann, J. pr. 89, 283).

Fermentation.—Lævulose ferments easily with yeast, with formation of alcohol and carbonic acid &c., but rather slower than dextrose, so that towards the end of a fermentation of invert sugar the solution is always lævorotatory (Bourquelot, A. Ch. [6] 9, 245; C. R. 101, 958; C. R. 100, 1404 and 1466; Maumené, C. R. 100, 1505; 101, 695; Leplay, C. R. 101, 479; Gourgon

a. Dubourg, C. R. 110, 865).

Compounds with acids. Chlorosulphonic acid gives a tetra-sulphate (Claësson, J. pr. [2] 20, 1 a. 18). Acetyl chloride yields an acetochlorhydrin (Colley a. Wachowitsch, B. 13, 2390; Bl. [2] 34, 326). With acetic anhydride, acetic Bl. [2] 34, 326). acid, and zinc chloride lævulose yields pentacetyl lævulose (Erwig a. Koenigs, B. 23, 672). Benzoyl chloride yields pentabenzoyl lævulose, m.p. 85° (Skraup, M. 10, 389).

Compounds with bases. See Action of alka-

lis above.

Compounds with metallic salts. Compounds with potassium or sodium chloride have not been described. A compound with lead chloride is known (Herzfeld a. Winter, B. 19, 390).

Compounds with cyanogen. By treatment with strong hydrocyanic acid, lævulose yields a cyanhydrin C,H₁₈O₆N, this crystallises from water in tabular monoclinic crystals, m.p. 110°-115°, and has a slight dextrorotatory power; fuming hydrochloric acid converts it into the lactone of fructoheptonic acid C,H₁₂O, and this, heated with hydriodic acid and phosphorus, yields a heptolactone C,H,2O2, b.p. 220°, which does not solidify in a freezing mixture. When the lactone is further treated with hydriodic acid and phosphorus an acid is obtained boiling at 210°, which is identical with methyl-butyl-acetic axid as symbolically prepared (Kiliani, B. 18, vor. IV.

3966; 19, 221 and 1914; Kiliani a. Düll, B. 23, 449). Tetra-oxy-butane-tri-carboxylic acid COOH.(CH.OH), C(OH)(COOH), is obtained by oxidising fructoheptonic acid by nitric acid (Düll, B. 24, 348).

Hydrazine compounds. Lævulose yields an osazone which appears to be identical with, and is prepared in the same manner as, dextrose

osazone (q.v.).

Anilides. By heating an aqueous solution C12H1, NO5; it is easily obtained in crystals which melt at 147° with partial decomposition (Sorokin, B. 19, 513; J. R. 1886, 129; 1887, 377). With hydroxylamine lævulose combines to form lævulosoxim (Wohl, B. 24, 993). It strongly resembles dextrosoxim in appearance, m.p. 1185, it is lævorotatory, and reduces ammoniacal silver solutions on warming, whilst dextrosoxim does so in the cold.

Galactose. Lactose. C₆H₁₂O₆. Galactose

does not occur in nature.

Formation.—By the action of dilute acids on many carbohydrates and similar substances of vegetable and animal origin. Lactose yields dextrose and galactose (Pasteur, C. R. 42, 347; Fudakowski, B. 9, 42; C. C. 1877, 6). Raffinose yields lævulose, galactose, and dextrose. a-Galactin, B-galactin, paragalactin, lactosin, agar-agar, carragheen moss, plum gum, peach gum, gum arabic, gedda gum, and many other similar vegetable substances, yield galactose amongst other products of their hydrolysis (Müntz, Bl. [2] 37, 409; A. Ch. [6] 10, 566; C. R. 94, 453; 102, 624, 681; A. Meyer, B. 17, 690; Bauer, J. pr. [2] 30, 375; L. V. 1898, 33 a. 214; Hädicke, Bauer a. Tollens, A. 238, 302; Kiliani, B. 13, 2304; 15, 36; v. Lippmann, B. 17, 2238; Scheibler, B. 1, 58; 6, 612; 17, 1729; O'Sullivan, C. J. 1884, 41; 1391, 1029; Hädicke a. Tollens, A. 238, 308; Schulze a. Steiger, B. 19, 827; 20, 290; L. V. 36, 391; Schulze, B. 25, 2213; Maxwell, Am. C. J. 12, 51 a. 265; v. Lippmann, Deutsche Zuckerind., 1887, 455; B. 20, 1001; Stone, Am. 12, 435). Galactose is also producel by the long-continued action of dilute sulphuric acid on certain nitrogenised constituents of the brain (Thudichum, Ann. of Chem. Med. [2] 209; Thierfelder, Z. P. C. 14, 209; Brown a. Morris, C. J. 1890, 57). This sugar was called by the discoverer cerebrose; B. a. M. proved it to be galactose.

Preparation.—A solution of lactose (milksugar) containing in every 100 c.c. 25 g. dry lactose; 2 to 3 grams H SO, is digested in a flask on the water-bath for 2; to 3 hours, the water being kept boiling all the time and care being taken by continual shaking of the flask to avoid colouring. The acid is, on cooling the solution, carefully neutralised with baryta water and the BaSO, separated by filtration. On evaporating the filtrate to a strong syrup in a vacuum solidification takes place in a short time. The solid crystallised mass is digested several times with dry methyl alcohol; this dissolves out the dextrose and other products of the action of the acid on lactose and the hydrolysed sugars, and leaves galactose as a white crystalline powder.
This is dissolved in the least possible quantity of boiling water, again allowed to solidify, and again treated with boiling methyl alcohol. If

the whole of the lactose hydrolysed, and it generally is under the conditions indicated, the residue is white, nearly pure, galactose. It can be rendered pure by recrystallisation from water. If the crystals are allowed to form slowly from solutions not too concentrated they are distinct and well-defined (Rendell, N. Z. 4, 163; Soxhlet, J. pr. 2, 21, 269). The sugar can also be prepared from certain gums of the arabin group. For this purpose the most suitable are those having a lævorotatory power and yielding the larger percentage of mucic acid on oxidation with nitric acid. The dextro-gums also yield the sugar, but in less quantity (O'Sullivan, C. J. 1884, 41; 1891, 1029). A solution of the gum selected—gum arabic, Turkey, Levantine or East India gum -is made containing about equal weights of water and gum; this is heated in a flask in a water-bath as high as the boiling bath will take it, and 5 grams H,SO, diluted to 15 c.c. added for every 100 c.c. of the gum solution employed. The mixture is digested for 30 to 35 minutes, then rapidly cooled, and alcohol, S. G. 0.83, added as long as a pp. is produced. This pp. is again dissolved in water, and again ppd. One or two repetitions of this process frees the pp. from arabinose. A solution of the same strength as that above described for the original gum is then made of the pp. and heated as high as the boiling water of a water-bath will take it, and then for every 100 c.c. of the solution 5 grams H2SO4 diluted to 15 c.c. added. At the end of about 3 hours' digestion the solution is cooled, carefully neutralised with baryta water, and alcohol, 0.82, added as long as a pp. is thrown out. The alcoholic solution becomes clear on standing; it contains galactose, and on concentration in a vacuum yields crystals of the body in a fairly pure state. They may be completely purified by washing with methyl alcohol, recrystallising the residue from ethyl alcohol, and finally from water.

Formula and synthesis. See general part. Properties.—Galactose crystallises in hexagonal plates, m.p. 163°-164° (after drying at 100°) (E. Bourquelot, J. Ph. [5] 18, 51), or, according to v. Lippmann (B. 18, 3335), in microscopic six-cornered prisms and needles, m.p. 168°. Galactose is easily soluble in water, almost insoluble in methyl alcohol. Solutions of galactose rotate a ray of polarised light to the right, the specific rotatory power for the D line being given by the formula [a]_D = 83·883 + 0·0785P - 0·209t (Meissl, J. pr. [2] 22, 97); this formula agrees well with numbers found by other observers. Freshly-prepared solutions exhibit the phenomenon of bi-rotation (Pasteur, J. pr. 68, 428). Parcus a. Tollens (A. 267, 160) observed an angle of [a]_D = 117·48° in a 10 p.c. solution at 20°, 7 minutes after commencing solution. K=100, and varies slightly with the conditions of determination.

Action of acids. By prolonged heating with dilute acids, galactose yields aceto-propionic and formic acids (Kent a. Tollens, A. 227, 228; B. 17, 668; Conrad a. Guthzeit, B. 18, 2905); sulphuric acid and chloro-sulphuric acid yield galactose-sulphuric acids (Hönig a. Schubert, M. 7, 455; Classon, J. vr. [2] 20 a. 18).

M. 7, 455; Claësson, J. pr. [2] 20 a. 18).

Action of alkalis. Alkalis colour galactose yellow; the action of milk of lime produces saccharia (Cuisinier, v. Lippmann, Zuckerarten).

Fusion with KHO gives rise to acetic and oxalic acids.

Reduction. — By reduction with sodium-amalgam, galactose yields dulcitol and other alcohols (Bouchardat, A. Ch. [4] 27, 79).

Oxidation.—Bromine oxidises galactose to galactonic acid (lactonic acid) C₆H₁₂O₇, isomeric with gluconic acid (Hlasiwetz a. Habermann, A. 122, 96; Kiliani, B. 13, 2307; 18, 1551; Bauer, J. pr. [2] 30, 379); which when heated loses 1 mol. water, and is converted into the lactone; it yields with phenylhydrazine a hydrazide which melts at 200°-205° with decomposition (Fischer a. Passmore, B. 22, 2731). Nitric acid oxidises galactose to mucic acid, isomeric with saccharic acid. As mucic acid is only slightly soluble in water, the yield of mucic acid, which is normally 75 to 78 p.c., is sometimes made use of for the estimation of galactose; but the yield varies considerably with the condition of experiment (Pasteur, C. R. 42, 349; Kent a. Tollens, A. Ch. 227, 228). Neutral or alkaline solutions of cupric hydroxide are reduced by galactose; carbonic, formic, glycollic, and lactic acids are produced (Habermann a. Hönig, M. 5, 208).

Fermentation.—According to Kiliani (B. 18, 2305) and Koch (B. 20, Ref. 145), galactose does not ferment with yeast, whilst v. Lippmann (B.17, 2238; 20, 1001), Bourquelot (C. R. 106, 283), and Pasteur find that it does ferment. Stone a. Tollens (A. 249, 257; B. 21, 1572) find that it ferments with beer yeast as well as with Mucor racemosus, but slower than dextrose. Certain bacteria yield lactic acid (Pranzmovaski a. v.

Tieghem, B. 12, 2087).

Compounds.—Galactose yields a crystalline compound with sodium chloride (Fudakowsky, Bl. [2] 30, 433). With a solution of baryta in methyl alcohol, galactose yields the compound (C₀H₁₁O₀),Ba₂.BaO (Fudakowsky, *l.c.*). With acetic anhydride, a pentacetate is produced C₀H₁O(C₂H₂O₂), m.p. 62°-66° (Fudakowsky). According to Erwig a. Koenigs (B. 22, 2207) this body melts at 142°, crystallises in rhombic prisms, reduces Fehling's solution, is dextrorotatory, and differs in all its properties from Fudakowsky's galactose pentacetate. aniline, galactose forms an anilide, crystallising in long triclinic prisms, and lavorotatory (Sorokin, J. R. 1887, 377; B. 19,298). With hydroxylamine, galactose yields the oxim C, H13NO, the crystals melt at 175°-176°, and are easily soluble in water and weak alcohol (Rischbieth, B. 20, 2673). With ortho-diamido-benzene, galactose yields galactose-o-diamido-benzene C₆H₄(NH)₂.C₆H₁₀O₅, melting at 246° with decomposition. Galactose-γ-diamido-benzoic acid COOH.C₄H₃(NH)₂-C₆H₁₀O₅ + H₂O crystallises in needles (Griess a. Harrow, B. 20, 3111). With hydrocyanic acid, galactose yields galactoheptonamide C,H₁₅NO,, m.p. 194°; it is converted into the barium salt of galactoheptonic acid C,H,Os by digesting with baryta. The free acid melts at 145° and crystallises in slender needles; it is reduced by hydriodic acid and phosphorus to the lactone of Lormal y-oxyheptoic acid (b.p. 231°) and a very small quantity of normal heptoic acid (Maquenne, C. R. 106, 286; Kiliani, B. 21, 915; 22, 521 a. 1885). With phenylhydrazine, galactese forms galactese-phenylhydrazide, orystaliteing in colourless

needles, m.p. 158°, and galactose-osazone, m.p. 193°-194° (Fischer, B. 20, 821). Fischer's first determination of the melting-point (B. 17, 579) was 182°, Scheibler (B. 17, 1731) gave 171°, and other observers have given lower numbers; this may be accounted for by the fact that unless the osazone be heated up very quickly it decomposes before it fuses. By the action of hydrochloric acid it yields galactose-osone (Fischer, B. 22, 87). Galactose also yields a diphenylhydrazide, m.p. 157° (Stahel, A. 258, 242). Galactose forms a penta-benzoate C₈H,O(C,H₈O₂)₃, m.p. 165° (Skraup, M. 10, 389).

Sorbose. Sorbinose C.H. 2O6. Sorbose is obtained from the juice of the mountain ash after allowing it to stand for some time. It is doubtful if it exists in the fresh juice as a free sugar, but it is probably formed by the decomposition of a glucoside (Pelouze, A. Ch. [3] 35, 222; Boussingault, C. R. 74, 939; Delffs, B. 4, 799; Byschl, J. pr. 62, 504; Freund, M. 11, 560). The expressed juice of the unripe berries of the mountain ash is allowed to ferment for eight or ten months, the clear solution is filtered from the organisms by which the fermentation was effected, and evaporated to a syrup, when the scrbose crystallises out on standing. It forms colourless rhombic crystals easily soluble in water. Its solutions are lavorotatory: for 10 p.c. solution, $[a]_D = -43.4^{\circ}$ (Wehmer a. Tollens, B. 19, 708). It is oxidised by chlorine and silver oxide to glycollic acid; it behaves with bromine water like lævulose, being practically unaltered (Kiliani a. Scheibler, B. 21, 3276); nitric acid oxidises it to oxalic, aposorbic and tartaric acids. By more careful oxidation with HNO. Kiliani a. Scheibler (l.c.) obtained trioxy-glutaric acid. Copper hydrate oxidises it to carbonic and formic acids and other bodies, amongst which is possibly glyceric acid (Dessaignes, J. pr. 89, 813; Hlasiwetz a. Habermann, A. 155, 129; Habermann a. Hönig, M. 5, 208). Hydriodic acid reduces it to hexyl iodide (Kiliani a. Scheibler, l.c.). It reduces Fehling's solution, and is coloured yellow by alkalis. Sorbose Sorbose ferments slowly with beer yeast (Stone a. Tollens, A. 249, 257). By prolonged heating with dilute acids it is decomposed, yielding, amongst other bodies, lævulinic acid (Wehmer a. Tollens, l.c.). With phenyl-hydrazine it yields an osazone C₁₈H₂₂N₄O₄, m.p. 164°, which is almost insol. in water, but fairly easily soluble in alcohol, from which it crystallises in fine microscopic needles (Fischer a. Tafel, B. 19, 1920; 20, 217 and 2566; Fischer, B. 20, 828).

Mannose. Seminose C.H.,O. This sugar does not occur free in nature.

Formation.—It is obtained by the action of dilute sulphuric acid on the so-called reserve cellulose (Reiss, B. 22, 609, 3218) and on salep juice (Tollens a. Gans, A. 249, 256), and by the oxidation of mannitol (Fischer a. Hirschberger, B. 21, 1805; 22, 1155 a. 3218; Carlet, J. 1860. 250; Gorup-Besanez, A.118, 257; Dafert, B. 17, 227).

Preparation. — Sifted ivory nut shavings (1 pt.) are digested with 6 p.c. hydrochloric acid (2 pts.) for six hours on the water-bath, filtered hot, and the residue pressed and extracted with water. The brown solution contains the sugar. It may be obtained as hydraxide by adding

phenyl-hydrazine acetate, and this is decomposed by hydrochloric acid with reproduction of the sugar (Fischer a. Hirschberger, B. 22, 365 a. 3218).

Properties.—Mannose is precipitated from its alcoholic solution by ether; the syrup, on keeping under absolute alcohol, solidifies to a hard colourless mass which shows no sign of crystalline structure. Its solutions are dextrorotatory, $[a]_D = 13.0^{\circ}$ approx.; they are reduced by Fehling's solution, 1 c.c. of this solution being reduced by 4:307 mgm. mannose, i.e. $K = 110^{\circ}-112^{\circ}$. It is fermented by yeast.

Reduction.—Mannose is reduced by sodium-

amalgam to mannitol.

Oxidation.—Mannose is oxidised by bromine to mannonic acid, the lactone of which $(C_{\bullet}H_{10}O_{\bullet})$ crystallises in colourless needles, readily soluble in water and less in alcohol; it is dextrorotatory in aqueous solutions, $[a]_{\rm D}=53\cdot8^{\circ}$, and forms a phenylhydrazide $C_{12}H_{18}N_{.}O_{e}$, m.p. $214^{\circ}-216^{\circ}$. Its optical isomeride, l-mannonic acid, is obtained by the nitrile reaction from arabinose. Oxidised with nitric acid, manno-saccharic acid is obtained (Easterfield, C. J. 1891, 306; Fischer, B. 24, 539). It crystallises in colourless needles, m.p. 180°–190°, and $[a]_{\rm D}=201\cdot8^{\circ}$. With ammonia ityields a monamide, and with phenylhydrazine a mono- and di-phenylhydrazide. Kiliani's metasaccharic acid is its optical isomeride (l-mannosaccharic acid) (Fischer, l.c.).

Action of acids. Hot hydrochloric acid has little action on mannose, but if the heating be continued for some time, humus substances are

formed

Action of alkalis. Alcoholic potash precipitates a flocculent hygroscopic substance

from solutions of the sugar.

Compounds. - Lead acetate and ammonia produce a precipitate which becomes yellow on keeping. With phenylhydrazine it yields a sparingly soluble hydrazide, $C_{12}H_{18}N_2O_3$, crystallising in yellow slender prisms, m.p. 195°-200°, lævorotatory in dilute hydrochloric acid solution. By heating the hydrazide with excess of phenylhydrazine hydrochloride, sodium acetate, and water, it yields an osazone C18H27N4O4 crystallising in yellow needles, which is identical with phenylglucosazone. With di-phenyl-hydrazine it yields a di-phenyl hydrazide, m.p. 155°. With hydrocyanic acid it yields a compound which is easily decomposed, yielding mannose-heptonic acid; its lactone $C_1H_{12}O_2$, crystallises in needles, m.p. 148°-150°; by reduction with hydriodic acid normal heptoic acid is obtained. With hydroxylamine mannose yields an oxim C.H.2O.N, m.p. 184° (Reiss, l.c.; Fischer a. Hirschberger, B. 22, 1155). Aceto-chloro-mannose is obtained by the action of acetyl chloride (Fischer, l.c.).

Phlorose C_eH₁₂O_e. This has been shown to be identical with dextrose (Rennie, C. J. 887,

636; Fischer, B. 21, 988).

Crocose $C_eH_{12}O_e$. Kayser (B. 17, 2232) a. Rochleder a. Mayer (J. pr. 74, 1) obtained a dextrorotatory sugar by heating saffron (from Crocus electus or yellow shoots of Gardenia grandiflora) with dilute sulphuric acid. It forms rhombic crystals, which only reduce half as much copper oxide as dextrose. According to Fischer (B. 21, 988), it yields, with phenylhydrazine, as eassone identical with phenyl glucosestone.

Lokaose C₆H₁₂O₆ is obtained from lokao or Chinese green by boiling with dilute sulphuric soid, loksonic soid yielding lokanic soid and lokaose $C_{12}H_{48}O_{27} = C_{36}H_{36}O_{21} + C_6H_{12}O_6$. Lokaose forms minute acicular crystals, and is distinguished from dextrose by being optically inactive and having a lower reducing power (K = 50) (R. Kayser, B. 18, 3417).

Tabakose is said to exist in tobacco (Att-

field, Ph. 541).

Digitalose C, H14O, is obtained, together with dextrin, by the action of strong hydrochloric acid on digitalin; oxidised with bromine it yields digitalonic lactone C,H12O; hence the formula of the sugar (Kiliani, Ar. Ph. 230, 250;

B. 25, 2116).

Aromatic sugar. Phenyltetrose (Fischer a. Stewart, B. 25, 2555). Cinnamaldehyde cyanhydrin, when dissolved in chloroform and bromine added, yields phenyl-dibromo-oxybutyronitril CHPhBr.CHBr.CH(OH)CN; this when heated with hydrochloric acid yields phenylbromodioxybutyrolactone

CHPh CH.Br CH.OH, from which the acid CHPh.CHBr.CH(OH).COOH is easily obtained; on reduction this yields phenyltetrose OH.CHPh.CH(OH).CH(OH).COH, It yields a

phenylhydrazide, m.p. 154°

For Acrose, Formose, Glycerose, see Synthesis of the Sugars.

or arabinbiose

CLASS II. The '-on' sugars. (a) Dipentose.

Arabinon, diarabinose, $C_{10}H_{18}O_9$. A product of the partial hydrolysis of a series of gum acids. A 25 p.c. solution of any of the strongly rotating gum acids is heated to boiling, and 2 g. sulphuric acid for every 100 c.c. solution, previously diluted with 4 to 5 volumes water, added. At the end of 10 to 15 minutes' digestion the solution is quickly cooled and alcohol, sp.gr. '830, added as long as a precipitate forms. The clear alcoholic solution contains arabinon and arabinose, and some arabinon is taken down with the precipitate. and on again dissolving it in a little water and reprecipitating with alcohol, the arabinon often appears on the surface of the syrup in spherocrystals, and the alcoholic supernatant liquid contains it in quantity. It is easily soluble in water and methyl alcohol. Dilute ethyl alcohol also dissolves it, but absolute alcohol precipitates it as a syrup from a strong methyl alcohol solution. Ether also precipitates it. Its solutions are dextrorotatory, $[\alpha]_{D} = +202^{\circ}$ (c. 6.466), there

acids hydrolyse it, the sole product being ara- $\mathbf{C}_{10}\mathbf{H}_{18}\mathbf{O}_{9} + \mathbf{H}_{2}\mathbf{O} = 2\mathbf{C}_{1}\mathbf{H}_{10}\mathbf{O}_{8}$ Arabinon Arabinose

appears to be an increase with the concentration;

it reduces Fehling's solution; K = 58 (K - 57.5) if

2 mols. sugar reduce 9 mols. CuO). D=8.95.

It tastes sweet, and is easily diffusible. Dilute

(O'Sullivan, C. J. 1890, 59).

(b) Dihexoses.

binose,

Cane sugar. Saccharon, sucrose C₁₂H₂₂O₁₁. Occurrence. — Cane sugar is widely distributed throughout the vegetable kingdom, and is frequently accompanied by dextrose and lævulose. The most important sources are the sugar-cane (Saccharum officinarum); the sugar-beet (Beta

vulgaris); the sugar-maple (Acer saccharinum) and Sorghum saccharatum (Girard, C. R. 102, 103; Centrol. f. Agric. Ch. 1886. 683; Berthelot, C. R. 53, 583; A. Ch. [3] 55, 289; Buignet, A Ch. [3] 61, 233; Maxwell, L. V. 36, 15, Am. 12, 265; Reports of the Department of Agriculture, Washington, No. 6; The Sugar Industry of the United States, by Wiley, 1885; The Literature of the Sugars, H. L. Roth, London, 1890; Gössmann, A. 104, 335; Hermbstädt, Gehlen's J. f. Ch. Ph. Min., 8, 589 (1809); Beier, Jahresb. f. Zuckerfabr. 5, 189; Balland, J. Ph. [4] 25, 97; Corenwinder, C. R. 88, 1238; A. v. Wachtel, B. C. 1880, 344; Stone, B. 23, 1406; Wiley, C. N. 51, 88; Ann. Agronom. 11, 392; Schulze a. Seliwanoff, L. V. 1887. 403; Washburn a. Tollens, B. 22, 1047; Stingl a. Morawski, M. 7, 176; 8, 82; Déon, Bl. [2] 32, 125; Reali, G. 17, 325; Schulze, L. V. 1887, 403, 408; v. also the references under the heading Dextrose, Occurrence, and many others throughout the

range of chemical literature). Formation.—Icery (A. Ch. [4] 5, 350) and Jackson (C. R. 46, 55) found that in the early period of vegetation of many plants, invert sugar only occurs, and that cane sugar is formed later. Leplay (C. R. 46, 444) confirms this observation in the case of sugar-cane. Barley contains from 0.8 to 1.6 p.c. cane sugar, whereas germinated barley (malt) contains from 2.8 to 6.0 p.c. cans sugar; the starch of the endosperm being converted into cane sugar for the use of the growing plant (Kuhnemann, B. 8, 202, 387; Kjeldahl, Rés. du C. R. des Travaux du Lab. de Carlsberg. 1881, 189; O'Sullivan, C. J. 1886, 58). Brown a. Morris (C. J. 1890, 516) found that the cane sugar of germinated barley was chiefly localised in the embryo, whilst maltose appears in the endosperm; they consider that starch is first converted by diastase into maltose in the endosperm, and then, being transferred to the embryo, is converted into cane sugar for the purpose of growth. Colley a. Vakovitch (Bl. [2] 34, 326) may have obtained cane sugar by acting on barium lævulosate with acetochlorhydrose, but the evidence is not satisfactory. Aubert a. Giraud (D. P. J. 257, 298) state that by passing an electric current through acidified starch-paste at 100°C. cane sugar may be manufactured (?).

Preparation .- (For the commercial preparation of cane sugar see Thorpe's DICTIONARY OF APPLIED CHEMISTRY.) The best samples of commercial cane sugar consist of almost the pure sugar. Dry, clean, glassy sugar-candy is sufficiently pure for most purposes. Absolutely pure saccharon may be obtained by careful crystallisation from warm aqueous solutions; or a cold, clear saturated solution made with carefully-purified distilled water is mixed with an equal volume of purified ethyl alcohol, S.G. ·820, with continual stirring. On standing, saccharon crystallises out in pure glassy crystals.

Properties.-Cane sugar forms fine large monoclinic hemihedral crystals, S.G. $\frac{17.5}{17.5}$ 1.58047. (Schröder, B. 12, 562; Gerlach, D. P. J. 172, 31 and 286; Joule a. Playfair, C. J. 1, 130), which do not contain water of crystallisation. It is easily soluble in water; 100 parts of a solution saturated at 0° contain 65.0 pts. saccharon, at 20° 67 pts., and at 40° 75.8 pts. (Scheibler,

Z. V.' 22, 253). A large number of figures have been given to represent the relation between the quantity of sugar in solution and the specific gravity thereof (v. Lippmann's Zuckerarten; Brown a. Heron, C. J. 1879, 644; Périer, C. R. 103, 1202), and much apparent contradiction exists on the subject; but the bulk of this disappears when the processes by which the figures were obtained are inquired into. The general agreement is with the numbers of Balling, confirmed, within the limits of error, by Scheibler a. Mategczek (Z. V. 15, 586; 24, 827; 27, 32) and others. These are:—

Per cent. saccharon	Sp. gr. sol.
in sol.	17:50
5	1.01970
10	1.04014
20	1.08329
40	1.17943

The relation is sometimes more convenient when expressed as weights in measure. If a vessel be made to hold 100 g. H₂O at 15.5°, and such is the 100 c.c. vessel most generally in use, then a solution containing

20 g. saccharon in this 100 c.c. = sp. gr. 1·07686
10 ,, , 1·03857
5 ,, , 1·01931
1 ,, 1·00386

Hence, if the sp.gr. of a pure cane sugar is known the amount of sugar in the 100 c.c. can be arrived at broadly by dividing the S.G. — 1.000 by .00385. This number multiplied by 1000 has been called the D of the sugar. D = 3.85 for saccharon. The use of the number gives too tow an indication for the higher specific gravities and too high a one for the lower ones, being almost accurate for solutions containing between 10 and 20 g. sugar per 100 c.c. It is a useful, practical factor when its meaning and value are understood, being at most 3 per 1000 incorrect.

Concentrated sugar solutions possess a higher boiling-point than water, an 80 p.c. solution boils at 112°, a 90°8 p.c. solution at 130° (Gerlach, D. P. J. 172, 31 and 286).

Solutions of pure cane sugar crystallise quickly, but if foreign matters are present the crystallisation is very much slower, and the forms of the crystals vary with the character of the impurity; some impurities entirely prevent crystallisation. These facts play a most important part in the refinery (see art. in Dictio-NARY OF APPLIED CHEMISTRY and v. Lippmann's Zuckerarten, p. 115). Solutions of cane sugar are highly diactinic (Hartley, C. J. 1887, 59). Cane sugar is soluble in methyl and ethyl alcohol, acetone, and glycerin in proportion to the amount of water they contain, being almost insoluble in them when they are anhydrous (Scheibler, Z. V. 22, 246; B. 5, 843). Solutions of cane sugar and the fused sugar are optically active, rotating the plane of polarisation to the right; crystallised sugar is not optically active. The oldest value given for specific rotatory power is $[a]_1 = +73.8^{\circ} =$ $[a]_D = +66.6^{\circ} (24:21.67::73.8^{\circ}:66.6^{\circ}).$

This is only an approximate average number, much of the same value as the D described

¹ Z. V.=Zeitschrift des Vereins für die Rübensucker in-Austrie des deutschen Reiches.

above. It is the apparent specific rotatory power, and may be called the working value. Tollens (B. 17, 1757), Schmitz (B. 10, 1419), Girard a. de Luynes (C. R. 80, 1855), and Calderon (C. R. 83, 393), from numerous observations worked out an absolute value. Tollens gives $[a]_{D} = 66.386 + 0.015035P - 0.0003986P^{2}$, where P is the p.c. of sugar; Schmitz's numbers are $[a]_D = 64 \cdot 156 + 0.051596q - 0.00028052q^2$, where q = p.c. of water. The optical activity diminishes with the concentration of the solution, so that for a 100 p.c. solution (i.e. for the dry sugar, if the curve follows the same course as the observations) the specific rotatory power is $[a]_D = 64^{\circ}$ approximately, while very weak solutions, as 1 p.c. for example, it is $[a]_D = 67^\circ$ (Pibram, B. 20, 1849; Nasini a. Villavecchia, Gaz. 22, 1, 97). The temperature of observation has but a very slight influence on the optical activity (Dubrunfaut, A. Ch. [3] 18, 99; Andrews, M. S. [4] 3, 1866). The specific rotatory powers of solutions of the sugar in mixtures of water and ethyl alcohol, methyl alcohol, and acetone respectively are slightly greater than that of aqueous solutions; if the latter be taken at $[\alpha]_p = 66.67^\circ$ they are respectively $[a]_D = 66.83^{\circ}, 68.63^{\circ}, \text{ and } 67.40^{\circ}$ (Tollens, B. C. 1881, 570). The optical activity of fused sugar is less than that of its solutions, and diminishes according to the length of time it has been kept fused. This is, no doubt, due to the presence of decomposition products. The specific rotatory power of aqueous solutions is altered by the presence of most foreign substances; alkalis and alkaline earths diminish it (Thomsen, B. 14, 1649; v. Lippmann, Zuckerarten, p. 157; Farnsteiner, B. 23, 3570). Lead acetate does not appear to have any influence, while ammonia increases the rotation (Ost, N. Zeitsch. f. Rübens .-Ind. 9, 42). The dispersion of cane-sugar solutions is almost the same as that of quartz, so that the rotation of the plane of polarisation caused by a sugar solution may be almost completely neutralised by a plate of left-handed quartz of the correct thickness. Grimbert (J. Ph. [5] 16, 295) gives $[a]_{\rm p} = 66.45^{\circ}$ and $[a]_{\rm o} = 52.85^{\circ}$, so that the dispersive power for these two wave-lengths is 1.257. The refractive index for sugar solutions has been determined for seven lines of the spectrum (Obermeyer, Landolt-Börnstein, Physikalisch-chem. Tabellen, Berlin, 1883, p. 213; Kanonnikoff, B. 16, 3047). Gladstone (C. J. 1891, 589) finds the molecular refraction of cane-sugar solutions to be 118.7 for the A line, and the molecular dispersion between A and H lines 4.78.

Action of heat. Dry cane sugar melts at 160°, and solidifies on cooling, if care be taken and the temperature does not exceed 160°, to a colourless, glassy mass; but if the temperature be maintained the fused mass becomes slightly coloured, and on cooling yields the substance known as barley-sugar, this after a time becomes crystalline; but even with the greatest care it seems impossible to fuse the sugar without producing some decomposition. With the appearance of the colour dextrose and lævulosan are probably the chief products; but although much work has been done on the subject, the exact nature of the alteration is not settled. A little furfural is formed at times (Gélis, A. Ch. [3] 57, 234; C. R. 51, 331; Schiff, B. 20, 540). When the tempe-

rature is raised to 200° distinct decomposition sets in, caramel is formed, gases are evolved, and finally a residue of charcoal remains. Among the gases are carbon dioxide, carbon monoxide, marsh gas, ethylene, acetylene; among the volatile products are water, acetone, formic, acetic, and propionic acids, aldehyde, furfural, acrolein, benzoic aldehyde, and a bitter substance assamur (Völckel, A. 85, 59; 86, 63; 87, 303; Fremy, A. 15, 278; Schiff, B. 20, 540; A. 238, 380; Reichenbach, A. 49, 3). Caramel has the same composition as cane sugar, less one or more molecules of water. It is soluble in water and partly soluble in alcohol, and probably consists of a mixture of several bodies. Baryta and basic lead acetate give precipitates with it. It combines with aniline, reduces some metallic salts, and yields an acid with chlorine. It is largely used as a colouring matter in cookery, brewing, &c. (Gélis, A. Ch. [3] 52, 386; 65, 190 a. 496; Graham, A. Ch. 65, 190; Peligot, A. Ch. [2] 67, 172; Völckel, A. 75, 59; 85, 74; Pohl, J. pr. 82, 148; Schiff, B. 4, 908; Wachtel, Org. d. Centr.-Ver. 17, 930).

It seems proved that when dry saccharon is heated at 100°, even for a long time, no change takes place, and, as has been said, the sugar can be heated to the melting-point, 160°, without producing any marked decomposition, but it would appear that rapid heating to this temperature can produce a mixture of dextrose and lævulosan, without any change in weight. If, however, a mixture of 100 pts. of the sugar and about 5 pts. water is heated at 150° for some time, a mixture is produced which contains little or no unaltered saccharon, but which consists, in part at least, of a sugar said to be optically inactive, and to be a compound of bi-rotating dextrose and Other bodies must also be present, probably dextrose and lævulose. The compound sugar is, no doubt, present, but inasmuch as it has never been isolated it cannot be said with certainty that it is inactive (Morin, C. R. 86, 1033; Berzelius a. Mitscherlich, J. Ph. 3, 4, 216). The so-called inactive sugar is easily converted by boiling with water into dextrose and lævulose. Neutral aqueous solutions of saccharon are slowly inverted on boiling. They are, however, stable in the cold, if sterilised, and can be concentrated under diminished pressure at 50°-60° without undergoing much change (Kreuster, Z. V. 25, 521; Béchamp, A. Ch. [3] 54, 28; Morin, C. R. 86, 1,083; Horsin-Déon, Bl. [2] 32,121; Gunning, Z. V. 27, 895). By heating to 150°-200° in a sealed tube, humus substances, formic acid, pyrocatechin, and other bodies are formed (Löw, Z. 1867, 510; Hoppe-Seyler, B. 4, 15).

Action of acids. All dilute acids, even carbonic acid, hydrolyse (invert) cane sugar, causing the assimilation of one molecule of water and the production of equal quantities of dextrose and lævulose,

Saccharon Dextrose Leevulose $C_{12}H_{22}O_{11} + H_2O = C_6H_{12}O_6 + C_6H_{12}O_6$

This mixture is called invert sugar. The experiments made on this phenomenon may be classified under two headings: (a) the action of different acids acting under the same conditions on the sugar solutions; (b) the action of the same acids under varying conditions. With

regard to those coming under the first heading, Ostwald's experiments are the most complete (J. pr. [2] 29 385; 30, 93 a. 225; 31, 307; Koral, J. pr. [2] 34, 109). He examined the action of thirty-two acids, and finds that there is a close relation between the constants of inversion and the affinities of the acids. See also Fleury (D. P. J. 219, 436; J. Ph. [4] 22, 423). With regard to reactions coming under the second heading, it is found that inversion proceeds more rapidly in hot solutions than in cold ones (several formulæ have been given to express this relation, v. Arrhenius, Zeit. phys. Ch. 4, 226; Urech, B. 20, 1836); that increase in the percentage of acid increases the rapidity of inversion; that, the percentage of acid to water being constant, increase of volume decreases the time; and that the percentage of acid to sugar being constant, increase of volume (dilution) increases the time (Löwenthal a. Lenssen, J. pr. 85, 321 a. 401; Urech, B. 13, 1696; 15, 2130; B. C. 1881, 570; 1883. 501). The presence of neutral salts sometimes increases and sometimes diminishes the rapidity of inversion; for a detailed account see Spohr (J. pr. [2] 32, 32 and 33, 265). If the heating be long continued, or if too strong acids be used, most of the lævulose and part of the dextrose is destroyed, and humus substances, acetopropionic and formic acids are produced (Mulder, J. pr. 21, 219; Malaguti, A. 17, 52; Stein, A. 30, 82; Conrad, B. 11, 2178; Tollens, B. 14, 1951; Sestini, L. V. 26, 285; 27, 163; Conrad a. Guthzeit, B. 18, 439; 19, 2569 a. 2844; Tollens a. Grote, A. 175, 181, 206, 237 a. 233. The inversion of cane sugar cannot be carried out by heating with acids without destroying some lævulose (q. v.), unless the temperature and quantity of acid be kept within welldefined limits (Jungfleisch a. Grimbert, C. R. 108, 144; Wohl, B. 23, 2084). Nicol (Fr. 14, 180) and Clerget (A. Ch. [3] 26, 175) give instructions for the preparation of pure invert sugar. According to the latter, 5 c.c. of strong hydrochloric acid is added to 50 c.c. sugar solution, containing 8.175 g. saccharon, and heated for ten minutes at 68°. Under these conditions the sugar is completely inverted, and the products are unacted upon. They are identical in optical activity and cupric reducing power with the sugar obtained by the action of invertase (see Action of ferments) under the most favourable conditions. There is no doubt that invert sugar is a mixture of lævulose and dextrose in equal quantities (Jungfleisch a. Grimbert, C. R. 108, 144; 107, 390), as its properties agree in every particular with that of a mixture of these two sugars in molecular proportions. Concentrated sulphuric acid dissolves cane sugar at the freezing temperature, without the production of colour, but slight heating causes it to swell up to a black, frothy mass; a blue fluorescent body is formed (Simmler, C. C. 1862, 378). Other acids act in a similar way (Gmelin-Kraut, Hand buch d. Ch.; van Kerckhoff, J. pr. 69, 48).

Action of alkalis. Dilute alkalis do not act

Action of alkalis. Dilute alkalis do not act on cane sugar in the cold, and but slowly on heating (Michaelis, J. pr. 56, 430). By heating with potassium hydroxide and but little water, decomposition takes place, carbon dioxide, acetone, and formic, acetic, propionic, and oxalic acids are formed (Gottlieb, A. 52, 122). Potash

or baryta under certain conditions yield a large quantity of lactic acid (Schützenberger, Bl. [2] 25, 289). Milk of lime and strontia also decompose cane sugar by prolonged heating (Fremy, A. 15, 278; Benedict, A. 162, 303; Niedschlag, Deutsche Zuckerind. 1887. 159; Pinner, B. 16, 1728).

Oxidation. Cane sugar is easily oxidised by almost all the common oxidising agents. Free oxygen does not act on it, but in the presence of platinum black it does; ozone also oxidises it (Gorup-Besanez, A. 110, 103). Chlorine or bromine yields gluconic acid, glucose, and other substances (Reichardt, B. C. 1880, 559; Grieshammer, A. Ph. [3] 15, 193); the same reaction takes place in the presence of lead or silver oxide (Hlasiwetz a. Habermann, A. 155, 128; Herzfeld, A. 220, 353). Iodine and potassium carbonate yield a little iodoform (Millon, C. R. 21,828). The more powerful oxidising reagentssuch as strong chromic acid, chlorate of potash, strong sulphuric acid, &c.—act so violently as to cause explosions. Strong nitric acid acts very energetically; dilute nitric acid yields saccharic and oxalic acids; anhydrous nitric acid, or a mixture of strong nitric and sulphuric acids, yields a nitrate (Sobero, C. R. 24, 247; Schönbein, P. 70, 104; Carey-Lea, Bl. [2] 10, 415; Tollens, B. 15, 1828; Salkowski, B. 15, 1738; Heintz, A. 51, 185; P. 61, 315; Reinsch, Jahrb. f. Pharm. 18, 337; Hornemann, J. pr. 89, 304; Tollens, Chem. Zeit. 11, 1178). Fehling's solution is not reduced by cane-sugar solution, neither are alkaline solutions of other oxides, except ammoniacal silver solutions, which are reduced slowly in the cold and quicker on warming; but many neutral or acid metallic solutions are reduced (Tollens, Z. V. 32, 712; B. 15, 1828; Salkowski, B. 15, 1738). Dilute chromic acid oxidises cane sugar to oxalic, formic, and carbonic acids; potassium permanganate yields carbonic acid and water; by careful oxidation in the cold, oxalic and formic acids are also produced (Heyer, Ar. Ph. [3] 20, 336 a. 430; Brunner, B. 12, 549). Maumené (Bl. [2] 22, 2; 30, 99) says that three acids, di-, tri-, and hexa-pinic acids C2H4O4, C₃H₄O₃, and C₆H₁₂O₃, are formed under certain conditions.

Action of ferments. Cane sugar is not directly fermentable by organised ferments, with the exception of Monila candida; it requires first to be hydrolysed (inverted), and for this purpose every organism which has the power of assimilating cane sugar produces an organised ferment or enzyme called invertase, which first inverts it (Bernard, Lecons de Physiologie Exper., Paris, 1856; Hansen, Meddelelser, 1888, 2, 143; Brown, C. J. 1886. 178). The action of invertase was first studied by Kjeldahl (Meddelelser, 1881, 3, 186); O'Sullivan and Tompson (C. J. 1890, 834) have exhaustively investigated the subject. They find the change follows the same time curve as a catalytic reaction; the time necessary to perform a given percentage of hydrolysis varies directly with the amount of invertase; the most favourable concentration of the sugar solution is 20-25 p.c., and the most favourable temperature 55°. Hydrolysis proceeds slowly at the freezing-point; it ceases at about 65°; alkalis immediately stop the reaction, whereas very minute quantities of acid

aid it. Invertase is uninjured by the reaction, and is capable of inverting about twenty-two times its weight of cane sugar per minute for an unlimited period- The dextrose formed is in the bi-rotatory state; on standing, the invert sugar has the same optical activity as that obtained by the careful action of dilute acids. See also Barth (B. 11, 474), Mayer (B. C. 1881, 784; 1882, 850), Müller (Ann. Agronom. 12, 481), Tamman (Zeit. phys. Ch. 3, 25). Invertase acts at a temperature higher than is sufficient to kill the yeast whence it is derived. For production of alcohol, acids, &c., under the influence of organised ferments, see FERMENTATION.

Compounds with acids. Nitrosaccharon $C_{12}H_{18}(NO_2)_4O_{11}$ is prepared by treating the powdered sugar with a mixture of strong H_2SO_4 and fuming HNO, in the cold (Schönbein, P. 70, 167; Sobero, C. R. 24, 247). It is a white doughy mass, insoluble in water, soluble in alcohol and ether. Boiling water decomposes it:

m.p. 20°. It is explosive.

With arsenious acid a compound

C₁₂H₂₂O₁₁As₂O₃ is said to be produced. The

composition is improbable.

By the action of acetic anhydride and glacial acetic acid, or acetic anhydride and sodium acetate, a series of acetates have been obtained. In these the group C₂H₃O takes the place of H. We have the monoacetate $C_{12}H_{21}(C_2H_3O)O_{11}$ (Schützenberger a. Naudin, Bl. 12, 206); the (Schützenberger & Nadum, B. 12, 2007, the tetracetate $C_{12}H_{18}(C_2H_3O),O_{11}$; the hexacetate (Herzfeld, N. Z. 3, 155); the heptacetate (Schützenberger, Bl. 12, 204; C. R. 61, 485); and the ootacetate (Herzfeld, B. 13, 267; Démole, C. R. 89, 481), all more or less resinous substances, diminishing in solubility in water in proportion to the number of acetyl groups they contain, and increasing in solubility in alcohol and ether with the same change in composition. The octacetate is said to crystallise in needles, m.p. 67°. They are all decomposed by the action of acids and alkalis into acetic acid and the products of the inversion of saccharon. It is probable that by careful treatment with alkalis in equivalent quantities (sufficient to neutralise the acetic acid formed) saccharon is again produced. Benzoyl chloride yields with saccharon a hexabenzoate $C_{12}H_{16}(C_7H_5O)_8O_{11}$, m.p. 109° (Baumann, B. 19, 3220; Skraup, M. 10, 389).

Compounds with bases. By mixing sodium ethylate with a solution of cane sugar and precipitating with alcohol, a compound $C_{12}H_{21}NaO_{13}$ is obtained (Pfeiffer a. Tollens, A. 211, 285). A similar potassium compound has been prepared (Soubeyran, A. 43, 223; Brendeche, An. Ph. [2] 29, 73). Cane sugar combines with many bases, forming saccharates which are decomposed into the sugar and carbonate by carbon dioxide, and are but slightly soluble in water. The most important of these are barium saccharate C₁₉H₂₂O₁₁.BaO, strontium saccharate C₁₂H₂₂O₁₁.SrO, and C₁₂H₂₂O₁₁.2SrO, the former produced by mixing solutions of its constituents; it crystallises with 5H,O, the latter obtained by bringing together its constituents in boiling solution. Calcium saccharates: C₁₂H₂₂O₁₁.8CaO 3aq is but slightly soluble in water; C₁₂H₂₂O₁₁.2CaO 2aq and C₁₂H₂₂O₁₁CaO are soluble in water and decomposed partly by boiling into the tri-saccharate and free sugar.

 $C_{12}H_{22}O_{11}$. $CaO + 2H_{2}O$, $C_{12}H_{22}O_{11}$. 2CaO, and (C₁₂H₂₂O₁₁),3CaO have also been described. Lead acetate alone does not precipitate cane lead acetate and ammonia C12H18Pb2O11 and C12H18Pb2O11; iron and copper yield saccharates; it is doubtful if magnesia forms one (Soubeyran, A. 43, 223; Stromeyer, Ar. Ph. [3] 25, 229; Brendeche, Ar. Ph. [2] 29, 73; Péligot, J. pr. 13, 379; 15, 76; A. [3] 54, 377; Landolt a. Degener, Z. V. 32, 325; Scheibler, B. 15, 2945; 16, 985; v. Lippmann, Organ d. Centr. Ver. 18, 37; B. 16, 2764; Benedict, B. 6, 413; Harperath, Chem. Zeit. 1886. 323; Boivin a. Loiseau, C. R. 58, 60; also old editions of Watts' Dict.). Ammonia gas at 150° forms brown amorphous bodies; solutions of ammonia yield similar bodies (Thénard, C. R. 52, 444; Payen a. Schützenberger, Jahresber. f. Zuckerfabr. 1861-62, p. 191, 192; Laborde, J. 1874, 883).

Compounds with metallic salts. Several compounds of saccharon with sodium chloride, bromide, and iodide have been prepared (Péligot, A. 30, 71; Maumené, Bl. 15, 1; Gill, C. J. 1871, 269). Such compounds are C₁₂H₂O₁₁.Nacl.2H₂O; 2C₁₂H₂O₁₁.3Nacl.4H₂O; H. O. Napp. H. O. $C_{12}H_{22}O_{11}$.NaBr. $1\frac{1}{2}H_{2}O$; and $2C_{12}H_{22}O_{11}$.3NaI.3 $H_{2}O$; the latter is a very stable and definite compound, and is, like the others, easily prepared by boil. ing a mixture of the sodium salt and saccharon in equivalent quantities, and allowing the solution to stand in a quiet place over sulphuric acid. These bodies form distinct and definite crystals. No definite compounds of the halogen salts of potassium and ammonium with saccharon have been described, but there can be little doubt, from the work of Gill, that such are obtainable. Lithium salts seem to give lithium compounds. Compounds of saccharon with sodium-acetate, nitrate, iodate, and phosphate are not obtainable (Gill, l.c.). A double salt of copper suland saccharon, C12H22O11.CuSO. 2aq phate crystallises from a saturated solution of the constituents (Barreswil, J. Ph. [3] 7, 29). The sodium mercury chloride compound $2C_{12}H_{22}O_{11}$. NaCl. $HgCl_2$ crystallises in minute crystals on the slow evaporation of a solution of its constituents in weak alcohol (Boullay, Bl. 12, 292). A borax compound 3C12H22O11.Na2B4O7.4H2Ó has also been described (Stürenberg, Ar. Ph. 18, 279). Many other compounds have been described, but they present no material interest.

Saccharon, like many other organic substances, prevents the precipitation of many metals from their solutions by ammonium, sodium, and potassium hydroxide; in some cases the prevention is complete, in others only a portion of the metal is retained in solution (Lussaigne, C. R. 14, 691; Groth, J. pr. [1] 92, 175;

Pellet, J. Fabr. 18, 22).

With phenyl-hydrazine saccharon yields only lævulose and dextrose compounds (q. v.).

Lactose. Lacton, lactobiose, milk sugar $C_{12}H_{22}O_{11}.H_{2}O(a)$ and $C_{12}H_{22}O_{11}(\gamma)$.

Occurrence.—Lactose has been long known. It occurs in the milk of the mammalia and also in the fruit of Sapotillier (Achras Sapota) (Bouchardat, Bl. [2] 16, 26). Lactose constitutes 8-5 p.c. of the milk of women, sheep, goats, and cows, 6 p.c. of asses' milk, and a slightly greater proportion of mare's milk (Fleischmann, Das Molkereiwesen, Braunschweig, 1875; Kühne, Lehrb. d. physiol. Ch. p. 573).

Preparation.—Skim milk, or, better, milk from which the fat has been separated by mechanical means, is coagulated with rennet and the whey digested for some time with calcium carbonate (chalk) and aluminium hydroxide. The insoluble matter is then filtered out and the filtrate concentrated in a vacuum to a syrup; this, on standing, yields crystals of lactose. During concentration some matter becomes insoluble; this should be filtered out before final concentration. The crystals are purified by repeated re-crystallisation from solutions concentrated at the temperature of boiling water; on cooling C12H22O11.H2O crystallises. This is lactose a. If the saturated solution is evaporated rapidly on the water-bath with continual stirring, small crystals of C12H22O11 are obtained; lactose y. These are not hygroscopic. If the a modification is dried at 130° a hygroscopic mass is left, which is known as the B modification. There are said to be other modifications, but in solution after boiling they all have the same optical activity and are identical (Enling a. Rüf, B. C. 1882, 346; Centbl.

f. Ag. Ch. 1885, 130; J. Kunz, Ph. [3] 15,443). Properties.—Ordinary lactose forms large, rhombic, hemihedral crystais, having a sp.gr. 1.53-1.54 (Schröder, B. 12, 562) and containing 1 mol. H.O, which is lost by drying at 130°. It is soluble in six parts of cold water and in two and a-half or less of boiling water, from which on cooling it slowly crystallises. Its solutions rotate the plane of polarisation to the right, and its specific rotatory power for the D line is $[\alpha]_D = 52.53^\circ + (20 - t) \times 0.055$ (Schmöger, B. 13, 1927; Hesse, A. 176, 98; Erdmann, J. 1855, 661; Denigès a. Bonnans, J. Ph. [5] 17, 363 a. 411). The variation with the concentration is very little, and, as is obvious from the above formula, the effect of temperature is slight. numbers for the [a]D are calculated for the crystals $C_{12}H_{22}O_{11}.H_2O$. For the dry sugar $C_{12}H_{22}O_{11}$ I have found $[a]_D = 55.5^{\circ}$ with a Jellet-Cornu instrument (Schmidt a. Hans, Ch.), sodium flame, and absolutely the same number $[a]_j = 61.6^\circ$ with a Soleil-Scheibler, taking 100 divs. = 38.4°.

 $(24:21.67::61.6^{\circ}:55.5^{\circ})$. $55.5^{\circ}-5.25$ p.c. = 52.59: my observations were made at 15.5°, Schmöger's at 20°. Freshly-prepared solutions of crystallised lactose exhibit the phenomenon of bi-rotation, the angle immediately after solution being in the proportion of 8:5 to the constant value (Urech, B. 16, 2270; Dubrunfaut, C. R. 42, 228; Schmöger, l.c.; Parcus a. Tollens, A. 257, 160). The rotatory power does not vary with the concentration. The dispersive power—that is, the ratio between the specific rotatory powers for the C and D lines—is 1:1·259 ([α]_o = 41·58) (Grimbert, J. Ph. [5] 16, 295 a. 345). The modification γ when freshly dissolved in water exhibits the phenomenon of semi-rotation—that is, the specific rotatory power is only about s of the normal, to which it attains on standing a few hours (E. O. Erdmann, Fortschritte der Physik, 1855, 18; B. 13, 2180; Schmöger, B. 13, 1915). The modification β on solution has immediately the normal optical activity; the presence of alkali lowers the

¹ Journal des Fabricants de Sucre,

activity (Schmöger, l.c.; Urech, B. 17, 1543). Schmöger give the specific gravity of solutions, containing given percentages of the crystals from which by calculation we get D = 3.956 for a solution containing 2.5 g. dry sugar in 100 c.c. at 20°, 3.921 for a 10 g. solution and 3.913 for a 15 g. one. My observations are, temperature 15.5°:—

1 gram in 100 c.c. sp.gr. $\frac{15^{*5}}{15^{*6}}$ 1·0040; D = 4·00 $\frac{5}{6}$ grams, , , 1·0199; D = 3·98 $\frac{10}{2}$, , , 1·0397; D = 3·97 $\frac{10}{2}$, , , 1·0785; D = 3·93

D=3.99 for solution below 10 g. per 100 c.c. at 15.5°. The K=79.2 for dry sugar, i.e. 0.5723 anhydrous lactose reduce 1 g. CuO from Fehling's solution (Jones, Analyst, 1889, 81). According to Soxhlet 0.5 g. sugar=74 c.c. Fehling's solution, hence K=74; in this case 1 g. CuO=6127 lactose: by weighing the precipitated CuO as Cu by Allihn's method, 1 g. sugar=1736 CuO, or 1 g. CuO=0.5760 g. sugar, K=78.8, agreeing with the number given by Jones. According to Roderwald a. Tollens (B. 11, 2076) the working number 1 g. CuO=5705 g. anhydrous sugar. I am inclined, when the conditions hereafter to be described are followed, to place the K at 78.9, i.e. 1 g. CuO=0.5758 g. sugar. In this case 7.5 mols. CuO are reduced by 1 mol. sugar.

Action of heat. Hydrous crystallised lactose (a) remains constant at 100°; above this water is slowly given off, and at 130° the dehydration is complete without decomposition if the water is eliminated slowly. Anhydrous lactose commences to colour at 170°-180°, loses water, and forms the so-called lactocaramel C.H.O., which is easily soluble in water but insoluble in alcohol, and yields compounds with lead and copper oxide. Lactose melts at 203.5° to a brown liquid, and by further heating it is decomposed with separation of carbon (Lieben, J. pr. 68, 409). Heated in a sealed tube with water to 90°-100° it is coloured, and more so at 180°-200°, when a glucose is produced which may be precipitated by alcohol, and, by further heating, carbonic acid, a little pyrocatechin, and other bodies are formed (Vohl, A. 105, 334; Hoppe-Seyler, B. 4, 16; Munk, H. 1, 357).

Action of acids. By heating with dilute

Action of acids. By heating with dilute acids lactose is hydrolysed (inverted), yielding galactose and dextrose

$$\begin{array}{c} {\bf C_{12}H_{22}O_{11}+H_2O=C_{e}H_{12}O_{e}+C_{e}H_{12}O_{e}} \\ {\bf Lactose} \end{array}$$

one molecule of water being taken up. The reaction takes some hours to complete; during the hydrolysis the reducing power and optical activity increase (Pasteur, C. R. 42, 228; Fudakowski, B.8, 559; 9, 42; 278 and 1602; 11, 1069; Bourquelot, N. Zischr. f. Rubenz-ind. 16, 71; Kent a. Tollens, A. 227, 221; Rindell, N. Zischr. f. Rubens-ind. 4, 163). The chief products of the reaction are as stated, but the conditions have not yet been described by which the optical activity and K of the resulting substances will exactly correspond with those of a mixture of equal parts of galactose and dextrose. Other bodies are undoubtedly produced. By the continued action of acids for several days, aceto-propionic and formic acids are produced (Tollens a. Boderwald, A. 206, 231; Conrad a. Guthzeit,

B. 19, 2575). Concentrated sulphuric acid does not blacken lactose in the cold.

Compounds with acids. A mixture of sulphuric and nitric acids give lactose pentanitrate C₁₂H₁₇(NO₂),O₁₁, insoluble in water but soluble in alcohol and ether, m.p. 139.2° and exploding when struck by a hammer on an anvil; lactose tri-nitrate C₁₂H₁₉(NO₂)₃O₁₁, very slightly soluble in water, but easily in alcohol and ether, m.p. 87°; and lactose tetranitrate $C_{12}H_{18}(NO_2)_4O_{11}$, m.p. 80°-81° (Gé, J. R. 1882, 253; B. 15, 2238). With chlorosulphonic acid lactose yields dextrosetetra-sulphonic acid chloride (Claësson, J. pr. [2] 20, 1 and 18). Organic acids act very slowly on lactose, but by prolonged action they yield ethers of lactose or of dextrose and galactose (Berthelot, Chim. org. 2, 279, 295). A series of acetates, beginning with the mono-body and ending with the octo-one, have been described. $C_{12}H_{14}(C_2H_8O)_8O_{11}$ is obtained by boiling lactose with an excess of acetic anhydride or by heating a mixture in the proper proportions of the sugar, acetic anhydride, and fused sodium acetate at 100° (Herzfeld, B. 13, 265). This body is insoluble in water and in ether, soluble in benzene, acetic acid, and in alcohol. It crystallises from a mixture of alcohol and acetic ether. $[a]_D = +31^\circ$ in alcoholic solution. The mono- and di- acetate are obtained by incompletely decomposing the octo-acetate by alkalis (Démole, C. R. 89, 481). Tetracetate is formed at the same time as the octo- acetate, when lactose is boiled with acetic anhydride (Schützenberger a. Naudin, Bl. 12, 208). It is easily soluble in water, $[a]_D = +50^\circ$. Hexacetate C₁₂H₁₆(C₂H₃O),O₁₁ crystallises from a mixture of alcohol and acetic ether in white needles, m.p. 86° (Herzfeld, N. Z. R. 8, 156). It is doubtful whether or not lactose is reproduced when these bodies are saponified by treatment with alkalis. There seems to be no reason why it should not be so, if the alkali is carefully employed.

Action of alkalis. Hot alkalis colour solutions of lactose yellow to brown, and form lactic acid, pyrocatechin, and other products (Hoppe-Seyler, B. 4, 347; Nencki a. Sieber, J. pr. [2] 34, 503; Uroch, B. 17, 1543). Fused with KHO it yields carbonic, oxalic, and a little succinic acids (Hlasiwetz a. Barth, A. 138, 76).

Compounds with alkalis. Lactoseates are formed by the action of potash or soda on lactose in alcoholic solutions, or by adding alcohol to the mixed aqueous solutions. They are amorphous, easily decomposed precipitates having a composition expressed by the formulæ C₁₂H₂₁NaO₁₁ and C₁₂H₂₁KO₁₁ (Fremy, A. 15, 278; Hönig a. Rosenfeld, B. 12, 45). Calcium, barium, and lead lactoseates are obtained by dissolving the base in lactose solution and precipitating with alcohol. The long-continued action of lime produces isosaccharin and metasaccharin (Cuisinier, M. S. [3] 12, 520; Kiliani, B. 16, 2625). By heating with ammonia, brown, amorphous, nitrogenous products are obtained (Thénard, C. R. 52, 444). Aniline yields two crystalline compounds, C_{so}H₄₉NO₂₁ and C_{so}H₅₄N₂O₂₀₁ which reduce Fehling's solution (Sachsse, B. 4, 835; L. V. 16, 441).

Oxidation.—Oxygen and ozone have no action on lactose in the cold; hot solutions are oxidised by oxygen in the presence of platinum black (Gorup-Besanez, A. 110, 86 a. 103; Reiset

a. Millon, A. Ch. [8] 8, 285). Alkaline permanganate solutions oxidise lactose easily (Laubenheimer, A. 164, 283). Chromic acid yields aldehyde (Guckelberger, A. 64, 98). Nitric acid first inverts lactose and then yields mucic and saccharic acids; from 36 to 40 p.c. of mucic acid is produced; by long-continued heating tartaric and oxalic acids, &c., are obtained (Liebig, A. 113, 1; Dubrunfaut, C. R. 42, 228; Kent a. Tollens, A. 227, 227). Alkaline solutions of copper oxide are reduced by lactose, and from amongst the products of the reaction Bödecker a. Struckmann (A. 100, 264) isolated gallactinic acid $C_{14}H_{10}O_0$; pectolactinic acid $C_{14}H_{10}O_{12}$. lactic and glycollic acids are also produced (Habermann a. Hönig, B. 17, 351). Silver oxide oxidises lactose solutions with production of oxalic, glycollic, and lactonic (galactonic) acids (Kiliani, B. 13, 2307). Chlorine or bromine in the presence of silver oxide yield lactonic (galactonic) acid C_eH₁₂O₇ (Barth a. Hlasiwetz, A. 119, 281; 122, 96). By very careful oxidation with bromine water Fischer a. Meyer (B. 22, 861) have obtained lactobionic acid C12H22O12; it is a colourless, strongly acid syrup, easily soluble in water, but only sparingly in alcohol. It does not reduce Fehling's solution; it decomposes metallic carbonates, forming salts. calcium $(C_{12}H_{21}O_{12})_2Ca$, barium $(C_{12}H_{21}O_{12})_2Ba$, lead $(C_{12}H_{21}O_{12})Pb$, and other salts may be prepared in this way; they are insoluble in alcohol, but soluble in water. When heated with dilute mineral acids, lactobionic acid is decomposed into galactose and gluconic acid. With iodine and sodium bicarbonate lactose yields a little

iodoform (Millon, C. R. 21, 828).

Fermentation.—There is no known enzyme capable of inverting lactose; it does not ferment with yeast (Berthelot, A. Ch. [3] 50, 332 a. 362; Fitz, B. 11, 42). It easily undergoes lactic acid fermentation, especially in milk (Fitz, L.c.; Richet, C. R. 86, 550; Schmidt-Mülheim, B. 15, 2631). It undergoes alcoholic fermentation in kumys or kefir fermentation (Struve, B. 17, 314; Vieth, A. 12, 2), and also under the agency of certain bacteria (Rotondi a. Zechini, see v. Lippmann, Deutsche Zuckerind. 1887, 1091).

Phenylhydrazine yields lactose phenylhydrazide $C_{18}H_{28}O_{10}N_2$, soluble in water and alcohol, insoluble in ether and lævorotatory, and lactose osazone $C_{24}H_{22}N_4O_9$, crystallising in yellow needles; soluble to some extent in hot water, and m.p. 200° (Fischer, B. 17, 583; 20, 830; 20, 2566). By the action of hydrochloric acid it yields lactose osone (Fischer, B. 21, 2631).

Maltobiose, Amylon, C12H22O11; Maltose. C₁₂H₂₂O₁₁.H₂O. De Saussure (Bibl. Britannique, 56, 1814, 333; P. 1819, 29, 58) may have obtained this sugar when studying the products of the spontaneous decomposition of starch-paste when exposed to the air at 20°-25°. Guérin-Varry (A. Ch. 60, 32; 61, 66) described accurately how the body could be prepared in the crystal-lised state and some of its properties, but failed to recognise it as a distinct substance; Jacquelain (A. Ch. 63, 167) added nothing to Varry's work; Dubrunfaut (A. Ch. [3] 21, 78) prepared the body according to Varry's method, observed that it was less soluble in alcohol than dextrose, and had an optical activity three times as great, i.e. $[a]_1 = 58 \times 8 = 174$, a figure so high as to lead

one to believe that he had not a pure body in his hands. He recognised it as a distinct body, and called it maltose; he, however, looked upon it as a glucose, triglucose; O'Sullivan (C. J. 1872, 576; 1876, 478) showed that it was not an 'ose' but an 'on' sugar, and from this and a fuller description of its characters gave it a place among chemical entities.

Occurrence.—It is present in some commercial glucoses and in beer (Valentin, J. S. A. 24, 404); probably in bread; in amé, a rice-extract prepared in Japan (Yoshida, C. N. 43, 29), and in germinated cereals to the extent of 1 to 2 p.c. (O'Sullivan, C. J. 49, 58). After feeding with amylaceous substances it is found in the intestinal canal, but the blood of the mesenteries contains only dextrose. If injected into a vein it is found in the urine; if injected subcutaneously, it is partially converted into dextrose (Phillips, B. C. 1882, 127). Some of the reducing sugars of blood may be maltose.

Formation.—Diastase converts starch and the dextrins into maltose (O'Sullivan, L.c.). The ferment of saliva, ptyalin, of the pancreas, and of the liver, act in the same way (Nasse, J. Th. 1877, 62; Musculus a. v. Mering, Z. 2, 403). The same ferments act on glycogen, maltose being amongst the products; dextrose is present when saliva is employed (Kütz, Pf. 24, 8). All mineral acids and many organic acids act on starch and glycogen; maltose is amongst the products.

Preparation.-100 g. purified starch are mixed as completely as possible with 300 c.c. water at 40°, and then poured with continuous stirring into 2 litres of boiling water. The paste is cooled to 60° and the extract from 20 g. pale malt added to it; the mixture is kept at 60°-63° for four or five hours, and the solution may be allowed to cool and stand for a few days. At the end of that time it is evaporated, best in a vacuum, to 200 c.c., and then boiled with 2 litres alcohol, sp.gr. 820. On cooling and standing, the clear liquid is decanted off the undissolved syrup and put aside in a corked flask; at the end of six days the sides of the flask will be found to be covered with a crystalline crust of maltose. Or the undissolved syrup may be treated with strong alcohol as long as any matter is taken up, the whole of the clear alcoholic solutions are mixed, the alcohol distilled off and the residue concentrated to a syrup; this on standing in a cool place becomes filled with crystals of maltose, especially if a few crystals from a previous preparation be stirred in. These may be washed with dry methyl alcohol; if this is used hot, a portion of the crystals dissolves and the solution soon yields much sugar. The crystallisation is hastened by adding half a volume of ethyl alcohol (.810 sp.gr.) or a little ether with some crystals of a previous preparation (O'Sullivan, l.c.; Schulze, B. 7, 1047; Soxhlet, J. pr. [2] 21, 277; Herzfeld, A. Ch. 220, 209; Cuisinier, J. 1884, 1803). The body may be purified by recrystallisation from methyl alcohol or water.

Properties.—Maltose crystallises out of water in plates which contain $O_{12}H_{21}O_{11}$. $H_{2}O$; out of alcohol (sp. gr. \circ 810) it crystallises in crusts and at times in cauliflower-like or warty aggregations which contain $O_{12}H_{22}O_{11}$. The hydrous body easily parts with its water at 100° in a

current of dry air. The anhydrous body is but slightly soluble in strong ethyl alcohol; it is more soluble in methyl alcohol. The hydrous body dissolves more freely in both solvents. Both bodies are very soluble in water, but less so than dextrose. The solution is optically active, and the activity for a solution containing 10 g. in 100 c.c. is $15^{\circ}-20^{\circ}$ less when freshly prepared than after standing. The activity becomes constant in ten to twelve hours in the cold or immediately on boiling (Meissel, J. pr. [2] 25, 120).

The specific rotatory power for dry maltose is $[a]_1 = 154^\circ - 155^\circ$, $[a]_D = 139^\circ - 140^\circ$ at $15\cdot 5^\circ$ in solutions containing 10 g. per 100 c.c. and under (O'Sullivan, C. J. 85, 771; 45, 5); Meissel (J. pr. [2] 25, 114) gives the formula $[a]_D = 140\cdot 735 - 01837 P - 095 T$ to represent the specific rotatory power of dry maltose at 0°, in which P = percentage of maltose in solution and T the temperature of observation. On calculating the value of the sugar in a 10-g. solution at 15·5° from this formula we get $[a]_D = 138\cdot 9 \ (v. \text{ also Soxhlet}, l.c., \text{ and Herzfeld, } A. 220, 206)$. The dispersive power, i.s. the ratio of $[a]_D : [a]_C$ is 1·262 (Grimbert, J. Ph. [5] 16, 295). A solution of maltose containing 10 g. dry substance in 100 c.c. at 15·5° has a sp. gr. $\frac{15\cdot5^\circ}{1550} =$

1.0395. Maltose reduces alkaline copper solution. O'Sullivan (C. J. 35, 771) gives its reducing power K=62.5, i.e. 62.5 parts dextrose reduce as much copper oxide as 100 parts of Soxhlet (J. pr. 21, 227-317) says maltose. 0.5 g. maltose in 1 p.c. solution = 64.2 c.c. undiluted Fehling solution (1 c.c. = 005 g. dextrose), i.e. K = 64.2, and 67.5 c.c. if the solution is diluted with four times its volume of water. i.e. K=67.5. He further states that 100 parts of dry maltose = 113 parts Cu, hence 141 parts CuO; from this, we have K = 63.94. There appears to be fair evidence that the K varies within certain limits according to the mode of manipulation, but when we see that the K should be 63.17 if it be supposed that 1 molecule maltose reduces 6 molecules CuO, and that the observed numbers closely approach this, we may take it that this is the true number when errors of manipulation are avoided. If the Cu2O be separated, the filtrate treated with acid gives a further reduction with Fehling's solution which, if added to the first, equals the reduction of dextrose (Herzfeld, A. B. 589, 220). Alkaline mercuric cyanide solutions are also reduced by maltose. Copper acetate is not reduced by it (Barfoed, Org. Analysis, 214); dextrose reduces this reagent. When maltose reduces copper oxide, glycollic and a mixture of acids of unknown composition are produced (Habermann a. Hönig, M. 5, 208).

Maltose is hydrolysed to dextrose according to the equation $C_{12}H_{22}O_{11} + H_1O = 2C_6H_{12}O_6$ by the following reagents: the mineral acids and many organic ones; pancreatic secretion and portions of the small intestine (Brown a. Heron, C. N. 42, 63; Bourquelot, C. R. 97, 1000 and 1822); a ferment developed in Aspergillus niger and in Mucor mucedo; the former contains also a ferment capable of inverting sucrose, the latter yields only the one (Bourquelot, l.c.). Under the action of ordinary beer yearts malt-

ose ferments, i.e. is broken down into carbonic acid, alcohol, &c., apparently without being previously inverted. Other organisms yield lactic acid and other products; v. Fermentation.

Maltose withstands the hydrolytic action of sulphuric acid with five times the power of sucrose; 3 hours' digestion with a 3 p.c. solution gives complete inversion (Meissel, l.c.). 0.5 p.c. solution of lactic acid does not hydrolyse it, even at 110°; 1 p.c. oxalic acid acts at that temperature; carbonic acid at 100°, and under a pressure of six atmospheres, is without action (Bourquelot, J. de l'Anat. et de la Physiol. 22, 161-204; Urech, B. 18, 3074).

Nitric acid converts maltose into sacchario acid and finally oxalic acid. Chlorine does not act on it as energetically as it does on dextrose or sucrose; the product is neither gluconic nor glycollic acids (Meissel, l.c). Herzfeld (B. C. 1883) obtained an acid which he called maltonic acid, by the action of bromine on maltose; this is probably identical with Fischer and Meyer's (B. 22, 194) maltobionic acid C₁₂H₂₂O₁₂, which they prepared by acting on maltose in aqueous solution with bromine in the cold for two or three days. This acid is broken down into dextrose and gluconic acid by digestion with sulphuric Treated at 110° with acetic anhydride and acetic acid, maltose yields monacetyl maltose $C_{12}H_{21}(C_2H_2O)O_{11}$ (Yoshida, l.c., a. Steiner, C. N. 43, 52); with sodium acetate and acetic anhydride the octacetyl derivative C12H14(C2H2O)8O11 is produced, the optical activity of which is [a]_p = 81·18° (Herzfeld, A. 200, 206). With benzyl chloride it yields penta-benzyl maltose, m.p. 110°-115° (Skraup, M. 15, 359).

Sodium $C_{12}H_{21}NaO_{11}$, calcium $C_{12}H_{20}CaO_{11}H_1O$, strontium $C_{12}H_{20}SrO_{11}H_2O$, and barium $C_{12}H_{20}BaO_{11}.H_2O$ maltose are prepared as easily decomposable amorphous precipitates when alcoholic solutions of maltose and the corresponding bases are mixed (Herzfeld, *l.c.*). Compounds with the alkaline chlorides or bromides have not been prepared.

When 1 part maltose, 2 parts phenylhydrazine hydrochloride, 3 parts sodium acetate, and 15 parts of water are heated together for 1½ hours, phenylmaltosazone crystallises out on cooling in fine yellow needles, m.p. 190°–191°. The equation C₁₂H₂₂O₁₁ + 2N.H₂.C₂H₃ = C₂₄H₃.N₄O₅ + 2H₂O + H₂ represents the reaction. Only 30 p.c. of the maltose employed is obtained as osazone (Fischer, B. 17, 583). With y-diamido-benzoic acid, maltose yields malto-y-diamido-benzoic acid according to the equation

The body crystallises in white microscopic needles or narrow plates; its barium salt is a gummy mass (Griess a. Harrow, B. 20, 2712).

Isomaltose. Fischer (B. 23, 3687) prep ared a saccharon by the polymerisation of dextrose. Previous attempts in this direction had been made by Musculus (Bl. 18, 66), Muscu us a. Meyer (C. R. 92, 528), Hönig a. Schuber M. 7, 455), and they obtained by the action of strong sulphuric acid a dextrin-like body C.H. 90s.

which slightly reduced Fehling's solution, did not ferment with beer yeast, and was reconverted into dextrose by the action of dilute sulphuric acid. Grimaux a. Lefèvre (C. R. 103, 146) obtained a like result by evaporating in vacuo dextrose with dilute hydrochloric acid. Gautier (Bl. 22, 145) obtained a body C₁₂H₂₂O₁₁ by the action of hydrochloric acid on an alcoholic solution of dextrose; this body reduces Fehling's solution but slightly, does not ferment with beer yeast, and could not be reconverted into dextrose. Grimaux and Lefèvre considered that in their product they could detect maltose by the osazone. Scheibler a. Mittelmeier (B. 23, 3075; 24, 301) isolated an unfermentable syrup from commercial glucose which reduces Fehling's solution and yields an osazone $C_{2,1}H_{2,2}N_{1}O_{0}$, m.p. $152^{\circ}-153^{\circ}$. They find it is only produced when the heating with acid has been prolonged, and that it may be prepared by the action of acids on dextrose. This body has been previously named gallisin (Cobenzl, Roseneck a. Schmitt, B. 17, 1000 and 2456; Anthon, D. P. J. 151, 213; Mehring, Deutsche Vierteljahrsschrift für öffentl. Gesundheitspflege, 14, Heft 2). Fischer's substance also yielded the same osazone, m.p. 150°-153°; he calls the body isomaltose, and prepares it as follows: One hundred g. glucose (commercial dextrose) are digested with 400 g. hydrochloric acid solution sp. gr. 1.19 at 10°-15° for 15 hours. Alcohol throws out of the solution a small precipitate, which is not formed if the temperature of the reaction is kept below 10°. To the clear solution an excess of ether is added, when a colourless amorphous precipitate is obtained which is filtered out and washed with a mixture of alcohol and ether. This body is dissolved in water, neutralised, boiled to expel alcohol and ether, and fresh beer yeast added. After 18 hours all the dextrose is destroyed (?); the solution has still a high reducing power, and contains isomaltose and other bodies.

The isomaltose is separated as osazone; the solution is heated with phenylhydrazine acetate in the water-bath for 11 hours. A little glucosazone separates and is filtered out, and on cooling the filtrate deposits a yellow precipitate consisting of a mixture of glucosazone and isomaltosazone; the mother-liquor, by further heating and similar after-treatment, yields more osazone. The whole of the osazone obtainable is boiled with 100 c.c. water and the solution filtered, the filtrate contains isomaltosazone, which is deposited on cooling in needles; 2 g. of this are obtained from 100 g. glucose, the actual yield is, no doubt, considerably higher than this. By the action of hydrochloric acid the osazone is converted into the osone, which, when boiled with 4 p.c. hydrochloric acid, yields glucosone and glucose.

Lintner claims to have found isomaltose amongst the transformation products of starch by diastase (Woch. Brauerei, 9, 245), and in wort and beer (Zeit. ges. Brauw. 1891, 281), but it is very doubtful if this is the same body as that above described. Schiffer (C. C. 1892, 2, 825) has also isolated isomaltose from the diastase transformation products of starch. But the whole evidence is unsatisfactory, as the factors upon which the identity of the bodies can be recognised are not stated. Lintner and Düll (Zeit. ges. Brauw. 1892, 145) prepare it as fol-

lows: 250 g. potato starch are mixed with 500 c.c. diastase solution at 55° containing 0.5 g. diastase (J. pr. 34, 378) and 2 litres water at 75°C. After complete solution another 0.5 g. diastase is added, and the reaction allowed to proceed for three hours at 65° – 69° . [a]_D = 170°. for matter in solution. The solution is evaporated to a syrup, saturated with 80 p.c. alcohol, and poured into hot alcohol. Sufficient alcohol must be used so that every 100 parts of 80 p.c. alcohol do not contain more than 10 pts. dry substance. After cooling, the clear solution is decanted and the alcohol distilled off. The residue is diluted to a 20 p.c. solution and yeast added in order to destroy the maltose; this is found to take place in about 20 hours. The fermented solution is filtered, decolourised with animal charcoal, evaporated to a syrup, and precipitated with 85 p.c. alcohol; there must be 100 c.c. of 85 p.c. alcohol for every 5 g. of dry substance. The solution is evaporated and the syrup treated in the same way, but with 99 p.c. alcohol and so that 100 c.c. of alcohol are present for every 3 g. dry substance. The clear alcoholic solution contains isomaltose and but a trace of dextrin, which may be removed by fractionating with alcohol. In this way 20 p.c. of the starch is obtained as isomaltose.

Lintner (Zeit. ges. Brauw. 1892, 6) finds the reducing power of iso-maltose to be 84 p.c. of that of maltose, i.e. K = 53.1, and its optical activity to be $[a]_D = 139^\circ$. He has not been able to crystallise it, and finds it under certain conditions fermentable by yeast and converted by diastase into maltose (Zeit. ang. Ch. 1892, 263).

Trehalose. $Mycose C_{12}H_{22}O_{11}.H_{2}O$.

Occurrence.—It has been isolated from ergot of rye (Wiggers, A. 1, 173; Mitscherlich, 56, 15); trehala-manna, the hollow cocoons of the larve of an insect (Larinus maculatus) (Berthelot A. Ch. [3] 53, 232; 55, 272 and 291); and from various fungi (Müntz, C. R. 76, 649). Berthelot at first considered it a new sugar and called it trehalose, but afterwards found it was identical with the mycose of Mitscherlich, and the Saccharum spermodiæ of Wiggers.

Preparation.—Any one of the materials mentioned is extracted with boiling alcohol of moderate strength; the alcohol is distilled off the clear solution. The residue is taken up with water and precipitated with basic lead acetate. The precipitate is washed with water and pressed; it is then suspended in water and submitted to a current of H.S. The clear filtrate from the lead sulphide is evaporated to a syrup, when the sugar crystallises and can be purified by recrystallisation. I may say that I do not find this mode of treatment answer very well; the filtrate from the PbS is, no doubt, clear, but the precipitate with the lead salt is bulky and unwieldy. I much prefer in this case, as in others of the same kind, fractional precipitation with alcohol; a few experiments will show the strength of alcohol best suited to holding or precipitating the sugar. (See also Apping.)

Properties.—The sugar crystallises in rhombs (C₁₂H₂₂O₁₁H₂O); it gives up its water at 100°, m.p. 109°; it is sweet, easily soluble in water and boiling alcohol, insoluble in ether. Specific rotatory power in aqueous solutions.

[a]_b = 199° (Berthelot); [a]_b = 192.5° (Mitscherlich); [a]_b = 197.28° (Apping). These factors are unsatisfactory, and cannot be relied on to identify the sugar. It does not reduce Fehling's solution. Acids convert it very slowly into dextrose (Berthelot), there being no other product; but considering that five hours are required for complete hydrolysis, it is not possible that only pure dextrose could be formed. With strong nitric acid it yields a nitro- compound; with weaker acid no mucic acid is formed, but saccharic acid and finally oxalic acid. With acetic and butyric anhydrides it yields compounds identical with those produced from dextrose. Maquenne (C. R. 112, 947) says acetic anhydride yields $C_{12}H_{14}(C_2H_3O)_8O_{11}$. Alkalis do not act on it. On adding basic lead acetate, or an ammoniscal solution of the acetate, to its solutions a lead compound is precipitated. It appears to slowly undergo alcoholic fermentation in contact with beer yeast, but this has not been established with certainty. Phenylhydrazine does not form a compound with trehalose (Fischer, B. 17, 583).

Agavose C₁₂H₂₂O₁₁ (?) is obtained from the juice of Agava americana; it is crystalline, soluble in water, and optically inactive (?). It is hydrolysed to a sugar or mixture of sugars $[a]_i = -14.4^{\circ}$. It reduces Fehling's solution; K=62.5, agreeing nearly with maltose in this respect. It does not yield mucic acid on treatment with nitric acid. It is probably fermentable under the influence of some saccharomyces and yields alcohol, &c. (Michaud a. Tristan, Am.

14, 548).

Cyclamose $C_{12}H_{22}O_{11}$. A sugar obtained from Cyclamen europæum, having a lævorotation $[a]_D = -15\cdot15^\circ$ or $-11\cdot4^\circ$; it is hydrolysed by dilute acids, when the lævorotation increases to $[a]_D = -66.54^{\circ}$ (Michaud, C. N. 53, 232).

Parasaccharose C₁₂H₂₂O₁₁. This, together with another uncrystallisable sugar, is said to be obtained from cane sugar by the action of a yeast (Jodin, C. R. 53, 1252). It crystallises easily, and is easily soluble in water and difficultly in alcohol. It is dextrorotatory, having [a] = 108°, and it reduces Fehling's solution; K = 50 (about). Hot dilute sulphuric acid does not act on it, whilst hot dilute hydrochloric acid diminishes the optical activity and increases the K. On long digestion the solution becomes brown, and finally complete decomposition takes place

Melibiose C12H22O11 is obtained by the careful hydrolysis of raffinose by acids or invertase (see Raffinose). Its specific rotatory power is $[a]_D = +126.8^\circ$. It forms a hydrazide $C_{10}H_{20}O_{10}N$ as yellow microscopic crystals, m.p. 1465, and an octacetyl derivative C₁₂H₁₄O₁₁Ac₂ (Scheibler a. Mittelmeier, B. 23, 1438). Invertase converts

it into galactose and dextrose.

Tewfikose C12H22O11 occurs in the milk of the gamoose (Bos Bubalus). It is prepared by precipitating the milk by 4 p.c. acid mercuric nitrate (Wiley, Am. 6, No. 5 (?), neutralising the filtrate with NaHO, filtering out the precipitate, separating the mercury from filtrate by H.S., and concentrating the slightly acid solution to the crystallising point. A slight precipitate formed during evaporation must be separated. It is purified by two or three recrystallisations. The solutions are dextrorotatory, $[a]_D = 48.6^\circ$; it

reduces Fehling's solution; K=73.6, hence less than lactose; if 7 molecules CuO were reduced by 1 molecule of sugar, the K would be 73.4, a near approximation; and D = 3.94 for the crystals, hence they are anhydrous. Acids hydrolyse it very easily, the only product being dextrose (Pappel a. Richmond, C. J. 1890, 758). The substance is, no doubt, an '-on' sugar C₁₂H₂₂O₁₁, and is, like amylon, di-dextrose, and should not therefore yield mucic acid on treatment with nitric acid.

(c) Trihexoses.

Baffinose C₁₈H₃₂O₁₆5H₂O, first obtained from a Tasmanian Eucalyptus manna by Johnston (C. J. 1, 159). This was the melitose of Berthelot (A. Ch. [8] 46, 66), who now says (C. R. 103, 533) that this term should be applied only to a loose combination of raffinose with eucalinwhich is decomposed by re-crystallisation. From the evidence it would appear that the body first described as melitose was impure raffinose, and that the impurity even was not eucalin. It occurs also in cotton seed (Berthelot, l.c.; Ritthausen, J. pr. [2] 29, 357; Böhn, J. pr. [2] 30, 37); in beetroot (Loiseau, C. R. 82, 1058; Tollens, A. 232, 169; B. 18, 26; Rischbieth a. Tollens, A. 232, 172; B. 18, 2611); in beet juice (v. Lippmann, B. 18, 3087); in barley (O'Sullivan, C. J. 1886, 70); in young wheat-buds (Richardson a. Crampton (B. 19, 1180); and again in Eucalyptus manna (Tollens, A. 232, 201); probably also in Soja beans. Loiseau considered it a product of the refining process, but v. Lippmann that it was ready formed in beet juice.

Preparation. - A molasses containing a fair quantity of raffinose is selected, and the chief part of the cane sugar removed as monostrontium saccharate; on heating the solution raffinose and cane sugar bistrontium saccharates are The compounds are decomposed precipitated. by CO₂ and the sugar solution obtained again precipitated with strontia, when the cane sugar is precipitated as monostrontium saccharate, and rattinose remains in solution and may be purified by crystallisation (Scheibler, B. 18, 1409). Raffinose may be extracted from cottonseed cake and Eucalyptus manna by alcohol, and purified by crystallisation. See also Tollens (l.c.); Tenne (Zeit. d. Ver. 31, 795); Schaaf (Z. V. 33, 699); Wolff (Deutsche Zuckerind., 1887, 1413); Berthelot (l.c.); Lindet (C. R. 110, 795); Gun-

ning (C. C. 1891, ii. 798).

Properties.—Raffinose crystallises in cauliflower-like segregations consisting of thin microscopic rhombic prisms, which contain The H₂O is in greater part C₁₈H₃₂O₁₆.5H₂O. eliminated in a vacuum over sulphuric acid, and the remainder easily at 100° (O'Sullivan). heated too quickly it melts, and then cannot be dried without decomposition. Berthelot obtained crystals containing 6H,O (C. R. 109, 548; Scheibler, B. 18, 1779; Rinne in Rischbieth Dissert. über Raffinose, Göttingen, 1885). It is easily soluble in water, very slightly in strong alcohol, and slightly in methyl alcohol, being much more soluble than cane sugar in this solvent. The dry sugar melts at 118°-119° (Scheibler, B. 19, 2868). Its solutions are optically active, the specific rotatory power of: crystallised raffinose for the D line being [a]_D = 104.5° in 10 p.c. solution; [a]; = 114.7° or [a]; =

135·1° for the dry sugar. O'Sullivan observed $[a]_j = 185·3°$. No bi-rotation has been observed. It does not reduce Fehling's solution. D = 3·712 (Tollens, B. 18, 2616), but this is due to some arisunderstanding; D = 3·956 (O'Sullivan, C.J. 1886, 70).

Action of acids. Hot dilute acids hydrolyse raffinose; galactose, lævulose, and probably dextrose are formed (Hädicke a. Tollens, Z. V. 37, 17; A. 238, 808; Beythien a. Tollens, A. 255, 214; Maquenne, C. R. 112, 799). By very careful hydrolysis, Scheibler a. Mittelmeier (B. 22,1678) have obtained a saccharose and lævulose as the products of the reaction; the saccharose is a new body, which they have called melibiose (q. v.). The mixture of saccharose and lævulose has a specific rotatory power $[\alpha]_D = 50^\circ$. Complete inversion of raffinose takes some hours. vertase also hydrolyses raffinose (O'Sullivan). The products of hydrolysis are at first lævulose and melibiose (Scheibler a. Mittelmeier, B. 22, 8118), and melibiose is slowly acted on. When heated for some time with sulphuric acid, lævulinic acid is a product (Rischbieth a. Tollens, A. 232, 195).

Action of alkalis. By boiling aqueous solutions of strontia and raffinose, di-strontia raffinose $C_{15}H_{22}O_{16}(SrO)_2.H_2O$ is produced. Other compounds with baryta $C_{15}H_{22}O_{16}.BaO$, lime $C_{15}H_{22}O_{16}.SCaO.2H_2O$, lead oxide $C_{15}H_{32}O_{16}.SPbO$, and soda, $C_{15}H_{31}NaO_{16}$ and $C_{15}H_{31}NaO_{16}.NaOH$, have been prepared (Beythien a. Tollens, B. 22, 1047; A. 255, 195; Rischbieth a. Tollens, A. 232, 172).

Oxidation.—Nitric acid oxidises raffinose, producing 30 p.c. mucic acid, also saccharic and

oxalic acids.

Fermentation.—Raffinose is slowly fermented by beer yeast (Rischbieth a. Tollens, A. 232, 242; Tollens, A. 232, 169; O'Sullivan, C. J. 1886. 73). Weak yeast only partially ferments (Tollens, I.c.; Berthelot, C. R. 109, 548); the melibiose being with difficulty hydrolysed and fermented.

With phenylhydrazine a compound is formed, m.p. 187°-189° (Rischbieth a. Tollens, l.c.).

Melezitose $C_{18}H_{32}O_{18}.2H_{2}O$ is obtained from Persian manna, a product of Alhagi maurorum (DC.), it crystallises in rhombic prisms, m.p. 147–148° (when anhydrous); the specific rotatory power of the anhydrous body is $[a]_D = 87.7^\circ$. On inversion it yields turanose and dextrose. Turanose $C_{12}H_{22}O_{11}$ $[a]_D = 65^\circ-68^\circ$, yields dextrose by the further action of acid (Alekhin, J. R. 21, 407).

(d) Hexahexoses.

Gentianose $C_{sc}H_{e2}O_{sp}$. Obtained by A. Meyer (H. 6, 185), from the root of Gentiana lutea, by extraction with alcohol, precipitation with ether and re-crystallisation from alcohol. It has a sweet taste, dissolves easily in water, m.p. 210°, does not reduce Fehling's solution, and terments with yeast. It is hydrolysed by dilute acids; before inversion the optical activity is $[a]_D = 38.6^\circ$; after, it is 20.2° , and has a reducing power equal to that of dextrose. Strong sulphuric acid chars it, like cane sugar.

Stachyose is obtained from the root of Stachys tuberifera. It has a sweet taste, is dextrorotatory, having [a]_D = 148·1° in 9 p.c. solution. It forms triclinic crystals. When in-

verted it yields galactose, dextrose, and lævulose. Its formula appears to be $C_{22}H_{21}O_{22}+6H_2O$ (Planta a. Schulze, B. 23, 1692; 24, 2705), the products of the hydrolysis being the same as those of raffinose.

QUALITATIVE AND QUANTITATIVE DETERMINATION OF SUGARS.

If the material to be examined is a solution, it must be made neutral, clear, and colourless if it is not already so.

If the material is a dry substance, it must be ground fine. If not sufficiently dry to admit of grinding, it must, if possible, be dried in vacuum over sulphuric acid, or, where this is not possible, in the open, at a low, gradually increasing temperature.

The finely-ground substance is extracted with alcohol sp. gr. 860-880. The solution is neutralised and the alcohol distilled off. The residue is taken up with water. The examination then comes under the head of a solution, and the treatment is the same.

The solution is turbid and coloured after neutralisation. The turbidity may be removed by simple filtration; if not, a little aluminium hydroxide (prepared by precipitating a solution of alum with excess of ammonia and washing free from ammonia and sulphuric acid) may be employed. A little (well washed) Swedish filter paper-pulp is at times found useful. The turbidity and colour may be removed together by animal charcoal, lead sulphide, barium sulphate, &c. Animal charcoal, either prepared from blood or bones, the former by preference, thoroughly extracted with hydrochloric acid and washed free therefrom, is best employed in all cases of investigation. The crude substances are employed to remove the colour, &c., from sugar syrups in the manufacture of refined cane sugar. These decolouring agents must always be used in the least possible quantity, and with the understanding that the charcoal especially is capable of taking up some sugar as well as colouring matter. Dried, freshly-made bone charcoal does not take up more than 006 p.c. of its own weight from pure cane-sugar solution (Casamajor, C. N. 41, 66). In quantitative work, when the decolourising agents have to be used it is well to have a knowledge of the influence of the agent on the sugar or sugars under exami-Charcoal retains sugar, other subnation. stances influence the optical activity. Part of the sugar is again given up on washing the agent with water, or the agent is placed on a filter and a portion of the solution to be examined passed through it and rejected before the portion intended for examination is collected. This is done on the supposition that the charcoal is sooner saturated with sugar than with colouring matter.

Having thus the sugars in clear solution, the means at our disposal for identification are:—

1. Taste of solution.

Form of crystals and melting-point thereof.
 Depression of the freezing-point (Raoult's method of determining molecular weights).

Specific rotatory power.

5. Reducing power.

6. Action of unorganised ferments: ensymes.

7. Action of organised ferments.

- 8. Action of acids.
- 9. Action of alkalis.
- Action of phenylhydrazine, and meltingpoints of the resulting osazones and hydrazides.
- The solution tastes sweet if a sugar is present. It must not be forgotten, however, that some other bodies are also more or less sweet.
- 2. On evaporating this solution (best in a vacuum) to a syrup, if there is no preventive material present, crystallisation takes place. The microscopic appearance of the crystals will often be sufficient to identify the sugar; indeed, the habit of crystallisation will often suffice. substance or substances may be present which altogether prevent the crystallisation of the syrup, although it may contain a crystallisable sugar. This substance may be an inorganic or organic salt, or another body, or, indeed, another sugar. Treatment with strong or absolute ethylic or methylic alcohols may at times eliminate some or the whole of the substances preventing the crystallisation, and dissolve the sugar in so clean a state as to admit of the formation of crystals on concentration.

3. Depression of the freezing-point (Raoult's method). For references to Raoult's work see Victor Meyer, B. 21, 536; Auwers, id. 701; Tollens a. Meyer, id. 1566; and Brown a. Morris, C. J. 1888, 610. When crystals are obtainable from the solution as described above, a determination of the molecular weight by this method will decide to which group it belongs: a hexose or a pentose, a dihexose or dipentose, a trihexose. In the recorded observations water of crystallisation appears not to have received proper attention. Tollens a. Meyer record observations for dextrose C₆H₁₂O₆, so do Brown a. Morris. Water of crystallisation is not taken into account; the former treat of $C_{12}H_{22}O_{11}$, $H_{2}O$ for lactose, the latter omit the $H_{2}O$, while both employ crystallised raffinose C₁₈H₂₂O₁₈.5H₂O, mol. = 594, with the result that T. a. M. find the observed molecular weight between 544 and 644, and B. a. M. between 518 and 533. In these experiments with raffinose and lactose, it is not possible to say whether it is intended that the crystallisation water functions with the solid sugar or with the water; the calculations make it function with the sugar, but the results, as might have been expected, are unsatisfactory—it does not function with the sugar. It is perfectly clear that water of crystallisation of a sugar must be looked upon as water in observations with this method—that the quantity of dry sugar should be the factor dealt with. If the crystals contain water of crystallisation it should be determined and allowed for. A determination of the D will give some information as to whether a dry sugar or one containing water of crystallisation is being dealt with. The D of the different sugars in the dry state is given under each sugar. Between 8 and 15 g. dry substance is dissolved in water and the solution made up to 100 c.c. at 15.5°. A determination of the specific gravity of this will give a means of calculating the D, and the number of g. of water to substance employed in the solution. About 60 c.c. of this solution are introduced into a thin glass (120 c.c.) beaker. The

mouth of the beaker is loosely closed with an indiarubber plug, in which three holes are bored. One of these, in the middle, admits of the introduction of a thermometer; the second, at the side, carries a stirrer, of very thin glass rod or platinum, best in the form of a flat coil, so that when the stirrer is moved up and down in the liquid the thermometer will be in the middle of the coil; and the third, also at the side, to admit of the introduction of a small solid particle of the frozen solution or of ice. The thermometer must be graduated for 2° or 3° above and below 0° to the 20th of a degree. By means of a telescope, if the divisions are open, and they should be so, readings to .005° can be made with a fair amount of accuracy. The beaker thus charged and fitted is placed in a freezing mixture of ice and salt—which should not be at too low a temperature, -4° to -5° , not more than $2^{\circ}-3^{\circ}$ below the freezing-point of the solution—contained in a strong beaker of 500-600 c.c. capacity. In a short time the temperature in the inner beaker will be observed to sink below 0°, and, on rapidly moving the stirring rod up and down, to go gradually lower and lower until a point is reached when the solution begins to freeze. This may be accelerated by adding a particle of the solution previously frozen; when freezing begins the mercury of the thermometer rapidly rises and soon becomes stationary. This is the freezing-point of the solution c, and is the observation upon which the molecular weight of the substance employed is calculated.

$$M = \frac{19}{A}, \qquad A = \frac{c \times y}{x \times 100},$$

x=g. substance; y=g. water, and 19 is a number calculated by Raoult for water as a solvent from observations with well-defined compounds. As examples, Brown a. Morris (*l.c.*) with a solution containing 8.258 g. sucrose and 94.93 g. water observed a freezing-point $c=0.490^\circ$, thence

19

$$M = \frac{.49 \times 94 \cdot 93}{8 \cdot 258 \times 100} = 337 \cdot 5$$
. $C_{12}H_{22}O_{11} = 342$.

Hence, if an unknown sugar were under observation there would be no doubt left as to the molecular weight. O'Sullivan (C.J. 1890, 59) observed for arabinon: sp.gr. sol. 1.02554, $y = 96.088 \text{ g., } x = 6.466 \text{ g., } c = 0.535^{\circ}$; thence

$$19 + \frac{96.088 \times .535}{6.466 \times 100} = 239.2,$$

the molecular weight of the new sugar. dipentose requires 282. The indications agreed with further confirmatory observations. When the solution contains more than 12 g. per 100 c.c. the results are low; with most sugars a solution containing from 5 to 8 g. substance per 100 c.c. gives the best results, and with more dilute solutions the results are again low. Raoult's latest observations (C. R. 114, 264) for sucrose, made with apparatus constructed to meet the exact requirements of the case, the concentration of the solution being 5.839 g. sucrose in 100 g. water, work out M = 323, against 342, the true molecular weight. The results are, even under the best conditions, low; but the indication is sufficiently good to enable us to decide to which group the sugar we are dealing with belongs. If, instead of the factor 19, 19.8 were substituted, the results would agree with the accepted molecular

weight of sucrose, and it is probable that this number or 20 will be finally decided upon as the constant.

4. Specific rotatory power. If the form of the crystals or their habitat does not indicate the sugar present, a determination of the specific rotatory power may give the desired information. This is the quantitative expression of a property possessed by the sugars in common with many other substances. When light from any source is examined through a Nicol prism—i.e. polarised—no change is evident; but if the light, after passing through one prism (the polariser) is examined by another, it is found that in certain positions of the examining prism or analyser, as it is called—no light passes. If the analyser is made to carry an index, and is so mounted as to rotate with its axis at right angles to a disc graduated to degrees, it will be found that, if the position of the two prisms is so arranged that the index is at 0° on the graduated circle when no light passes, light will pass on the least rotation of the analyser to the right or left, and if the movement is continued another position is found on the disc in which the light is again cut off. This is at 180°, so that the circle should be graduated to 180° right and 180° left. Now, if a cell or tube with flat glass caps be filled with a solution of, say, sucrose and introduced between the two Nicols, and the source of light be a Bunsen burner flame coloured yellow with sodium chloride, when the index of the analyser is at 0° it will be observed that the light passes, and that on rotating the analyser a certain number of degrees to the right a point is found (which is not 180°) at which the light is cut off; and if note be taken of the exact number of degrees, and the analyser rotated further, it is found that the light is not again cut off at 180°, but the same number of degrees beyond 180° as the first extinction was beyond 0°. This is the optical activity of the sugar solution. It is the power of rotating the plane of polarisation of the ray passing through the polariser so as to admit of its passing through the analyser when it would not pass had the sugar solution not been introduced. The amount of rotation of the analyser to again find the direction of the plane of polarisation, as indicated by the index on the graduated disc, is the measure of the activity of the sugar solution, and the direction of the movement of the index is the direction in which the rotation has taken place; in the present case (i.e. when the index is moved on the disc in the direction of the hands of a watch) right or +. If the sugar solution be now hydrolysed by dilute acids or invertase, and again introduced between the Nicols, it is found that the analyser must be rotated to the left in order to find the position of extinction-i.e. the direction of the plane of polarisation of the rotated ray. This is left rotation, and is indicated by the sign -On further rotation of the analyser light again passes, and finally there is a second extinction as far beyond 180° as the first extinction was beyond 0°. From this it is clear that the second reading may indicate right-handed rotation as well as left. Say the first extinction was 10° left of 0° (-10°), the second would be +170°, 10° less than 180°; then the activity might be

-10° or +170°. On diluting the solution say one-half, if the activity is - the readings will be -5° and $+175^{\circ}$, if + they will be $+85^{\circ}$ and -95°. The exact measurement of this activity has been the subject of much investigation, and many plans have been devised to facilitate the operation. It is, however, beyond the scope of this article to deal more fully with the subject (for details see Das optische Drehungsvermögen organischer Substanzen, Landolt, Braunschweig, 1879, Engl. Trans.; Handbook of the Polariscope, Macmillan & Co., 1882; Watts' Dictionary, Suppl. 3, art. Light). In the experiments above described the light of a sodium flame was employed; it was observed early in the investigation of the phenomena that readings sufficiently accurate, especially if the amount of activity was small, could not be obtained. Hence other devices were introduced. In the literature we find the activity expressed $[a]_{R}$, $[a]_{J}$, or $[a]_{D}$ and $[a]_{B}$, $[a]_{G}$, $[a]_{Id}$, &c. $[a]_{R}$ values are not now used. They were

[a] values are not now used. They were obtained by employing white light, and interposing between the polariser and the source of light a plate of red glass. They are of no value for our purpose, and may be neglected.

[a], values are frequently used, but unless they are referred to a definite standard they, too, are of no value. If any ordinary source of white light be substituted for the sodium flame in the experiments above described, it will be observed that on rotating the analyser to find the effect produced by the sucrose solution no position of the analyser can be found in which all the light is cut off, but at one stage marked blue light passes, and, on further rotation of the analyser, red. This is due to the fact that the sugar solution acts unequally on the different rays of which the beam of white light is composed, the red rays being rotated least and the violet most, so that when the analyser is rotated the ray the plane of polarisation of which is rotated least is first cut off. The first cut off rays are the red, the complement of those cut off passing, the dominant being blue; on further rotation the dominant becomes red. On rotating the analyser carefully it is found that the transition from the blue to the red is well marked; this is the ay value; it is the, so-called, transition tint, the ray complementary to the median yellow, jaune moyen; hence aj. For the same strength of sugar solution the position of the analyser is not the same for every source of white light. The stage at which the transition appears with candle-light is different from that at which it appears with gas-light; the stage in this case differs again from that obtained with the electric light, and even the position for direct sunlight differs from that of the light reflected from a cloud. Hence, when a pair of Nicols are employed as described above, even when the source of light is given, a cannot be looked upon as a standard value. This value has, however, been given to it by the French instrument makers.

Soleil invented an instrument with which lamp- or gas-light is employed (afterwards improved by Dubosoq, and known as the Soleil-Dubosoq (see Watts' Dictionary, 1st ed. 3, 674, a. Suppl. 8, 1199), by which the activity of a solution is measured by divisions of a scale instead of degrees; 100 of these divisions were

made equal to the amount of rotation produced by 1 mm. quartz for the a_j ; this was considered to be 24°. Of course, if this is taken as a standard it is of full value; it may not be truly a_1 even for the quartz, but it is a_1 referred to a definite measure, the 100 divisions of the instrument being 24°. This was complicated by the fact, since pointed out by Scheibler, that quartz from different sources has not the same optical activity; hence we have various observers giving different values for the amount of sucrose in 100 c.c. solution, which when observed with the Soleil-Duboscq instrument, in a tube 200 mm. long, equals the activity of 1 mm. quartz. If, however, we look upon the 100 divisions of the instrument as $=24^{\circ}\alpha_{j}$, we have a standard value for the expression. I believe this is the value given to it by the French observers. The writer employed it in the same way. Another instrument of much the same construction is the Soleil-Ventzke-Scheibler (v. Watts, l.c.). In this the 100 divisions of the scale are made to represent the activity of 26.048 g. saccharon in 100 c.c. solution; we have only to inquire as to the $[a]_j$ and $[a]_D$ of cane sugar to arrive at a value in degrees of the 100 divisions for these two expressions. From these two factors we get 100 divisions, Soleil-Duboscq = 24° a, and $21.67^{\circ}\alpha_{D}$, and 100 divs. S.V.S. = 38.34° a, and $34.55^{\circ}\alpha_{D}$. The angular value for any other instrument in use in which the scale is given in divisions can be calculated for a_j or a_D by ascertaining the number of divisions required to compensate a solution of sucrose containing 10 g. in 100 c.c. at 15.5°, and taking the apparent specific rotatory power of the sugar for the concentration to be $[a]_j = +73.8^{\circ}$ and $[a]_D = 66.6^{\circ}$, the value of the divisions in degrees for both rays can be calculated. The direction of the rotation is indicated in these instruments by the direction of the movement of the 0 point to find the compensation necessary; if from left to right it is right or +, if in a contrary direction left or -. With the other instruments in use sodium flame is the source of light employed; the readings are therefore an. The scale is a circle graduated to degrees on a disc, beginning with 0 at the top and graduated in half degrees and degrees to 180° to the right, and in the same way to the left. The analyser rotates in the axis of this disc, and carries an index with vernier, which admits of readings being made to minutes. The arrangement is such that the polarised ray before entering the active solution has the appearance of a circular disc divided into halves by a perpendicular diameter. The position of the plane of polarisation is not the same in both semidiscs, so that when the light is completely cut off at one side it passes at the other, and vice versa. To find these positions the analyser has to be rotated several degrees, but between the two positions of complete cut off there is a position of the analyser in which the two semidiscs are equally illuminated; this is a wellmarked position distinctly ascertainable. The 0 point of the index and of the graduated scale are made to correspond exactly with it. It can always be found to within less than a minute. When the active substance is introduced the equality of the illumination of the semi-discs is

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destroyed; the analyser is rotated until it is again restored. The amount of rotation as shown by the graduations gives the activity in degrees and minutes, or degrees and decimals, according to the make of the vernier. These are the half-shadow instruments, of which the Jellet-Cornu and Laurent are examples (see Watts, l.c.). Other devices have been employed to facilitate an accurate determination of the position of the plane of polarisation of the ray before and after rotation; a description will be found in Landolt and in Watts (l.c.). Recently an instrument has been introduced in which the half-shadow contrivance is adopted, white light (a strong gas-flame) being employed. The scale is the same as that of the Soleil-Ventzke-Scheibler instrument; readings with it can be made with great accuracy. It is manufactured by Schmidt & Haensch, Berlin.

When it is necessary to determine the activity for rays of other refrangibility, say for the lithium or the thallium flame, it is only necessary to colour the Bunsen flame with these metals in the same way as in the case of sodium above. The readings are then α_{IA} and α_{Th} . The α_{B} , α_0 , α_0 , &c. for any active substance can be found by employing a pair of ordinary Nicols as described above—white light, best a ray of sunlight, being employed—and examining the light after passing the analyser, with a spectroscope. If the positions of the lines of the solar spectrum are marked, the amount of rotation for the active substance introduced between the polariser and the analyser is found, when on rotating the analyser the thickest part of the black band is on the marked position of the line of the spectrum. This is read off on the

scale connected with the analyser.

With any of these instruments we can determine the optical activity of a sugar and thus identify it. The sugar is crystallised; it is dried, and 10 g. or thereabouts carefully weighed out. This is dissolved in water, and the solution made up to 100 c.c.¹ at 15·5°. A portion of this is introduced into a tube of definite length, and an observation made with any of the instruments mentioned above. The formula $[a] = {}^{-100}$ gives the apparent specific rotatory power for the concentration. a = the angular disturbance observed, l = length in decimetres of tube employed, and o = grams substances in 100 c.c. Let us take an example, 10 g. sucrose, dissolved and made up to 100 c.c. Disturbance observed in a 200 mm. tube, with a Soleil-Dubosoq instrument, = 61·5 divs.; now for a_1 100 div. = 24°. Then, substituting the values in the above formula we have

$$\frac{61.5 \times .24 \times 100}{2 \times 10} = [\alpha]_{j} = +73.8^{\circ}$$

100 divs. = 21.67° α_D ; then

$$\frac{61.5 \times .2167 \times 100}{2 \times 10} = [\alpha]_{D} = +66.6^{\circ}.$$

The scale of the Soleil-Ventzke-Scheibler instrument is so graduated that 100 divisions = the activity of a solution of sucrose containing

¹ This is practically the usual 100 c.c. of a volumetric analysis. For more accurate manipulation see Landolt. This is, however, unnecessary for the identification of the sugar—our purpose.

26.048 g. in 100 c.c. in a 200 mm. tube. The apparent specific rotatory power of sucrose in a solution of this concentration is $[a]_D = 66.33^\circ$ and $[\alpha]_j = 73.61^\circ$. Substituting these values in the formula we get

x.100 $\overline{2 \times 26.048} = 66.33$;

x = the value of 100 divisions in degrees $\alpha_{\rm p}$ $= 34.55^{\circ}$, and

$$\frac{x' \cdot 100}{2 \times 26.048} = 73.61$$
;

x' = the value of 100 divisions $a_1 = 38.34^{\circ}$.

It is of importance that the values in degrees of the scales of these two instruments should be clearly understood. We find it stated that a quantity of sugar varying from 16.49 to 16.35 g. in 100 c.c. require, in a 200 mm. tube, a compensation with a Soleil-Duboscq instrument = 100 divisions; 16.35 g. is the best-recognised factor; from my observations it is still too high.

$$\frac{100 \times \cdot 24 \times 100}{2 \times 16 \cdot 35} = [a]^{1} = 73 \cdot 39^{\circ}, \text{ and}$$

$$\frac{100 \times \cdot 2167 \times 100}{2 \times 16 \cdot 35} = [a]_{D} = 66 \cdot 27^{\circ},$$

figures slightly too low for the apparent specific rotatory power of sucrose for the c, hence 16.35 g is too high. The number, no doubt, should be 16.28, and this gives factors agreeing well with the observed $[\alpha]_D$ and $[\alpha]_j$ for the c.

A solution containing 10 g. sucrose in 100 c.c. requires a compensation in a 200 mm. tube of 38.5 divisions f a Soleil-Ventzke-Scheibler instrument.

$$\frac{38.5 \times 3834 \times 100}{2 \times 10} = [\alpha]_1 = 73.8^{\circ} \text{ and}$$

$$\frac{38.5 \times 3455 \times 100}{2 \times 10} = [\alpha]_D = 66.5^{\circ},$$

numbers agreeing accurately with the specific rotatory power for the c. The optical arrangements of these instruments admit of the employment of white light.

If either of the sodium-flame instruments is employed the angular disturbance is introduced directly into the formula. With the 10 g. in 100 c.c. solution of sucrose, the reading in a 200 mm. tube is

 $a_D = +13.3^{\circ}$, then $\frac{13.3 \times 100}{2 \times 10} = [a]_D = 66.5^{\circ}$. The use of this factor is obvious. We have a solution of a known sugar of unknown strength, and want to find the c (g. in 100 c.c.). An observation is made in a tube of known length, $a_D \text{ or } a_j$; then $\frac{a.100}{l.c}$ - the determined value of

[a]D or [a], a the ray for which the observation is made, and [a] the specific rotatory power for that ray, according to the light employed and

concentration, $c = \frac{\alpha \cdot 100}{[\alpha] l}$.

It has been shown that if a solution contains two or more active substances, the observed activity is the algebraic sum of the activities of the constituents. The specific rotatory power of each individual sugar is given in the description

5. The reducing power. As has been stated, most of the sugars possess the power of redu-

cing the oxides of the higher metals and some organic colouring matters. Various propositions have been made to employ this property as a means of qualitatively and quantitatively determining the sugars, and various solutions have been proposed for that purpose; but speaking with many years' experience I may say that they may all be discarded in favour of alkaline copper solution, Fehling's solution; and even with regard to the value of this a great diversity of opinion exists. But there is a general agreement that if the conditions under which it is used are constant the results are constant. For literature see Becquerel, A. Ch. [2] 47, 15; Trömmer, A. 38, 360; Müller a. Hagen, Pf. 23, 221; Neubauer, Fr. 1, 378; Maly, Fr. 10, 383; Seegen, C. C. 1875, 223; Fehling, A. 72, 106; Claus, J. pr. [2] 4, 63; Neubauer, Ar. Ph. [2] 71, 278; Patterson, C. N. 25, 149; Loiseau, C. R. 1873. 26; Soxhlet, J. pr. [2] 21, 227; Krause a. Städeler, C. C. 1854. 936; Gräger, Fr. 7, 490; Buswitz, B. 11, 1445; O'Sullivan, C. J. 1876, 2, 125; Märcker, O. 7, 699; Brunner, Fr. 11, 32; Bayley, C. N. 37, 211; Allihn, J. pr. [2] 22, 55; Salomon, B. 14, 2711; Digener, Z.V. 31, 349; Schwarz, A. 84, 84; Mohr, Fr. 12, 296; Perrot, B. 9, 19; Ulbricht, B. 10, 128; Jean, C. R. 73, B. 5, 15; tollard, 2. 9, 820; Weil, Fr. 11, 284; Arnold, Fr. 20, 331; Volhard, A. 190, 1; Soldaini. B. 9, 1126; Possoz, C. R. 1874, 721; daini, B. 9, 1126; Possoz, C. R. 1874, 721; Pellet, J. Fabr. 19, 22; Pavy, C. N. 39, 1004; Löwe, Fr. 9, 20; Vogel, J. Ph. [2] 1, 245; and others of less value. As a result of a study of all this literature, I may say that I have come to the conclusion that if the Fehling's solution is carefully made with pure materials, and employed as described below, the reducing power of any sugar can be determined with constancy and accuracy, and when the conditions are fulfilled the reduction is molecular—i.e. a definite number of copper oxide molecules are reduced by a definite number of molecules of the sugar.

Fehling's solution is made as follows: 34.61 g. pure recrystallised copper sulphate are dissolved in 400 c.c. H₂O or thereabouts, 173 g. sodium potassium tartrate (Rochelle salt), and 74° grams freshly-prepared sodium hydroxide (commercial stick soda) are dissolved to 450 H₂O; when the solution is cold it is gradually added to the copper solution in small portions at a time, so that the pp. at first formed is re-dissolved. When both solutions are mixed the volume is made up to 1 litre. It is very generally stated that this solution does not keep very well. I found it constant in its properties for more than twelve months when kept in carefully-stoppered bottles, sealed with paraffin, in a cool, dark place. Some recommend that both the solutions be made up to 500 c.c., kept separately and mixed in equal volumes when required for use. This is probably the better plan, but unless the solutions are kept in well-stoppered bottles sealed they will deteriorate as badly as if mixed. This solution is used as (a) a qualitative test for the reducing sugars.

O. = Organ des Centralvereins für Rübenruckerfabrioation in der österreichisch-ungarischen Monarchie.
 Journal des Fabricants de Sucre.
 Redewald a. Tollens (B. 11, 2076) employ 60 g., and they say the increase of alkali to 70 g. diminishes the reducing power of lactore about 1 p.o.

is also used (b) quantitatively, to determine the reducing power and the quantity of a sugar.

(a) Qualitatively. A portion of the solution supposed to contain sugar, prepared as described, is heated to boiling in a small beaker or test-tube, and is added to 5 c.c. of the alkaline copper solution previously diluted with its own bulk of water, and heated to boiling. The production of a red pp. of Cu₂O indicates the presence of a reducing sugar. There are substances besides the sugars capable of reducing Fehling's solution. The absence of these must be established before it can be inferred that the reduction is due to sugar.

(b) Quantitatively. The reducing power of a sugar, or of a solution containing reducing sugars, can be determined volumetrically or gravimetrically, with Fehling's solution. It is now very generally admitted that the gravimetric method gives the only reliable results.

Volumetrically. The sugar solution, prepared as described, should be diluted (0.5 to 1.0 g. reducing sugar in 100 c.c.). This is introduced into a burette, and the Fehling's solution into a second one. Of the latter a definite quantity, say 10 c.c., is measured into a porcelain dish of about 60 to 100 c.c. capacity, and diluted with four times its own bulk of water. The dish is then placed over a small Bunsen, and the contents heated to boiling. The sugar solution is then run in a c.c. at a time until the blue colour of the copper solution has nearly all disappeared, then drop by drop until it has quite gone. The first experiment may only give approximate results; in a second a quantity of sugar slightly less than was found necessary in the first experiment is measured into a small beaker, heated to boiling, and mixed rapidly with the boiling dilute copper solution. If the whole of the copper is reduced, a few drops more of the Fehling's solution are added from the burette. A few experiments of this kind will give fairly approximately the amount of copper solution a certain quantity of the sugar solution is capable of reducing. That the Fehling's solution is not in excess is seen by the absence of blue on subsidence of the Cu2O, and may be confirmed by filtering a little and testing for copper with a dilute solution of potassium ferrocyanide in acetic acid. The absence of a brown colouration indicates the absence of CuO. If the sugar is in excess, more CuO solution is decolourised, but after boiling the sugar in the alkaline solution the amount reduced is not a measure of the quantity of sugar. The final experiment must be performed after two or more observations, when the relative value of the two solutions has been approximately ascertained, as described above, by adding the whole, or nearly the whole, of the sugar at a boiling temperature to the copper solution at once. If only a few drops of the sugar solution are required to remove the last traces of CuO, the error is not great; 10 c.c. of the Fehling's solution, made as directed, contain 1100 g. CuO; 396 CuO = 180 dextrose, i.e. 5 mols. CuO (79.2 + 5 = 396) are reduced by 1 mol. dex-396:110 = 180:050180, then dextrose = 10 c.c. Fehling's solution. Soxhlet (l.c.) states that the value of the CuO solution in dextrose varies with the way in which the experiment is performed; that 10 c.c. undiluted Fehling's solution are reduced by

0.04753 g. dextrose, while it takes 0.0494 to reduce the same bulk diluted with four measures of water. The same applies to the other reducing sugars, only in the case of lactose dilution has no effect, while in that of maltose dilution diminishes the amount of sugar required. This is, no doubt, correct under the conditions stated, but it is not necessary in dealing with a reaction of this kind to deviate from recognised conditions such as those laid down above. When these are adhered to it is found that the reducing power of the sugar very closely approaches molecular interaction; in fact, so much so that the molecular relation most closely approximated to by the experimental data may be taken as the normal reducing power of a sugar. These normals for the betterknown sugars are:-

10 c.c. Fehling's solution = 0.0500 g. dextrose.*

" 0.0500 g. galactose.

" 0.0500 g. lævulose.

" 0.06334 g. lactose.†

" 0.07196 g. maltose.†

" 0.04630 g. arabinose.

" 0.08701 g. arabinon.

Soxhlet says the reducing power of lævulose is different from that of dextrose. This is not supported; his error is due to his mode of preparing invert sugar.

† Sugar free from water of crystallisation.

Gravimetrically. The Fehling's solution is prepared as described; 25-30 c.c. of it are measured into a beaker of 130-140 c.c. capacity, and about 50 c.c. well boiled water added. The beaker is then placed in a boiling-water bath, consisting of a larger beaker in which water is kept boiling. At the end of five or six minutes. when the dilute copper solution has acquired the temperature of the bath, a known weight or measure of the sugar solution is heated to boiling, and added. If at the end of a few minutes the blue colour is completely destroyed, it can be restored by adding more Fehling's solution, but it is much better to so arrange matters by a few experiments that the amount of sugar solution employed will reduce the Fehling's as nearly as possible, leaving only sufficient blue, on subsidence of the Cu₂O, to make it evident that the copper solution is in slight excess. After twelve to fourteen minutes' boiling the clear supernatant liquid is decanted on to a filter, the Cu₂O is washed by decantation, and finally on the filter. If the Fehling's solution is in large excess it is difficult to wash the filter free from unreduced copper oxide (blue colouration), but if the excess is only slight there is no indication that the filter retains copper compounds other than Cu₂O. Ignition for four or five minutes in an open porcelain crucible converts the red precipitate into black copper oxide CuO, in which form it is weighed. From this the sugar is calculated by multiplying by the constants (see next page). It may be again pointed out that the factors given by various authorities differ more or less from these (see the K of individual sugars), but I believe these are the constants, the differences amongst the authors being due to manipulation, to the state of dilution and alkalinity of the solutions, and to the temperature of the mixed sugar and CuO solution. An imperfect knowledge of the composition of the material is a source of error. In dealing with some solutions, other

```
1 mol. sugar to 5 mols. CuO: 1 g. CuO=0.4545 g. dextrose : K=100
                             1 g.
                                   " = 0.4545 g. galactose : K = 100
               5
                 ,,
                        ,,
                             1 g.
               5
                                      = 0.4545 g. lævulose : K = 100
 ,,
               7.5 "
                             .1 g.
                                      = 0.5758 g. lactose
                                                            K = 77.67
                                   ,,
 ,,
        ,,
                         ,,
                                                            K = 63.16
                             1 g.
                                      = 0.7196 g. maltose
               6
                                   ,,
 ,,
        ,,
                         ,,
               4.5 "
                             1 g.
                                      =0.4209 g. arabinose : K=108.8
               4.5 "
                             1 g.
                                      = 0.7910 g. arabinon : K = 57.5
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substances may be precipitated with the Cu₂O. These should be ascertained, determined, and allowed for. In dealing with solutions in which ammonia is found, or liberated by the alkali, Cu₂O is retained in solution. Sundry propositions have been made to avoid the disturbing influence of the alkali in Fehling's solution, but they seem to have met with no favour. Soldani (B. 9, 1126) proposed copper carbonate in potassium bicarbonate; Possoz (C. R. 1874, 721) and Pellet (J. fb. 19, 22) other solutions of copper in alkaline carbonates. Löwe (Fr. 9, 20) recommends the use of a solution of CuO in glycerol, but so far little can be said of the value of the solution. In cases where the sugar solution contains other substances than reducing sugars capable of giving a precipitate with Fehling's solution without reduction, the whole precipitate may be dissolved in hydrochloric acid, made strongly alkaline, and added to an excess of ammoniacal solution of silver nitrate. The copper suboxide reduces an equivalent quantity of silver oxide. The silver in the precipitate may be estimated as chloride (Jean, C. R. 73, 1397). As has been stated above, many other solutions and modifications of Fehling's solution, have been proposed for employment in determining the sugars, but they appear to be of no value.

Pavy (l.c.) recommends the addition of sufficient ammonia to the Fehling's solution to hold the Cu₂O in solution; the disappearance of the blue is then the indicator of the reaction. The reducing powers for sugars other than dextrose for this solution have not been ascertained; even for dextrose it is doubtful (v. Hehner, An.

6, 218).

Three solutions of mercury salts have been proposed as agents by which sugars can be determined by their reducing power—the one known as Knapp's solution, the other as Sachsse's solution, and the third as Hager's solution. Neither of them are of high value, except in certain circumstances, but Hager's least of all.

Knapp's solution is made by dissolving 10 g. HgCy, in about 600 c.c. water, then adding gradually 100 c.c. sodium hydroxide solution, specific gravity 1.145, and diluting the clear solution to 1 litre; 10 c.c. of this solution equal 0.025 g. dextrose; but according to Soxhlet (l.c.) this is not true, and the quantity varies with the conditions under which the reducing solution and reducible one are brought in contact, and even with the indicator used in determining the absence of mercury. The solution is used as follows:-40 c.c. of it are heated to boiling in a flask, and the sugar solution, which should not contain more than 0.5 p.c. reducing sugar calculated as dextrose, run in as quickly as possible until the whole of the mercury is ppd. as such, the mixed liquids being kept continually boiling. The complete prespitation of the Hg is ascertained by testing for

it with ammonium sulphide, or alkaline tin solution. Less mercury is reduced by adding the sugar gradually than when the addition is made all at one time (Brunner, J. pr. [2] 21; Soxhlet 1.c.). According to the latter, when the reducing solution is added all at one time, and alkaline tin solution employed as a test for the absence of Hg from the solution, 0.200-202 g. dextrose reduce 100 c.c. of Knapp's solution.

Sachsse's solution is made to contain in 1 litre 18 g. mercuric iodide, 25 g. potassium iodide, and 80 g. potassium hydroxide. The mercury salt is dissolved in a solution of the potassium iodide, the solution of the potassium hydroxide added, and the mixture made up to the litre. A measured quantity, 40 c.c., of the solution is boiled in a porcelain dish, and the sugar solution, containing not more than 0.5 g. per 100 c.c., run in until a drop of the supernatant liquid ceases to give a brown colour with an alkaline solution of tin chloride. It is said that in using this reagent more mercury is reduced by adding the sugar slowly than by adding it all at one time, the reverse being the case with Knapp's solution. Soxhlet (l.c.) compares the value of these solutions for the reducing sugars as follows:-

Taking the reducing power of dextrose = 100, the reducing powers (K) of the other sugars are:

	Fehling (undiluted)	Knapp	Bachsse
Dextrose	. 100	100	100
Invert sugar .	• 96.2	99	124.5
Lævulose .	• 92.4	$102 \cdot 2$	148.6
Galactose .	. 93.2	83.0	74.8
Lactose (hydrated	l?) 70·3	64.9	70.9
Maltose (dry?).	. 61.0	63.8	65.0

These relations for the K of the sugars for Fehling's solution do not agree at all with the generally recorded numbers (v. K of individual sugars); there are no data to enable one to form an opinion on the value of the K for the other solutions.

Potassium ferricyanide has been proposed asan agent for determining the reducing power of the sugars by Gentele (Fr. 9, 458), but although Stahlschmidt (B. 1, 141), Stammer (D. P. J. 158, 40), and Sostmann (V. Z. 22, 170) have worked with it, we know nothing very definite about its value. It should yield definite results, as the complete reduction is definitely recognisable.

6. The action of unorganised ferments. Enzymes. Sucrose is converted by invertase into lævulose and dextrose; the only other sugar acted upon by this enzyme is raffinose (Kjeldahl, Fr. 22, 588; O'Sullivan, C. J. 1886. 70; and O'Sullivan a. Tompson, C. J. 1890. 834), and, as it has been shown, the invertive action is most active at a temperature at which yeast, from which the invertase is prepared, has no fermentative action on the products of the inversion; yeast can be employed instead of prepared invertase. The products of the action of invertase.

on raffinose are lævulose and melibiose, the latter yielding finally dextrose and galactose. Raffinose is but very sparely distributed. It may be considered that sucrose is present if the optical activity of a solution is materially altered by treatment with invertase and the copper oxide reducing power at the same time increased. The absence of raffinose can be determined by the non-production of mucic acid on treatment with nitric acid. Before applying the invertase test the solution to be tested must be boiled. If this is not done, and active diastase and any of the highstarch transformation products are present, the latter would be acted upon by the diastase, the opticity lowered, and the cupric reducing power increased. Diastase does not appear to act on any of the sugars, except isomaltose (?)

Other organisms, such as the moulds which grow on steamed rice, contain an enzyme capable of acting on maltose, but this substance has not been employed in the detection or estimation of maltose. There is no doubt, however,

it is available for the purpose.

7. Organised Ferments (see Fermentation). The hyphomycetes, saccharomycetes, and schizoinycetes act on the sugars and yield various products, and, no doubt, can be made available for the detection and estimation of them; but we have only to concern ourselves here with the action of ordinary beer yeast (saccharomyces cerevisiæ); the changes effected by the other organisms come under art. FERMENTATION. On adding pressed yeast to a very slightly acid solution, and keeping the mixture at a temperature of 20° or thereabouts, if a fermentable sugar be present, a frothy head will form on the surface of the liquid, carbon dioxide will be evolved, and alcohol formed. The specific gravity and optical activity of the solution diminishes. Certain substances known as antiseptics prevent this reaction—they must be proved to be absent. The fermentable sugars, in this sense, are described as such under the respective heads.

The quantitative value of this phenomenon has been studied by many workers, but it is still unsatisfactory. It has been made available with approximate accuracy in determining the total fermentable sugars in presence of other substances from which they cannot otherwise be conveniently separated. The solution should be of about sp.gr. 1.050-1.060, slightly acid, and should not contain much foreign matter. A measured quantity of this is taken, best 100 c.c. if available, introduced into a 150 c.c. flask, and 0.5 to 1 g. freshly-pressed yeast added. If necessary, a little yeast ash dissolved in the least possible quantity of hydrochloric acid, together with a little ammonium tartrate, should be added. A temperature of from 20° to 22° is best suited for the process. The gas evolved is washed by being made to pass through a few c.c. water in two successive wash-bottles. In some cases the fermentation is very slow and takes many days to complete itself, in others it is rapid and is finished in a few days; but in all cases it should be continued as long as gas bubbles through the wash-bottles on shaking the flask in which the fermentation is going on. As soon as this point is arrived at, the contents of the flask are washed with the water of the wash-bottles into a distilling flask, and submitted to distillation. Dilute alco-

hol distils over and is collected in a 100 c.c. flask. If only 20-25 c.c. wash water were used. the distillate, when it amounts to 80-85 c.c., will contain all the alcohol produced. This is then made up to 100 c.c. and its specific gravity taken. On referring to the alcohol tables, the weight of alcohol in the 100 c.c. is ascertained. From this a knowledge of the amount of fermentable sugars is arrived at. Sucrose yields 51 p.c. alcohol, dextrose 48-49 p.c. (Pasteur, A. Ch. [3] 58, 330), maltose 50.4-51.8 p.c. (O'Sullivan, C.J. 1876, 479). Hofmann, Graham, and Redwood (C. J. 5, 229) gave expression to the value of the specific gravity of distillate, in points or degrees of specific gravity lost by the saccharine solution fermented for various sugars. These factors are still employed in determining the original specific gravity of beer, from the alcohol it contains and the specific gravity of the residue after distillation. The residue left in the distilling flask mentioned above is with the washings made up to 100 c.c., the specific gravity of this solution subtracted from that of the solution before the yeast was added + 1-or 1000 if water be taken at 1000 instead of 1-will give approximately the specific gravity of a solution containing the amount of sugars or sugar fermented in 100 c.c. The average D of the sugars being known, the quantity fermented within 2 or 3 p.c. can be calculated from this. Inasmuch, however, as the value in specific gravity of the non-volatile products of the fermentation of the sugars has not been accurately determined, this mode of estimation can only be approximate.

It has been proposed to estimate the amount of sugar or sugars by determining the amount of carbon dioxide evolved during fermentation. Pasteur (*l.c.*) gives 0.4665 g. CO₂ as the product of 1 g. dextrose. Jodhauer (*Z.V.* 25, 308) state that dry maltose yields 49.54 p.c. CO₂ and dextrose 46.54 p.c. These figures are, no doubt, true also for all the fermentable '-on' and '-ose'

sugars.

Some of the sugars when in the pure state are not fermented by yeast, but the presence of a small quantity of an easily fermentable sugar sets up the action. Pure galactose is not fermented by yeast; the presence of a little dextrose, lævulose, or maltose sets up the process (Bourquelot, C. R. 106, 283; Tollensa. Stone, B. 21, 1572; Stone a. Tollens, A. 249, 257), the fermentation of galactose being as complete as that of dextrose if the suitable nourishment is Under the same condition sorbose present. ferments more slowly and less completely, arabinose and lactose very slowly and incompletely. Pure arabinose behaves like pure galactose. Sucrose is inverted by yeast, dextrose ferments easily, lævulose less so, maltose still less, and raffinose less still. The other sugars ferment only when these are present; hence it should be possible to detect the presence of individual sugars from the specific rotatory power and K by fractional fermentation. With some varieties of yeast lævulose ferments before dextrose (Dubourg, Again, some organisms are C. R. 110, 868). found to ferment the '-ose' sugars and to be incapable of fermenting the '-on' ones. A study of the action of these organisms will yield results valuable in the identification and estimation of the sugars.

8. Action of acids. All the di- and tri- sugars are converted into mono-sugars by the action of dilute mineral acids; this is accompanied by an alteration in the K and [a] of the solution. If this took place only with the high sugars, the reaction would be applicable in distinguishing the '-on ' and '-ose -on' sugars from the '-ose; but the K and [a] of these sugars are also affected by digestion with mineral acids, but much more slowly, so that the reaction is only available when carefully performed. If the sugar solution increase in K, and there is a material alteration in [a] on ten or twelve minutes' boiling with 2-3 p.c. sulphuric acid or hydrochloric acid, it may safely be inferred that a high sugar (di- or tri-) is present. Some organic acids act differently on the high sugars. Dilute citric acid inverts sucrose; it is without action, at least material action, in a short time on lactose (Jones, l.c.).

The invertive action of acids—except under well-defined conditions, such as the employment of hydrochloric acid by Clerget in hydrolysing sucrose—cannot, for the reason mentioned above, be employed in estimating these sugars.

Dextrose and substances capable of yielding it can be detected by acting on the material with nitric acid (1 pt. mat. 1 pt. H₂O and 5 pts. nitric acid, sp.gr. 1.15). Saccharic acid, recognisable by the microscopic appearance of its acid potassium salt, is produced (Sohst a. Tollens, A. 245, 1). Lævulose, galactose, sorbose, and arabinose do not yield this acid (Gans, Stone, a. Tollens, B. 21, 2148). Galactose, lactose, and raffinose, on treatment with nitric acid in the same way, yield mucic acid. This reaction has some quantitative value. Pasteur (C. R. 42, 349) obtained 75 to 78 p.c. mucio acid from galactose. The yield, however, varies very considerably with the quantity and strength of the acid employed. Kent a. Tollens (A. 227, 221) find on employing 1 pt. sugar, 12 pts. nitric acid, sp.gr. 1.15, allowing to crystallise in 1.5 to 2.0 pts. solution, and washing with 5 pts. water that galactose gives 77.4 p.c. mucic acid. Lactose treated in the same way gives 40 p.c., and raffinose (Rischbieth a. Tollens, A. 232, 172) 22-23 p.c. By using 1 g. dry sugar, or a quantity equal thereto, in a concentrated syrup, 4 g. nitric acid sp.gr. 1.25-1.24, heating gently until red fumes began to be evolved, then allowing the reaction to proceed until the violence was over, heating on a water-bath until red fumes ceased to be evolved, allowing to stand for 24 hours to crystallise, I obtained 73 p.c. mucic acid from galactose, 86.9 p.c. from dry lactose, and 28-30 p.c. from dry raffinose (O'Sullivan, C. J. 1886, 73). The mucic acid was collected on a tared filter, washed carefully with 50 c.c. water, dried in a vacuum over sulphuric acid, then at 100°, and weighed. By treating 0.5 g. mucic acid in the same way, the loss was 0.04-0.05 g. Corrected for this the above numbers become 80-81 p.c. for galactose and 40.5-41.5 p.c. for lactose. The number for raffinose was corrected.

The pentoses, and bodies from which they can be obtained, yield on digestion with strong hydrochloric acid, furfural, the detection of which is simple and definite. This reaction has been employed in the estimation of these sugars. It must not, however, be forgotten that the hexoses also yield furfural, although, no doubt, in very small

quantities (Günther a. Tollens, B. 23, 1751; de Chalmot a. Tollens, B. 24, 694; Stone, B. 24, 3019). The first difficulty to overcome was to determine the conditions under which the greatest yield of furfural was obtainable, the next to accurately determine the furfural. 2 to 5 g. of the material are digested with 100 c.c. hydrochloric acid, sp.gr. 1.06, and then distilling, returning the distillate until the reaction is complete; or the mixture of material and acid is submitted to distillation in a flask over a small flame, so that not more than 10 c.c. distillate pass over in five minutes, fresh acid being regularly added until a sample of the distillate ceases to give the reaction for furfural. The distillate is then neutralised with soda, acidified with a few drops of acetic acid, and made up to a definite volume. In this the furfural is estimated by a standard solution of phenylhydrazine, made by dissolving 2 g. phenylhydrazine chloride with 6 g. sodium acetate in 1 litre water. This is standardised with a solution of furfural of known strength (1 g. pure furfuramide in a little acetic acid to 1 ltr. water). Standardisation must be made before each set of determinations, as the phenylhydrazine solution loses its value in a short time. The titration is performed as follows: 25 c.c. of the distillate are mixed with a measured quantity of the phenylhydrazine solution, the mixture rapidly boiled and cooled; a little of the solution is filtered and boiled with twice its volume of Fehling's solution; if the phenylhydrazine is in excess reduction takes place, but if furfural is in excess there is no reduction. Trials are made until the equivalent amount of phenylhydrazine solution is found (Stone, l.c.). The dilute acidified solution is precipitated with phenylhydrazine acetate, the hydrazide collected on an asbestos filter and dried in a vacuum in a specially-constructed apparatus at 50°-60° until the weight becomes constant. The precipitation is always effected in the same volume of solution, so that allowance can be made for the solubility of the hydrazide. Arabinose yields on an average 48.72 p.c. and zylose 56.25 p.c. furfural (de Chalmot a. Tollens).

9. The action of alkalis. This can only be applied as a rough qualitative test. All the hexoses yield a brown colouration when boiled for a few minutes with a 3-4 p.c. solution sodium hydroxide. The '-on' sugars are not affected.

10. The action of phenylhydrazine. The melting-point of the osazones and hydrazides and composition thereof give good indication of the nature of the sugar whence they are prepared. For preparation and melting-point see individual sugars. Maquenne (C. R. 112, 799) finds that different sugars yield, with excess of phenylhydrazine, very different quantities of osazones. By heating 1 g. each sugar for one hour at 100° with 100 c.c. water, and 5 c.c. of a solution of phenylhydrazine acetate, containing in 1 litre 400 g. phenylhydrazine and 400 g. glacial acetic acid, cooling, collecting the precipitate, washing with 100 c.c. water and drying at 110°, he found sorbose gave 0.82 g., lævulose 0.70, xylose 0.40, dry dextrose 0.32, arabinose 0.27, galactose 0.23, rhamnose 0.15, lactose 0.11, maltose 0.11. There must, however, be some error or misunderstanding here, as there is no difficulty in getting 120 p.c. osazone from arabinose and more than 130 p.c. from lævulose.

SULPHACETIC ACID v. SULPHOACETIC ACID. SULPHAMIC ACID and SULPHAMATES SO₂.NH₂.OH and SO₂.NH₂.OM¹. (Amidosulphonic acid and Amidosulphonates.) By passing dry NH₃ over SO₃, Rose (P. 33, 81; 47, 471; 49, 183) obtained a compound SO₃.2NH₃. This compound, called by Rose 'sulphatammon,' is genedescribed as ammonium sulphamate rally SO₂NH₂ONH₄, but Divers a. Haga (C. J. 61, 948) have shown, fairly conclusively, that the compound is tri-ammonium imidosulphonate $(N(NH_4)(SO_2.ONH_4)_2 [= 2SO_3.4NH_3];$ this salt is described under Imidosulphonic acid and its salts (v. Sulphonic acids and derivatives, p. 600).

Barium sulphamate SO₂.NH₂.O Ba seems to have been obtained by Berglund (B. 9, 1896) by boiling an aqueous solution of barium imidosulphonate HN < SO₂.O > Ba (v. Imidosulphonic acid under Sulphonic acids and Derivatives, p. 600) till the solution reacted strongly acid, adding excess BaOAq, boiling so long as NH, came off, filtering, removing excess of Ba by passing in CO₂, filtering, evaporating the filtrate to a small bulk, and allowing to stand. The salt crystallises in long, lustrous needles; it is not changed by heating to 200°, nor by boiling in solution, even in presence of HClAq.

Berglund (l.c.) says that a solution of potassium sulphamate is obtained by the reaction of nascent hydrogen with potassium oxyamidosulphonate (v. Oxyamidosulphonic acid under SULPHONIC ACIDS AND DERIVATIVES, p. 602), thus: $NHOH.SO_2.OKAq + 2H = NH_2.SO_2.OKAq + H_2O.$ B. was unable to isolate the salt. Raschig (A. 241, 176) obtained the potassium salt SO₂.NH₂.OK by boiling potassium imidosulphonate NH(SO₂.OK)₂ with a little water for some time, removing the H₂SO₄ that was produced by CaCO, filtering from CaSO, evaporating, pouring off from K2SO4 that separated, and allowing the syrupy mother-liquor to crystallise. The salt separated in large colourless, rhombic crystals. In 1878 Berglund prepared several sulphamates, viz. the salts of NH₄, Cd, Ca, Co, Cu, Pb, Li, Mn, Ni, Ag, Na, Sr, Tl, and Zn (Lunds Universitets Arsskrift, 13; abstract in Bl. [2] 29, 422).

SULPHAMIC ACID SO. NH2.OH. This acid was obtained by Berglund (l.c.) by passing H₂S into a solution of the silver salt, filtering from Ag2S, and evaporating over H₂SO₄. It forms large, translucent, rhombic crystals; the ratio a:b:c = 9945:1:1.056 (Raschig, A. 241, 178). Easily sol. water, less sol. alcohol; aqueous solution only slowly decomposed to NH4.H.SO4 on boiling; more rapidly decomposed if HClAq is added; not decomposed by heating to 190° (B., l.c.). solution of the acid gives no pp. with BaOAq; on boiling with HClAq and BaCl,Aq a pp. of BaSO, gradually forms. Raschig (l.c. p. 209) obtained sulphamic acid by saturating a solution of hydroxylamine hydrochloride with SO2, allowing to stand for a little, and then evaporating on a water-bath (NH2OH.HClAq + H.SO2-OHAq

= NH₂.SO₂.OHAq + HClAq + H₂Ô).

SULPHAMIDE SO₂(NH₂)₂. (Sulphonamide.

Sulphurylamide.) The neutral amide of sul-

phuric acid is obtained by passing dry NH, into cooled SO₂Cl₂, diluted with 15 to 20 vols. CHCl₂, to complete saturation, dissolving the pp. so produced in water, making strongly acid by HNO₂Aq, ppg. Cl as AgCl by addition of AgNO₃Aq, filtering, neutralising by KOHAq, and again ppg. by AgNO₃Aq, filtering from SO₂NAg (v. Sulphimide, p. 587), adding more AgNO, Aq and then KOHAq, warming and then cooling quickly, and collecting the pp. that forms. This pp. consists chiefly of SO₂(NHAg)₂, but there is a small quantity of another Ag salt which must be removed. This is done by washing well, adding exactly enough HClAq to convert all Ag into AgCl, neutralising by NH,Aq, adding AgNOsAq, filtering from the pp. of the foreign Ag salt, adding more AgNO, Aq and then excess of NH, Aq, when pure SO2(NHAg)2 is ppd. The salt is decomposed by exactly the proper quantity of HClAq, AgCl is filtered off, the filtrate is evaporated in vacuo at a temperature not above 40°, and the liquid is allowed to remain in vacuo over $\rm H_2SO_4$, when $\rm SO_2(NH_s)_3$ separates in large, colourless crystals (W. Traube, B. 26, 607). Sulphamide is very sol. water, less sol. dilute alcohol, insol. alcohol, ether, and other ordinary organic solvents; it softens at 75° and melts at 81°, begins to give off NH₃ below 100°, no further decomposition occurring to 250°; above 250° gives off acid vapours, and is completely decomposed. An aqueous solution of sulphamide is neutral; boiled with acids it gives H.SO.Aq and NH₃. Solutions of alkalis split off NH₃ and form alkali salts of SO2.OH.NH2. Small quantities of H2SO4Aq are formed after prolonged boiling. By adding AgNOsAq to solutions of sulphamide, and then NH, Aq, a pp. of SO₂(NHAg)₂ is formed; other metallic derivatives of SO₂(NH₂)₂ exist, but have not yet been isolated satisfactorily.

IMIDO-SULPHAMIDE NH(SO₂,NH₂)₂ v. SULPH-DO-AMIDE (D. 587). M. M. P. M. IMIDO-AMIDE (p. 587). M. M. P. M. SULPHAMINE-BENZOIC ACID v. Amide of

O-SULPHO-BENZOIC ACID.

SULPHAMMONIC ACIDS AND SALTS v. SULPHUR OXYACIDS, NITROGEN DERIVATIVES OF, p. 619.

SULPHANILIC ACID v. vol. i. p. 154.

SULPHANTIMONATES, &c. For such salts as sulphantimonates, sulpharsenates, sulphophosphates v. Thio-antimonates &c., under ANTIMONATES &c.

SULPHATAMMON and PARASULPHATAM-MON v. Tri-ammonium imidosulphonate and Di-ammonium imidosulphonate under Imidosulphonic acid and salts, a section of SULPHONIC ACIDS AND DERIVATIVES, p. 600. SULPHATES and allied salts. So

Salts of sulphuric acid H₂SO₄. The compositions of the normal sulphates are expressed by the general formulæ M2ISO4, MIISO4, MIII2(SO4), and M^{IV}(SO₄)₂; the greater number of the acid sulphates come under such formulæ as MIHSO. $M^{II}H_2(SO_4)_2$, $M^{III}H(SO_4)_2$, $M^{IV}H_2(SO_4)_3$. All sulphates, whether normal, acid, or basic, may be represented by the formula nMO.mSO, xH2O, where MO stands for a basic oxide. The sulphates are prepared (1) by the interaction of H₂SO₄Aq with metals, metallic oxides, or hydroxides; some metals react with dilute H, SO, Aq, giving sulphates and H, others react only with

hot conc. H₂SO₄, giving sulphates and SO₂, or SO, and H.S, and sometimes H also; (2) by the interaction of H.SO.Aq with salts of volatilisable or decomposable acids, e.g. with chlorides, nitrates, or carbonates; (3) by the direct combination of SO, with metallic oxides; (4) by oxidising sulphides, by boiling with HNO, Aq, or by reacting with Cl in presence of water, or sometimes by heating in air, e.g. FeSO₄, CuSO₄, and ZnSO₄; (5) by double decomposition from other sulphates, e.g. BaSO₄, PbSO₄. Most of the metallic normal sulphates are soluble in water; BaSO₄ is insol. water, PbSO₄ and SrSO₄ nearly insol., and CaSO, very slightly soluble. The basic sulphates are generally insoluble in water. Sulphates of metals whose oxides are strong bases—the alkali and alkaline earth metals—are not decomposed by heat alone; the sulphates of metals whose oxides are weak bases are decomposed by heating, giving off SO2, or SO2 and O, and leaving oxides, or metals if the oxides are reducible by heat. Sulphates are reduced by heating with charcoal, either to sulphides or oxides, with evolution of CO and CO, and also SO, from the sulphates of weak bases. Many sulphates are decomposed completely by heating with HCl gas, giving chlorides (v. Hensgen, B. 9, 1671; 10, 259). Fusion with excess of alkali carbonate produces alkali sulphate and carbonate of the metal of the original sulphate. Many sulphates are reduced by strongly heating in a stream of NH, yielding sulphides, oxides, metal, or mixtures of these (v. Hodgkinson a. Trench, C. N. 66, 223). Several sulphates occur as minerals; e.g. BaSO₄ (heavy spar), CaSO₄ (gypsum), SrSO₄ (celestine), MgSO₄ (Epsom salts), &c.
Aluminium sulphates. The normal salt

Al₂(SO₄)₃. 18aq occurs native as feather alum; it is prepared by heating clay with conc. H,SO,Aq and boiling down the solution; also, according to Persoz (A. Ch. [8] 56, 102), by adding solution of alum or AlCl_s to a large excess of conc. boiling H₂SO₄, when anhydrous Al₂(SO₄), separates as a white powder. Gawalovski (C. O. 1885. 721) says that perfect octahedral crystals of Al₂(SO₄)₂. 17aq are obtained by saturating H₂SO₄Aq with freshly ppd. AlO₂H₂ and allowing to stand for some months. Crystallises with difficulty from water; crystallisation hastened by adding alcohol in which the salt is insol. (P., l.c.). Concerning ppn. of $Al_x(SO_4)_xxH_yO$ from aqueous solutions by H_2SO_4 v. Eremin, J. R. 20, 468 (abstract in C. J. 56, 347). When heated gives off all water; heated to redness gives off SO, and leaves Al.O. Various basic salts are known; they are obtained by heating solution of Al₂(SO₄), with AlO₂H₂, or by partially ppg. solution of the normal salt by NH₂Aq, or by partial reduction of the normal salt by Zn (v. Maus, P. 11, 80; Debray, Bl. [2] 7, 1; Athanasesco, C. R. 103, 271; Crum, A. 89, 156; Marguerite, C. R. 90, 1854; Löwe, J. pr. 79, 428; Bley, J. pr.

The double salts Al₂(SO₄)₃.M₂SO₄. 24aq, where M = NH₄, Na, K, Rb, Cs, Ag, or Tl, are alums. For expansions of alums v. Spring, B. 15, 1254, 1739; 17, 408.

Ammonia alum Al₂(SO₄)₂.(NH₄)₂SO₄. 24sq, prepared by adding (NH₄), SO₄ or NH₄Cl to Al. (SO₄), Aq, so that the salts are in equivalent quantities, crystallises in octahedra; S.G. 1.56;

S. 5.22 at 0°, 421.9 at 100°. Heated to 190°, 23H₂O is given off and Al₂(SO₄), (NH₄)₂SO₄, aq remains; NH₅ begins to come off at c. 193° (Lupton, C. J. [2] 18, 201).

Potash alum Al₂(SO₄)₃.K₂SO₄. 24aq, prepared by mixing solutions of the two sulphates and evaporating, crystallises in regular octahedra; S.G. 1.7; S. 3.29 at 0°, 9.52 at 10°, 22 at 30°, 81 at 60°, 90 at 70°, 857 at 100°. The crystals effloresce in air. At 190°, 23H₂O is given off (L., l.c.). Also obtained in cubical crystals (v. Polis, B. 13, 360). Solution in water has an acid reaction, and dissolves Zn and Fe, giving off H.

Cæsium and rubidium alums v. Bunsen, P. 119, 1; Godeffroy, B. 181, 176; Redtenbacher, J. pr. 95, 148. Setterberg (A. 211, 100) gives S. in water from 0° to 80° (cf. Cæsium,

vol. i. p. 658).

Silver alum v. Church a. Northcote, C. N. 9, 155.

Sodium alum v. Pouisin, Polytech. Centralbl. 1852.774; Zellner, S. 86, 183; and especially Augé, C. R. 110, 1139 (abstract in C. J.

58, 1059)

Thallium alum v. Lamy, Bl. [2] 11, 210. The double salts of Al₂(SO₄), with FeSO₄, MgSO₄, MnSO₄, and ZnSO₄ also crystallise with 24H₂O. For double salts with Fe₂(SO₄)₈, Mn₂(SO₄)₈, and Cr₂(SO₄)₈ v. Etard (BL [2] 31, 200). For a double salt with PbSO, v. Bailey (C. S. I. 6, 415).

Ammonium sulphates. The normal salt

(NH₄),SO₄ is found in certain volcanic districts; it is prepared by neutralising H₂SO₄Aq by NH₂Aq or (NH₄)₂CO₃Aq and evaporating. Crystallises in clear rhombic forms, is isomorphous with K₂SO₄ (Mitscherlich, P. 18, 168). S.G. 1.761 (Playfair a. Joule, C. S. Mem. 2, 401); for S.G. from 10° to 100° v. Spring (B. 15, 1940). Melts at c. 140° (Marchand, P. 42); decomposes above 280°; according to Johnson a. Chittenden (Am.S. [3] 15, 131), gives NH_4HSO_4 and $(NH_4)_2S_2O_7$, and at incipient redness gives SO, H,O, and NH,. Absorbs dry HCl, forming some NH₄Cl (v. Thomas, C. J. 83, 372). S. 71 at 0°, 73 65 at 10°, 76 3 at 20°, 78 95 at 30°, 81 6 at 40°, 84 25 at 50°, 86 9 at 60°, 89.55 at 70°, 92.2 at 80°, 94.85 at 90°, 97.5 at 100° (Alluard, C. R. 59, 500). Insol. absolute alcohol. For S.G. of (NH₄)₂SO₄Aq from 1 to 50 p.c. (NH₄)₂SO₄ v. Schiff (Å. 108, 338; 110, 74). H.F. [N²,H³,S,O⁴] = 284,800 (Thomsen, J. pr. 21, 477). The acid salt NH₄HSO₄ crystallises from solution of (NH₄)2SO₄ in hot conc. H₂SO₄ in long rhombic prisms, S.G. conc. 12,504 in long rhombic prisms, S.G. 1.787 (Schiff, A. 107, 83). S. 100 in cold water. Two other acid salts have been isolated: (1) (NH₄)₂H(SO₄)₂ [=8(NH₄)₂SO₄.SO₂.H₂O or 3(NH₄)₂O.4SO₂.H₂O] by adding the proper quantity of H₂SO₄ to (NH₄)₂SO₄Aq (Marignac, Ann. M. [5] 12, 38); (2) (NH₄)₂SO₄.7SO₂ [= (NH₄)₂O.8SO₂] by heating dry (NH₄)₂SO₄ with SO₂ to 100° in a closed tube and distilling off excess of SO₂ at c. 60° (Weber. B. 17, 2501). c. 60° (Weber, B. 17, 2501).

substantial and the common salt of the c

Sb₂(SO₄), is obtained by dissolving Sb₂O₂ in boiling conc. H₂SO₄Aq (c. 98 p.c. H₂SO₄), recrystallising from conc. H₂SO₄, and drying on a porous tile over H₂SO₄ (Adie, C. J. 57, 540; where an account of the action of H₂SO₄Aq of different concentrations, and of SO, on Sb,O, will be found, with references to older papers).

Barium sulphates. The normal salt BaSO. occurs native as heavy spar. Prepared by adding dilute H₂SO,Aq or dilute solution of a sulphate to solution of a salt of Ba, washing, and drying. A white solid; S.G. 4.525 (G. Rose, P. 75, 409; v. also Schröder, P. 106, 226; a. Wiedemann, P. M. [5] 15, 371). Obtained as a crystalline powder by fusing 1 part K2SO4 with 41 parts dry BaCl₂ in a closed crucible, and washing with water (Manross, A. 82, 848); also by heating dilute BaCl2Aq with slight excess of H₂SO₄Aq to 245° (Scheerer a. Drechsel, *J. pr.* [2] 7, 63). H.F. [Ba,O⁴,S] = c. 338,000 (*Th.* 3, 516). Insol. water; 1 part dissolves in c. 23,000 parts cold, and in c. 4,900 parts hot, HClAq S.G. 1.03, and in c. 9,200 parts HNO, Aq S.G. 1.02. Struve (Fr. 9, 34) gives S. of BaSO, in conc. H₂SO, as 5.69, and 15.89 in Nordhausen acid. BaSO, is partly decomposed by boiling with conc. solution of an alkali carbonate, or by fusion with alkali carbonate; Spring (Bl. [2] 44, 166) found that some BaCO, was formed by compressing BaSO, and Na₂CO₃ at a pressure of c. 6,000 atmospheres.

An acid salt BaSO₄.H₂SO₄ is formed by dissolving BaSO₄ in hot conc. H₂SO₄ S.G. 1.843, and heating to 100° (v. Garside, C. N. 31, 245).

Beryllium sulphates. The normal salt BeSO4. 4aq is formed by dissolving BeCO3 in dilute H2SO4Aq and concentrating the slightly acid liquid. For crystalline form v. Topsöe (W. A. B. 66 [2nd part], 5). According to Klatzo (J. pr. 106, 233) the salt crystallises with 7H₂O. Various basic salts are obtained by the action of BeCO₃ or Zn on solution of the normal salt.

Bismuth sulphates. The normal salt

Bi₂(SO₄)₃ is best prepared by dissolving Bi₂S₂ in conc. H,SO, heated to commencing vaporisation; on cooling the salt separates in lustrous, very deliquescent, needles (Hensgen, R. T. C. 4, 401; cf. Schultz-Sellack, B. 4, 13). For the action of heat on Bi.(SO₄), v. Bailey (C. J. 51, 680). An acid salt BiH(SO₄)₂, 3aq was obtained by Leist (A. 160, 29) by dissolving Bi₂O₃ in dil. H₂SO₄Aq under definite conditions. *Basic salts* are also formed by the reaction of dilute H2SO4Aq with Bi₂O₃ (v. Heintz, P. 63, 55, 567), and by the action of H₂O on the normal salt (cf. Hensgen, l.c.)

According to B. H. Adie (priv. comm.), acid salts BiH(SO₄)₂. xaq (x = probably $1\frac{1}{2}$ and 3) crystallise from solutions of Bi₂O₃ or Bi₂(SO₄)'s in H2SO, Aq of concentrations varying from H₂SO₄.H₂O to c. H₂SO₄.5H₂O; from acid of the concentration H₂SO₄.6H₂O, a basic salt crystal-

concentration 11204.SO4.H2O. lises, probably Bi.OH.SO4.H2O. The normal salt CdSO4. aq is obtained by dissolving Cd in dilute H₂SO₄Aq, and concentrating, by boiling, the acid solution (von Hauer, J. pr. 72, 372). other hydrates have been isolated. Various $[\text{Od}, \text{S}, \text{O}^4] = 221,550 \ (Th. 3, 516).$ CdSO, combines with NH₂ to form various compounds (v. H. Rose, P. 20, 152; Malaguti a. Sarzeau, A.Ch. [3] 9, 431; Isambert, C. R. 70, 456; Müller, A. 149, 70). Forms double salts with (NH₄)₂SO₄, MgSO₄, and Na₂SO₄; these salts crystallise with 6aq (von H., l.c.) Al₂(SO₄), is an alum. The double salt with

Cosium sulphates. Normal salt Cs.SO.; by neutralising H.SO.Aq by Cs.CO. and evapo-

rating. Short needles, not hygroscopic; insol. alcohol; S. 158.7 at -2°. The acid salt CsHSO., formed by reacting on Cs, SO, with excess of H2SO4, crystallises in small rhombic prisms

(Bunsen a. Kirchoff, P. 113, 342).

Calcium sulphates. The normal salt CaSO. occurs native as anhydrite, and the dihydrate CaSO, 2aq as gypsum, alabaster, and selenite. CaSO, is obtained in crystals by fusing K2SO, with excess of CaCl₂ and washing (Manross, A. 82, 348; cf. Mitscherlich, P. 21, 321). CaSO, 2aq is formed by ppg. fairly conc. CaCl2Aq by dil. H₂SO₄Aq or solution of a sulphate, evaporating, and washing the solid that separates. CaSO, 2aq is said to be formed by the interaction of CaCO, and S moistened with water (Polacci, G. 1874. 177, 245). CaSO, has S.G. 2.964, and CaSO, 2aq has S.G. 2.31 (v. also McCaleb, Am. 11, 35). S. for CaSO., ·19 at 0°, ·206 at 20°, ·214 at 40°, ·208 at 60°, ·195 at 80°, ·174 at 100° (Marignac, A. Ch. [5] 1, 274); S. for CaSO, 2aq 2119 at 16.5°, 2352 at 22° (Cossa, G. 1873. 135); S. for CaSO, 2aq in glycerin, 957 at ordinary temperature, S. increases as temperature rises (Asselin, C. R. 76, 884). According to Raupenstauch (C. C. 1888. 821) the solubility in water of gypsum increases to 32°, is constant from 32° to 38°, and decreases above 38°. Gypsum loses 2H2O when heated to 80° in a stream of dry air; when the CaSO. thus produced is moistened it takes up 2H2O, becomes crystalline, and expands. The dehydrated CaSO, obtained by heating gypsum to 160° takes up water very slowly; if the gypsum is heated to c. 300° the CaSO, formed combines with water with extreme slowness (v. Schott, D. P. J. 202, 52, 355, 513; cf. Plaster of Paris, under CEMENTS, in DICTIONARY OF APPLIED CHEMISTRY, vol. i. p. 468).

According to Potilitzin (J. R. 1893 [1] 201, 207) CaSO₄. 2aq slowly gives off water at 62°-65° till the hydrate 2CaSO₄, aq is formed; CaSO₄ prepared by heating CaSO₄, 2aq to 130°-170° absorbs H2O from ordinary air until 2CaSO. aq is formed when absorption of water ceases. The hydrate 2CaSO4.aq absorbs water from air saturated with moisture, forming CaSO4. 2aq. The water of crystallisation of gypsum is not equally firmly retained; P. writes the formula 2CaSO, 4aq; three-quarters of the H2O is much more readily given off than the other fourth. The hydrate 2CaSO₄, aq is more soluble in water

than the dihydrate CaSO₄. 2aq.
An acid salt CaH₂(SO₄)₂ is said by Berzelius to be formed by digesting CaSO, with H2SO, Aq at 80°-100°; it is decomposed by moist air to the normal salt and H2SO4. For double salts with alkali and alkaline earth sulphates v. Fassbender (B. 9, 1858; 11, 1968); Struve (Bl. [2] 12, 848); Ditte (C. R. 84, 86); Popp (A. Suppl. 8, 1); Hannay (C. J. 32, 399).

Cerium sulphates, The normal cerous sulphate, Ce₂(SO₂)₂.xH₂O (x being probably 6, 8, 9, and 12) is obtained in colourless crystals by evaporating a solution of Ce₂(CO₂)₃ in dilute H₂SO₄Aq, or a solution of CeO₂ in H₂SO₄Aq saturated with SO₂. The normal ceric salt Ce(SO₄)₂. 4aq forms, in yellow crystals, from a solution of CeO₂ in dilute H₂SO₄Aq. Cerosoceric sulphates, xCe₂(SO₄)₂·yCe(SO₄)₂·2aq, are obtained by dissolving CeO₂ in conc. H₂SO₄, and evaporating. Basic salts are formed by the interaction of water with the normal salts. Double salts of Ce2(SO4)3 with (NH4)2SO4, K2SO4, and Na₂SO, are known (v. Marignac, A. Ch. [4] 30, 57; Czudnowicz, J. pr. 30, 19; Hermann, J. pr. 92, 126; Zschiesche, J. pr. 107, 65; Rummelsberg, B. 6, 85; Jolin, Bl. [2] 21, 523; Wyrouboff, Bl. [3] 2, 745; Wing, Am. S. [2] 40, 856).

Chromium sulphates. The normal chromous salt CrSO, 7aq was obtained by Moissan (Bl. [2] 37, 296) by dissolving the acetate in warm dilute H₂SO₄Aq, crystallising, and drying between paper, all operations being conducted in an atmosphere of CO2 (for directions for making Cr(C₂H₂O₂)₂ v. Nitrogen, Preparation, vol. iii. p. 557). Blue crystals; isomorphous with FeSO, 7aq. S. 12:35 at 0°; sl. sol. alcohol. Absorbs O rapidly from air, and acts as a strong reducer. A monohydrate CrSO4 aq was obtained as a white crystalline powder by adding the acetate to excess of conc. H₂SO₄ (M., l.c.). Forms a double salt, CrSO₄.K₂SO₄. 6aq (Péligot, A. Ch.

[8] 12, 539). The normal chromic salt $Cr_2(SO_4)_3$ xH_2O exists in two forms. (1) Violet, x = 15 (Schrötter, P. 53, 516), = 18 (Siewert, A. 126, 97); by digesting 1 pt. CrO_3H_3 dried at 100° with 1 to 14 pts. conc. H₂SO₄, in a closed vessel, till a mass of greenish-blue crystals is formed, dissolving these crystals in water, ppg. by absolute alcohol, dissolving again in water, adding alcohol till ppn. just begins, covering the vessel with moist parchment-paper, and allowing to crystallise slowly. Regular octahedra; red-violet in reflected light, garnet-red in transmitted light; S. 120 in cold water (Schrötter, P. 53, 516; Löwel, A. Ch. [3] 40, 42). (2) Green, x=5. By heating the violet salt to 100° ; the salt melts, gives off water, and then solidifies to a green amorphous mass. Also formed by dissolving CrO₃H₃ in conc. H₂SO₄ at 50°-60°, and evaporating quickly. Easily sol. alcohol; whereas the violet salt is insol. alcohol.

Addition of a cold solution of a Ba salt ppts. all SO, from a solution of the violet form, but boiling is needed to complete the ppn. from the According to Etard (C. R. 84, green variety. 1089), the difference between the two forms is one of hydration only (but v. Chromium salts, vol. ii. p. 167; and v. also Recoura, C. R. 113, 857).

Basic salts are obtained by adding a little NH3Aq to solution of the normal salt, also by digesting the normal salt with CrO₃H₃, or excess of Cro.H., with H.SO.Aq (v. Schrötter, P. 53, 516; Siewert, A. 126, 97; Schiff, A. 124, 167, 172; Recoura, C. R. 112, 1439). According to Traube (A. 66, 87), an acid salt, Cr₂(SO₄)₃.H₂SO₄, is formed by heating either variety of the normal salt with excess of H2SO4 till vaporisation of the acid begins, and allowing to cool (Schrötter [l.c.] regarded this as an insoluble variety of the normal salt); insol. water, not acted on by cold KOHAq, heated strongly gives Cr₂O₂, SO₂, and O.

Cr₂(SO₄)₃ forms many double salts; those with the alkali sulphates have the composition

Cr₂(SO₄)₃,M₂SO₄,24aq, and are alums.
Ammonia chrome-alum is ppd. by adding (NH₄)₂SO₄ to a saturated solution of violet Cr₂(SO₄)₃; also formed by mixing (NH₄)₂SO₄Aq and Cr₂(SO₄)₂Aq, and evaporating; also by mixing Cr₂(SO₄)₂Aq (violet) with (NH₄)₂Cr₂Aq,

adding a little H₂SO₄, and allowing to evaporate. Crystallises in ruby-red octahedra; S.G. 1.738. Effloresces slowly in air. Solution in water is violet; at 75° it becomes green, but slowly returns to violet on standing for some weeks (regarding this change v. CHROMIUM SALTS, vol. ii. p. 167).

Potash chrome-alum is produced similarly to the ammonia salt; also by heating K2Cr2O, with conc. H2SO4, more quickly in presence of a reducer, e.g. SO₂. Crystallises in large dark-purple octahedra; S. 14.5 in cold water. Solution in water is grey-blue with tinge of red; heated to 70°-80° it becomes green and noncrystallisable, but returns to violet after standing for some weeks. For Soda chrome-alum v. Schrötter (P. 53, 513). For double salts of $Cr_2(SO_4)_8$ with $Al_2(SO_4)_3$, $Fe_2(SO_4)_3$, and $Mn_2(SO_4)_8$ v. Etard (Bl.~[2]~31,~200); for double salts of the form $Cr_2(SO_4)_3$.3M, SO_4 , where M=Li, K, or Na, v. Warnicke (P. 159, 572).

For description of other double salts of Cr₂(SO₄)₈ with alkali sulphates, v. Klobb, C. R.

117, 311 (Abstract in C. J. 64 [ii.], 573).

Cobalt sulphates .- The normal cobaltous salt CoSO, 7aq occurs native as biberite; prepared by dissolving Co, CoO, or CoCO₃ in dilute H₂SO₄Aq, and evaporating. Forms carmine-red crystals, isomorphous with FeSO₄, 7aq; S.G. 1.918 at 15° (Thorpe a. Watts, C. J. 37, 102). Unchanged in air; loses all water by heating, and gives CoSO, as a red powder, not decomposed easily by heat; S.G. 3.472 at 15° (T. a. W., l.c.). Klobb (C. R. 114, 836) obtained the anhydrous salt in crystals by mixing CoSO4.7aq with excess of (NH₄)₂SO₄ and heating in a partly-closed crucible, out of contact with the gases of the flame, till (NH₄), SO₄ was all volatilised. The hexahydrate CoSO₄, 6aq separates from solutions of Co or CoCO₃ in H₂SO₄Aq at 40°-50°; it is isomorphous with the corresponding salts of Mg, Ni, and Zn (Marignac). For other hydrates, with 4aq and aq, v. Fröhde (J. 1866. 244); Vortmann (B. 15, 1888). CoSO, 7aq is insol. alcohol; S. water 30.5 at 10°, 36.4 at 20°, 40 at 29°, 46.3 at 35°, 55.2 at 50°, 60.4 at 60°, 65.7 at 70° (stated as parts CoSO, dissolved by 100 parts water; Tobler, A. 95, 193). Thomsen (Th. 3, 516) gives H.F. [Co,S,O',7H,O] = 234,000. By electrolysing CoSO, Aq in presence of K, SO, and H2SO4, under special conditions, Marshall obtained Co₂(SO₄)₃ along with K₂S₂O₈ (C. J. 59,

Basic salts are formed by addition of a little NH, Aq to hot CoSO, Aq, also by digesting CoSO, Aq with CoCO, (v. Habermann, M. 5, 442; Athanasesco, C. R. 103, 271).

For double salts with FeSO4, MgSO4, MnSO4,

and ZnSO, v. Vohl (A. 94, 57).

Normal cobaltic sulphate Co2(SO4)3 was obtained by Marshall (C. J. 59, 767) by electrolysing CoSO₄Aq in presence of H₂SO₄, under special conditions. A blue crystalline powder, unchanged in a stoppered bottle in ordinary air, but decomposes rapidly in dry air; soi. water, with evolution of O; sol. conc. H2SO4; in HClAq forms CoCl, and gives off Cl.

Copper sulphates. Only salts corresponding with the oxide CuO have been isolated. The normal salt CuSO, is obtained by dissolving Cu in hot conc. H2SO4, decanting the hot solution, and allowing to cool; also by adding conc. H₂SO₄ to CuSO₄. aq; also by heating powdered | crystals of CuSO₄. 5aq to c. 230° as long as they lose weight. Klobb (C. R. 114, 836) obtained CuSO, in crystals by heating a mixture of the by hydrated salt and excess of $(NH_4)_2SO_4$ in a partly closed crucible till all $(NH_4)_2SO_4$ had volatilised. A white crystalline powder; S.G. 8:606 at 15° (Thorpe a. Watts, C. J. 37, 102). Partly decomposed at red heat, wholly at white heat, to SO₂, O, and CuO. Reduced by heating to redness with charcoal, giving Cu, SO2, and CO2. At a higher temperature some CuS is said to be formed; also reduced to Cu by heating in H; heated in PH, gives off H2O, and leaves a mixture of sulphide and phosphide of Cu. HCl gas forms CuCl₂ and H₂SO₄. CuSO₄ acts as a powerful dehydrating agent; it rapidly absorbs water, forming CuSO .. 5aq. H.F.[Cu,S,O'] =182,500 (Th. 3, 516).

The pentahydrate CuSO. 5aq (blue vitriol) is prepared by dissolving Cu in hot conc. H.SO., evaporating, and recrystallising from water the solid that separates. At. c. 20° there is a very slight reaction, with formation of Cu₂S and Cu₃O₄; at c. 270° the reaction is Cu+2H₂SO₄ $= \text{CuSO}_4 + \text{SO}_2 + 2\text{H}_2\text{O}$ (Pickering, C. J. 33, 112). According to Anthion (R. P. 81, 344), CuSO .. 5aq free from nitrate, is obtained by dissolving Cu in a warm mixture of dilute H2SO, Aq and HNO2Aq, and crystallising. (For preparation of commercial blue vitriol v. DICTIONARY OF APPLIED CHEMISTRY, vol. i. p. 608.) Large blue, triclinic crystals; S.G. 2.284 at 15° (Thorpe a. Watts, C. J. 37, 102). Solubility in water is given by Poggiale (A. Ch. [3] 8, 463) as follows:—

	S.		1	s.		
Temp.	CuSO. Saq	CuSO.	Temp.	CuSO. 5aq	OuSO.	
0°	81.61	18.20	60°	77.39	38.83	
10	36.95	20.92	70	$94 \cdot 40$	45.06	
20	42.31	23.55	80	118.03	53.15	
30	48.81	26.63	90	156.44	$64 \cdot 23$	
40	56.90	80.29	100	203.32	75.25	
50	65.83	34.14				

Gerlach (D. P. J. 181, 129) gives the following table :-

Percentage	8.G.	Percentage	s.G.
CuSO. 5aq		CuSO. 5aq	
2.	1.0126	14 .	1.0933
4 .	1.0254	16 .	1.1063
6.	1.0384	18 .	1.1208
8.	1.0516	20 .	1.1354
10 .	1.0649	22 .	1.1501
12 .	1.0785	24 .	1.1659

Insol. absolute alcohol; S. in 40 p.c. alcohol at 15°= 25 (Schiff). Fairly sol. glycerin. Effloresces in dry air, giving off 2H₂O; heated for some time to 100° loses 4H₂O, the last H₂O is removed at c. 230°. For dissociation-pressures of water given off at various temperatures v. Lescour (C. R. 102, 1466). Solution in HClAq gives CuCl, 2H,O on evaporation. The dry salt, in powder, absorbs HCl, the water of crystallisation being set free (Kane, A. 19, 1); several compounds of CuSO4, HOl, and H2O are formed according to Latschinoff (J. R. 1888 [1] 586, 657, 707; abstract in B. 22 Ref. 192). Combines with NH, forming CuSO, 5NH, all water of orystallisation being removed (L., l.c.).

Other hydrated copper sulphates are (1)

CuSO₄.6aq and (2) CuSO₄.7aq, obtained by Lecoq de Boisbaudran (C. R. 65, 1240), the former by placing a crystal of NiSO₄.4aq in a warm supersaturated solution of CuSO, 5aq, to which a few drops of H2SO4Aq had been added, and the latter by placing a crystal of FeSO, 7aq in the mother-liquor from the first-named hydrate; (3) CuSO₄. 3aq, by keeping CuSO₄. 5aq in dry air at 25° – 30° (Magnier de la Source, C. R. 83, 899); also by pouring cold conc. CuSO, Aq into H2SO, Aq, S.G. 1.7, and washing the pp. with small quantities of absolute alcohol (Thorpe a. Watts, C.J. 37, 104); (4) CuSO₄. 2aq by keeping the pentahydrate in vacuo at 20° for some days, or by pouring conc. CuSO, Aq into cold conc. H₂SO₄, and washing the pp. with absolute alcohol (T. a. W., l.c.); (5) CuSO₄, and by heating powdered pentahydrate at 110° till it ceases to lose weight (T. a. W., l.c.). T. a. W. (l.c.) give the S.G. of these hydrates, at 15°, as: monohydrate, 3.289; dihydrate, 2.953; trihydrate, 2.663. Lescœur (C. R. 102, 1466) obtained only the penta-, tri-, and monohydrates.

Double salts with the alkali sulphates, of the form CuSO₄. M₂SO₄. 6aq, are obtained by crystallising mixtures of the constituents in the proper proportions (v. Graham, P. M. 1835. 327, 417; Brunner, P. 15, 476; 32, 221; Pickering, C. J. 49, 1). Vohl (A. 94, 57) has described several triple salts of the form CuSO4.MSO, 2X2SO, 12aq, where M = Fe, Zn, &c., and X = an alkali metal; and also quadruple salts CuSO, 2MSO, 3X, SO, 18aq. Double salts of the form CuSO, MSO, xaq, x being 5 and 7, and M being Co, Fe, Mg, Ni, or Zn, are described (v. Rammelsberg, P. 91, 321; also Lefort, A. Ch. [3] 23, 95; von Hauer, P. 125, 638). Several basic salts of the form CuSO₄.xCuO. aq are known: -x=1, v. Roucher (J. Ph. [3] 37, 249); x = 2, v. Reindel (J. pr. 100, 1; 102, 204), Persoz (A. Ch. [3] 25, 257), Shenstone (C. J. 47, 375); x = 3, v. Smith (J. 25, 280), Pickering (C. J. 43, 336; 49, 1), Grimbert a. Barré (J. Ph. [5] 21, 414), Kuhn (Ph. C. 1847. 595), Kane (A. 19, 1); x=4, v. Smith (l.c.); x=7, v. Pickering (l.c.). For other basic sulphates v. Steinmann (B. 15, 1411), Brunner (P. 15, 476; 32, 221), Becquerel (C. R. 67, 1081), Casselmann (Fr. 4, 24).

Didymium sulphates (v. Marignac, A. Ch. [3] 58, 148; Cleve, Bl. [2] 21, 246; 39, 151; Hermann, J. pr. 82, 385; Frerichs a. Smith, A. 191, 348). The normal salt Di₂(SO₄)₃. 8aq crystallises in red, lustrous, monoclinic crystals from a solution of Di₂O₃ or Di₂(CO₃), in excess of H.SO.Aq; S.G. 2.878; S.H. 1948 (Nilson a. Pettersson, B. 13, 1459). Heated to 300° loses all water of crystallisation, leaving Di₂(SO₄)₃ as a red powder; S.G. 3.735; S.H. 1187 (N. a. P., l.c.); S. 43.1 at 12°, 11 at 50°, 2 at 100° (Marignac). For effect of heat at various temperatures v. Bailey (C. J. 51, 682). Hydrates with 6aq and aq crystallise from cold saturated solutions of Di₂(SO₄)₃; Marignac (l.c.) also obtained a hydrate with 9aq. Double salts of the form Di₂(SO₄)₃.M₂SO₄. Saq, where M is an alkali metal, are known. Basic salts are formed by heating the normal salt, also by adding a little NH2Aq to solutions of the normal salt.

The normal auro-auric Gold sulphates. sulphate AuSO, was obtained by Schottländer (A. 217, 312) by evaporating solution of AuO.H.SO, (v. infra) nearly to dryness at 250°, and drying in a desiccator over lime. Lustrous, scarlet prisms, that rapidly absorb moisture from the air, with partial decomposition. A basic salt AuO.H.SO, (auryl sulphate) was obtained (S., l.c.), as a yellow crystalline powder, by heating AuO.NO, with conc. H₂SO, at c. 200° (AuO.NO, was formed by dissolving AuO₃H₃ in HNO₃Aq S.G. 1·4, filtering through asbestos, and evaporating under reduced pressure over lime and soda). A double salt AuK(SO₄)₂ was formed by dissolving KHSO₄ in solution of AuO.H.SO₄, and evaporating at 200° (v. also Allen, C. N. 25, 85).

Indium sulphates (v. Winkler, J. pr. 94, 1; 95, 414; 98, 844; 102, 273; R. Meyer, A. 150, 429). The normal salt In₂(SO₄)₃ is obtained by dissolving In or InO₈H₃ in excess of H₂SO₄Aq, evaporating to dryness, and heating till excess of acid is removed; a white, very hygroscopic powder; S.H. ·129 (Nilson a. Pettersson, B. 13, 1459). S.G. 3.438 (N. a. P., C. R. 91, 232). By dissolving in water and evaporating, a hydrate with 9aq is obtained as a gummy solid. The solution of In or InO, H, in excess of H2SO, Aq yields a syrupy liquid on evaporation which probably contains an acid salt (? In₂(SO₄)₂, H₂SO₄). By adding (NH₄)₂SO₄Aq to solution of In₂(SO₄)₂, and evaporating, the double salt with the composition In₂(SO₄)₂.(NH₄)₂SO₄.24aq is obtained; this salt is an alum; it is sol. in $\frac{1}{3}$ part water at 16°, and in $\frac{1}{4}$ part water at 30°. On evaporating the mother-liquor, the salt In₂(SO₄)₂.(NH₄)₂SO₄. 8aq is obtained (Rössler, J. pr. [2] 7, 13). Double salts with K_2SO_4 and Na_2SO_4 , orystallising with 8aq, have also been obtained.

Iridium sulphates. By treating IrS₂ with conc. HNO₃Aq, Birnbaum (A. 136, 179) obtained an amorphous, brownish-yellow solid, easily sol. water, probably a sulphate of Ir. The double salt Ir₂(SO₄)₂.3k₂SO₄ was obtained by Lecoq de Boisbaudran (C. R. 96, 1406, 1551) by fusing KHSO₄ with Ir compounds, treating with water containing KHSO₄, dissolving the residue in hot water, and crystallising; green, translucent crystals, probably regular octahedra; almost insol. saturated KHSO₄Aq.

Iron sulphates. I. Ferrous salts. The

normal salt, FeSO4, is obtained by heating powdered FeSO4. 7aq very carefully to 800° till it ceases to lose weight (the heating is best done in a stream of H); at a slightly higher temperature SO_2 and O begin to be given off, and finally Fe₂O₂ remains. FeSO₄ is a white powder; S.G. 3.346 (Thorpe a. Watts, C. J. 37, 102). Several hydrates of FeSO, are known, ordinary green vitriol being the most common and best examined of these; this heptahydrate, FeSO, 7aq, is best prepared by adding iron filings to warm dilute H2SO4Aq until the acid is saturated and a little iron remains undissolved, boiling for a little, filtering into a vessel that has been rinsed with a very little conc. H₂SO₄, allowing to crystallise, washing the crystals two or three times with very small quantities of cold water, and then with alcohol till all acid is removed, drying at c. 80°, powdering the dry crystals, and strongly pressing between paper or linen. As thus prepared the salt does not readily oxidise in the air. If alcohol is added to the hot solution, filtered from undissolved iron, FeSO. 7aq is ppd. in minute crystals; if this pp. is washed with alcohol till acid is removed and then spread on filter paper, in dry air, till the alcohol has evaporated, the salt is obtained as a bluish white powder, which does not oxidise except in very moist air. Blue-green, monoclinic crystals; S.G. 1.8889 at 4° (Playfair a. Joule, C. J. 1, 138), 1.9854 at 16° (Pape, P. 120, 372). H.F. [Fe,S,O',7H²O] = 240,100 (Th. 3, 516). Heated to 100° gives off 6H₂O, and at c. 300° leaves FeSO. (v. supra). S. 60.9 at 10°, 70 at 15°, 114.95 at 25°, 227.27 at 46°, 263.15 at 60°, 270.27 at 84°, 370.37 at 90°, 333.33 at 100°. Gerlach (D. P. J. 181, 129) gives the following data (v. also Schiff, D. P. J. 1858. 38):—

.G. FeSO.Aq	Potge. FeSO., 7aq	Pctge, FeSO
1.0267	5	2.811
1.0537	10	5.784
1.0823	15	8.934
1.1124	20	12.277
1.1430	25	15.834
1.1738	30	19.622
1.2063	35	23.672
1.2391	40	27.995

Insol. glacial acetic acid and absolute alcohol. FeSO4. 7aq is fairly easily oxidised; exposure to air produces basic ferric sulphates. FeSO, Aq absorbs NO, forming FeSO, xNO (x probably = 1 or \frac{1}{5}; v. Gay, A. Ch. [6] 5, 145, and cf. Nitric oxide, vol. iii. p. 563). Other hydrates of ferrous sulphate are (1) FeSO4. 4aq, formed by crystallising FeSO, Aq at 80° (Regnault, A. Ch. [3] 1, 201), also by exposing finely-powdered FeSO. 7aq over H.SO., in an atmosphere of CO., until it ceases to lose weight (Thorpe a. Watts, C. J. 37, 102); S.G. 2.227 (T. a. W.); (2) FeSO. 3aq, by evaporating FeSO. Aq made strongly acid by H₂SO₄ (Kühn, S. 61, 235), also by dissolving FeSO4. 7aq in hot HClAq and crystallising (Kane, A. 19, 7); (3) FeSO, 2aq by boiling the heptahydrate with successive quantities of alcohol; S.G. 2.773 (T. a. W., l.c.); (4) FeSO, aq, by heating powdered FeSO, 7aq to 120° in H; S.G. 2.994 (T. a. W.; v. also Mitscherlich, P. 18, 152). Several double salts of ferrous sulphate are known; they are generally prepared by crystallising a mixture of the constituent sulphates in the proper proportions. The salts $FeSO_4.M_2SO_4.6aq$, where $M=NH_4$, K, Na, or Tl, form blue-green monoclinic crystals (v. Marignac, Ann. M. [5] 9, 19; Schiff, A. 108, 326; Tobler, A. 95, 193; Wilm, A. Ch. [4] 5, 56). For the salts FeSO, MSO, 14aq, where M=Mg or Zn, v. Schiff (l.c.), and Murmann (W. A. B. 27, 172). For various triple and quadruple salts containing FeSO₄, alkali sulphates, and sulphates of Mn, Mg, Zn, &c., v. Bette (A. 14, 278) and Vohl (A. 94, 57).

II. Ferric salts. The normal salt Fe₂(SO₄)₃ is obtained by adding 1 part conc. H₂SO₄ to 5 parts FeSO₄. 7aq dissolved in water, warming, and adding HNO₃Aq little by little till oxidation is complete, evaporating to a syrup, and then heating very cautiously so long as acid fumes are given off; on cooling Fe₂(SO₄)₃ remains as a white amorphous solid; S.G. 3.097 at 18° (Pettersson, Upsala, Nova acta, 1874). Fairly large quantities of FeSO₄. 7aq should be used, as evaporation of a dilute solution containing Fe₂(SO₄)₂ produces ppn. of basic sulphates. The hydrate Fe₂(SO₄)₃, 9aq occurs native as

coquimbite in Chili. On one occasion Bertels obtained rhombic leaflets of a hydrate with 10aq (J. 1874. 268). Fe₂(SO₄)₃ is slowly sol. in water; the presence of a very small quantity of FeSO. causes Fe₂(SO₄), to dissolve rapidly in water (Barreswil, *Ph. C.* 1845. 604). Fe₂(SO₄), is insol. in conc. H₂SO₄; it is reduced to FeSO₄ by most metals, more quickly if a little H2SO4 is present. Fe₂(SO₄)₃ forms several double salts. The compounds $Fe_2(SO_4)_3$. M_2SO_4 . 24aq, where $M = NH_4$ or K, are alums. Potash iron alum Fe₂(SO₄)₄,K₂SO₄. 24aq is prepared by dissolving 28 parts FeSO₄. 7aq in water, adding 5 parts conc. H₂SO₄, oxidising by HNO₃Aq, adding 8.75 parts K₂SO₄, and crystallising; or by mixing 10 parts powdered FeSO. 7aq with 4 parts KNO₃, slowly adding 5 parts conc. H₂SO₄, warming so long as NO₂ is given off, dissolving the residue in 4 times its weight of water at 80°, and allowing to stand at 0° for some time (Heintz, A. 44, 271). The salt slowly decomposes in the air; heated to 80° it gives Fe₂(SO₄), basic ferric sulphate, and H2SO4; a similar decomposition occurs when an aqueous solution of the salt is heated (Rose, P. 94, 459). Addition of KOHAq to a conc. solution of the salt till a slight permanent pp. is formed, followed by evaporation, produces a double salt of K2SO4 with a basic ferric sulphate (v. Richter a. Scheerer, P. 87, 73). Ammonia iron alum is formed similarly to the K salt; loses 23H₂O at 150° (v. Lupton, C. J. 28, 201). For double salts with Al₂(SO₄)₃, $Cr_2(SO_4)_3$, $Mn_2(SO_4)_3$ and $MnSO_4$, v. Etard (Bl. [2] 31, 200). Some double compounds of Fe₂(SO₄)₂ with (NH₄)₂SO₄, and with FeSO₄ and (NH₄)₂SO₄, are described by Lachaud a. Lepierre (C. R. 114, 915).

A great many basic salts have been described; generally prepared by the action of air on solutions of FeSO₄, 7aq, or by heating Fe₂(SO₄)₃Aq in sealed tubes (v. Athanasesco, C. R. 103, 271). Pickering (C. J. 37, 807) gives very full references to all the memoirs on those basic salts, and comes to the conclusion that only one exists, viz. 2Fe₂O₈SO₈; in a later paper (C. J. 43, 182) he gives reasons for writing the formula rather as 6Fe₂O₈SO₈.xH₂O₈ or Fe₂(SO₈)₈Fe₂O₈xH₂O.

6Fe₂O₃.3SO₃.xH₂O₄ or Fe₂(SO₄)₃.5Fe₂O₃.xH₂O₅ in Lanthanum sulphates. The normal salt La₂(SO₄)₃ is obtained by dissolving La₂O₅ in excess of H₂SO₄Aq, evaporating, drying the crystals of La₂(SO₄)₃, aq that separate, and heating to dull redness; a white powder, S.G. 3·6 (Nilson a. Pettersson, C. R. 91, 232); S.H. ·1182 (N. a. P., B. 13, 1459). Much more sol. cold than hot water; Mosander (P. 11, 406) gives S. 16·6 at 2°-8°, 2·35 at 25°, ·87 at 100°. The hydrate with 9aq, formed by the slow evaporation of La₂O₃ in excess of H₂SO₄Aq, crystallises in large, lustrous, hexagonal prisms (Topsoë); S.G. 2·853 (N. a. P., C. R. 91, 232). A hydrate with 6aq was obtained by Frerichs a. Smith (A. 191, 331). Several double salts with the alkali sulphates have been obtained; the principal are M.(NH₄)₂SO₄. 8aq, M.3K₂SO₄, and M.4K₂SO₄, where M = La₂(SO₄)₃ (v. Cleve, Bl. [2] 39, 151; 43, 56). A basic salt was obtained by F. a. S. (l.c. p. 860).

Lead sulphates. The normal salt PbSO₄ occurs in various lead ores. It is obtained as a white, heavy powder, by heating Pb with conc. H₂SO₄, by the interaction of warm H₂SO₄Aq

and PbO, or of PbO₂ and SO₂, and by ppg. solutions of lead salts by dilute H₂SO₄Aq or solution of a sulphate. PbSO4 is prepared in small rhombic crystals by melting PbCl2 with K2SO4, and washing with water. S.G. 6.2 (Schröder, P. Ergänz. Bd. 6, 622); 5.96 at 17° (Pettersson, Upsala, Nova Acta, 1874). H.F.[Pb,S,O'] = 216,200 (Th. 3, 516). Nearly insol. water; Rodwell (C. N. 11, 50) says that 31,062 parts water at 15° dissolves 1 part PbSO. Struve (Fr. 9, 34) gives S. in conc. H₂SO, as 13, and S. in Nordhausen acid as 4·19 (v. also Kolb, D. P. J. 209, 268). For solubility in dilute HClAq and HNO, Aq v. Rodwell (C. J. 15, 59); for S. in various salt solutions v. Löwe (J. 1859. 685); Staedel (J. 1863. 245); Storer (C. N. 21, 17); Dibbits (Fr. 1874. 137). Heated to redness, PbSO4 melts without decomposition, but at a white heat it gives PbO, SO2, and O (Boussingault, C. R. 64, 1159). Heated in a stream of H, it is reduced to Pb with evolution of SO, H₂S, and steam; heated with charcoal, PbO, PbS, or Pb is formed according to the proportions of the reacting substances. Reduced by Fe or Zn in contact with water. Rubbed with K2CrO4 produces PbCrO4; with KIAq yields PbI₂, and with NaCl is partly decomposed to PbCl₂ (v. Matthey, Ar. Ph. [3] 13, 233). For reaction when heated in NH₃ v. Rodwell (C. N. 15, 137). A double salt PbSO₄.(NH₄)₂SO₄ is formed by adding excess of H₂SO₄Aq to Pb(C₂H₃O₂)₂Aq, neutralising excess of acid by NH3Aq, boiling, and allowing to cool. An acid salt PbSO₄.H₂SO₄. aq was obtained by Schultz (P. 133, 137) by dissolving PbSO₄ in conc. H₂SO₄ and setting aside. A basic salt PbSO₄ PbO is formed by digesting PbSO₄ with NH, Aq (Kühn, Ar. Ph. [2] 50, 281). Lithium sulphates. The normal salt Li₂SO,

is obtained by dissolving Li₂CO₈ in H₂SO₄Aq, evaporating, drying the crystals of Li₂SO, aq that separate, and heating them to c. 135°. S.G. 2.21 at 15° (Brauner, P. M. [5] 11, 67). Melts at 818° (Carnelley, C. J. 33, 280). S. 85.34 at 0°, 34·36 at 20°, 32·8 at 45°, 30·3 at 65°, 29·24 at 100° (Kremers, A, 99, 47). Easily sol. alcohol; insol. SO, (Weber). H.F.[Li²,S,O¹] = 338,200 (Th. 3, 516). Heated in HCl gas below redness gives LiCl (Hensgen, B. 8, 125). The mono-hydrate Li₂SO, aq crystallises in monoclinic forms (Rammelsberg, P. 128, 311). S.G. 2.052 at 21° (Pettersson, Upsala, Nova Acta, 1874). Effloresces a little in air, gives up water at a little above 130°. An acid salt LiHSO, is obtained by dissolving Li₂SO, in H₂SO,Aq S.G. 1.6 to 1.7 (Schultz, P. 180, 149); prismatic crystals, melting at c. 120° (Lescœur, Bl. [2] 24, 516). Double salts with K₂SO, are obtained by evaporating mixed solutions of the constituent salts (v. Rammelsberg, A. 128, 311; Schabus, J. 1854. 323). For double salts with Rb₂SO₄ and Na₂SO₄ v. Wyrouboff (Bull. soc. mineral. de France, 1882. part 2). Wernicke (P. 159, 576) describes a salt 3Li₂SO₄.Cr₂(SO₄), formed, in green needles, by adding Cr2Os to molten LiHSO. Li2SO4 does not form alums, nor does it combine with the magnesian sulphates (Rammelsberg, l.c.; Scheibler, J. pr. 67, 485).

Magnesium sulphates. The normal salf MgSO, is obtained by dissolving MgO or MgCO, in H₂SO₄Aq (or by heating dolomite till CO₂ is

removed, treating with HClAq, which dissolves CaO, and dissolving the residual MgO in H.8O.Aq), evaporating, crystallising MgSO₄. 7aq, drying this salt, and heating it gradually to c. 300° till it ceases to lose weight. A white solid; S.G. 2.709 at 15° (Thorpe a. Watts, C. J. 37, 102); 2.77 to 2.795 at 14° (Pettersson, Upsala, Nova Acta, 1876). H.F. [Mg,S,O⁴] = 302,300 (Th. 3, 516). [MgSO',Aq] = 20,765 (Pickering, C. J. 47, 100). Decomposed to MgO, SO₂, and O at c. 360° (v. Bailey, C. J. 51, 682). Not acted on by HCl gas below incipient red heat (Hensgen, B. 9, 1671; 10, 259). The heptahydrate MgSO. 7aq (Epsom salts) occurs in many mineral springs, in sea water, and in some rocks. It crystallises at the ordinary temperature from solutions of MgO or Translucent, rhombic MgCO₃ in H₂SO₄Aq. prisms; from supersaturated solutions crystallises in hexagonal forms, and from solutions containing some FeSO, in monoclinic forms.(v. Marignac, Ann. M. [5] 12, 50). S.G. 1.678 at 15° (Thorpe a. Watts, C. J. 37, 102); 1:683 at 4° (Playfair a. Joule, C. J. 1, 138). H.F. [Mg, S, O⁴, 7H²O] = 326,400 (Th. 3,516). Melts at 70° (Tilden, C.J. 45, 267); gives up $6H_2O$ by prolonged heating at 150° - 160° , and all water of crystallisation at c. 280° (v. Pickering, C. J. 47, 100). The following data for solubility in water are calculated by Mulder from observations by Gay-Lussac, Tobler, and others :-

Temp.	ŝ.	Temp.	s.
0ô	25.76	60°	55.0
5	29.3	65	57·3
10	31·5	70	59.6
15	33.8	75	61.9
20	86.2	80	64.2
25	38.5	85	66.5
30	40.9	90	68.9
35	43.3	95	71.4
40	45.6	100	73.8
45	48.0	105	76.2
50	50.3	108.4	77.9
55	52.7		

Gerlach (Fr. 8, 287) gives the following:-

	, , , ,			
P.c. MgSO.	S.G. MgSO.Aq	P.c. MgSO.	S.G. MgSO,Aq	
1	1.01031	14	1.15083	
2	1.02062	15	1.16222	
8	1.03092	16	1.17420	
4	1.04123	17	1.18618	
5	1.05154	18	1.19816	
6	1.06229	19	1.21014	
7	1.07304	20	1.22212	
8	1.08379	21	1.23465	
9	1.09454	22	1.24718	
10	1.10529	23	1.25972	
11	1.11668	24	1.27225	
12	1.12806	25	1.28478	
13	1.13945			

:8. at 15° in 10 p.c. alcohol 39.3, 20 p.c. alcohol 21.3, 40 p.c. alcohol 1.62 (Schiff, A. 118, 365). Heated with NaCl gives off HCl, leaving Na2SO4, MgCl₂ and some MgSO₄ (Ramon de Luna, J. pr. 66, 256). Decomposed, almost entirely, by heating in steam, giving off H2SO4 and leaving MgO (Clemm, J. 1864. 764). Strongly heated with charcoal gives off SO₂ and CO₂ (or CO), and leaves MgO. Mixed with NaCl and heated in steam gives Na₂SO₄, MgO, and HCl.

Other hydrates of magnesium sulphate. (1)

MgSO₄ aq occurs native as kieserite, S.G. 2-281 at 16° (Pape, P. 120, 369); dissolves slowly in cold water, easily in boiling water, forming MgSO₄.7aq; prepared by heating the heptahydrate for a long time at 150°-160° (Pickering, C. J. 47, 100); S.G. 2445 (Thorpe a. Watts, C. J. 37, 102). (2) MgSO₄.6aq, by crystallising a solution of the heptahydrate above 40°; S.G. 1.734 (T. a. W., l.c.). (3) MgSO, 5aq, by drying the heptahydrate over conc. H₂SO₄; S.G. 1.869 (T. a. W., l.c.). (4) MgSO₄. 2aq, by boiling powdered MgSO4.7aq with absolute alcohol; S.G. 2.373 (T. a. W., l.c.). (5) According to Jacquelain (A. Ch. [3] 32, 201), the hydrates 4M.7aq, 4M.9aq, and 2M.5aq exist $(M = MgSO_4)$.

Acid salts. Schiff (A. 106, 115) obtained deliquescent, hexagonal plates, MgSO₄.H₂SO₄, from a solution of MgSO₄ in hot conc. H₂SO₄; according to Schultz (P. 130, 149), the crystals

are MgSO₄.3H₂SO₄.

Double salts. With K₂SO₄, or (NH₄)₂SO₄, to form compounds MgSO4.M2SO4.6aq; by crystallising mixtures of the component salts in proper proportion. For solubility and S.G. of solution of the K₂SO₄ compound, v. Tobler (A. 95, 193), Schiff (A. 113, 183), Gerlach (Fr. 8, 287). The compound of MgSO, and Na2SO, contains 4aq (cf. van't Hoff a. Deventer, B. 19, 2144). The compound MgSO, KCl. 3aq occurs native as kainite (v. Reichardt, Ar. Ph. 159, 204; Erlenmeyer, B. 2, 289). A double salt MgSO4.Al2(SO4)3.xH2O occurs as feather alum, and pickeringite. Some triple salts of MgSO. with CaSO, and K,SO, occur as minerals (v. Reichardt, Ar. Ph. 159, 204; Precht, B. 14, 2138).

Manganese sulphates. I. Manganous salts. The normal salt MnSO, is obtained by dissolving MnCO, or Mn in warm dilute H2SO, Aq, crystallising out MnSO. 5aq, drying this salt, and heating to c. 280° until it ceases to lose weight. A white salt, with very faint rose tint; S.G. 8:282 (Thorpe a. Watts, C. J. 37, 113). H.F. [Mn,S,O⁴] = 249,900 (Th. 3, 516). S. 56:5 at 6.3°, 60 at 18.7°, 68.7 at 37.5°, 67 at 75° (Brandes, P. 20, 575). Decomposed by heating to full redness, giving off SO₂ and O, and leaving Mn₃O₄; heated in steam is said to give an oxysulphide Mn₂OS (Arfvedson, P. 1, 50).

Hydrates of manganous sulphate. MnSO₄.7aq separates from solutions of Mn or MnCO₅ in H_2 SO₄Aq below 6° (Mitscherlich, P. 25, 287; Claassen, Ar. Ph. [3] 25, 310). Very pale rose-coloured, monoclinic crystals; isomorphous with FeSO₄.7aq. Very sol. water; S. at 18.5° c. 200 (Jahn, A. 28, 110). According to Etard (C. R. 86, 1399), solubility increases from 0° -55°, and decreases from 55° -145°; the changes are probably due to the formation of different hydrates. (2) MnSO, 5aq, obtained by crystallising between 7° and 20°, or by treating a conc. solution of MnSO, with 95 p.c. alcohol a conc. solution of MnSO, with 95 p.c. alcohol and allowing to crystallise slowly (Claassen, Ar. Ph. [8] 25, 310). S.G. 2·103 (Thorpe a. Watts, C. J. 37, 113). H.F. [Mn,S,O,5H²O] = 263,600 (Th. 3, 516). (3) MnSO. 4aq, obtained by crystallising between 20° and 30° (C., l.c.; Mitscherlich, P. 25, 287). S.G. 2·261 (Topsoö, C. C. 4, 76). This is the main constituent of ordinary manganese sulphate. Mulder gives solubilities as follows:—

Temp.	S. MnSO4. 4aq	1 Tem	S. MnSO. 4aq
0ô	55.4	50°	74.8
5	58.2	54	75.3
10	63.8	63.5	61.3
20	66.3	64	61.5
25	68.5	85	61.3
30	70.4	90	60.3
35	71.9	95	57.9
40	73.1	100	52.9
45	74.0		

Insol. absolute alcohol (cf. Schiff, A. 118, 365). (4) MnSO .. 2aq, obtained by boiling the powdered pentahydrate with absolute alcohol; also by pouring a saturated solution of the pentahydrate into conc. H.SO.; S.G. 2.526 (T. a. W., l.c.). (5) MnSO4. aq, obtained by heating the pentahydrate at 100° till it ceases to lose weight; S.G. 2.845 (T. a. W., l.c.).

Linebarger (Am. 15, 225) has obtained hydrates of MnSO₄ with 1, 2, 3, 4, 5, 6, and $7 H_2O$. Below -10° mixtures of the heptahydrate and ice separate from MnSO₄Aq; above 117° only MnSO₄ separates. The solubilities of the various hydrates are discussed; the more water in the hydrate, the more soluble it is.

Acid salts MnSO4.H2SO4 and MnSO4.3H2SO4 were obtained by Schultz (P. 130, 149) by dissolving the normal salt in hot conc. H2SO4.

Basic salts are formed by adding a little warm KOHAq to a large excess of boiling MnSO, Aq (v. Gorgeu, C. R. 94, 1425).

Double salts; (1) with sulphates of Cu, Fe, Mg, and Zn to form salts MnSO. MSO. xH.O (v. Rammelsberg, P. 91, 321; Vohl, A. 94, 73). (2) With Al₂(SO₄)₃, Fe₂(SO₄)₃, and Cr₂(SO₄)₃ v. Etard (C. R. 86, 1399; 87, 602). (3) With alkali sulphates to form MnSO4.M2SO4.4aq or 6aq (v. Marignac, Ann. M. [5] 9, 1; Pierre, A. Ch. [3] 16, 239; von Hauer, J. pr. 74, 431). For various triple salts of MnSO, with alkali sulphates and CuSO, MgSO, &c.; v. Vohl (A. 94, 57).

II. Manganic salts .- The normal salt $Mn_2(SO_4)_4$ is obtained by the action of hot conc. H₂SO₄ on MnO₂. Carius (A. 98, 53) recommends to prepare pure MnO₂ by passing Cl into NaOHAq containing MnCO₃ in suspension, to wash and dry the MnO2, to rub up to a paste with conc. H2SO4, and heat gradually to 110° till O is given off and the mass becomes greyviolet, then to heat to c. 135°-140° till the whole is liquid, to pour the solid that forms as the liquid partially cools on to warm unglazed porcelain, after H2SO4 has become absorbed to mix with a little conc. HNO, Aq and allow to drain on the porcelain, to repeat this treatment with HNO₃ six or eight times, and finally to heat at 150° till all HNO₃ is removed. Franke (J. pr. [2] 36, 451) obtained Mn₂(SO₄)₃ by heating 8 g. KMnO, with 100 c.c. conc. H₂SO, at 100° for some time. A dark-green powder; Franke describes it as green crystals. Decomposed above 160°, giving MnSO4. Insol. conc. H2SO4 or conc. HNO2Aq. Sol. conc. HClAq to a brown liquid, which gives off Cl when warmed. Mn₂(SO₄), is very hygroscopic; quickly decomposed by water, with separation of MnO2.xH2O (Carius, l.c.).

Forms double salts with Al₂(SO₄)₃, Fe₂(SO₄)₃, and Or₂(SO₄)₂ (v. Étard, C. R. 86, 1399). The double compounds of Mn2(SO4), with alkali sul-

phates— $Mn_2(SO_4)_3$, M_2SO_4 , 24aq—are alums. According to Fremy (C. R. 82, 1231), the salt Mn(SO₄)₂, manganese disulphate, corresponding with MnO2, is formed by decomposing KMnO1 by a considerable excess of cold conc. H₂SO₄ (cf.

Franke, J. pr. [2] 36, 453).

Mercury sulphates. I. Mercurous salts. The normal salt Hg₂SO₄ is obtained by gently warming equal weights of Hg and conc. H2SO4, removing the white solid so produced before the whole of the Hg is changed, washing with cold water, and drying at a low temperature. Divers a. Shimidzu (C. J. 47, 639) recommend to mix Hg with fuming sulphuric acid rich in SO, in a covered dish, in a cool place, adding a little more Hg from time to time till there is sufficient to saturate the SO, in the acid, but avoiding excess, and when the Hg is all converted into Hg₂SO₄ to set the dish in a warm place to expel SO₂. Hg₂SO₄ is also obtained by adding 6 pts. water to 18 pts. HgSO, and rubbing with 11 pts. Hg (Planche, A. 66, 168). Also by ppg. HgNO₃Aq by dilute H2SO, Aq or solution of a sulphate; or by rubbing together HgNO, and Na₂SO₄ and washing with cold water (H. Rose; Stoedeler, A. 87, 129). A white powder, consisting of monoclinic prisms (S., l.c.); S. G. 7.56 (Playfair a. Joule, C. S. Mem. 2, 401). Sl. sol. water; S. in cold water 2, in boiling water 33; more sol. dilute HNO3Aq; sol. conc. hot H2SO4, crystallising out on cooling. Hg₂SO₄ becomes grey on exposure to light (v. Buchner, Chem. Zeitung, 10, 759): hence the salt should be kept in the dark. Hg₂SO₄ melts to a reddish liquid, and decomposes at a higher temperature, giving off SO, and O, and also partially subliming. Slowly changed by boiling water to an acid salt, and a yellow powder which is probably a basic salt (v. Kane, P. 42, 367). An acid salt Hg₂SO₄.H₂SO is said to be formed by the prolonged reaction of conc. H₂SO₄ containing a drop of HNO₃Aq on Hg (Braham, C. N. 42, 163). A double salt $Hg_2SO_4.2HgSO_4$ was obtained by Brooks (P. 66, 63) by gently warming mercuro-mercuric nitrate (formed by warming 2 pts. Hg with 3 pts. HNO, Aq S.G. 1.2) with Na, SO, Aq.

II. Mercuric salts. The normal salt HgSO₄ is obtained by the reaction of 6 pts. boiling conc. H₂SO₄ with 5 pts. Hg, till SO₂ ceases to be evolved, and a little of the white solid produced gives no pp. of HgCl when dropped into dilute HClAq; the white mass is gently warmed till acid fumes cease to be given off. A white powder; S.G. 6.466 (Playfair a. Joule, C. S. Mcm. 2, 401). Becomes yellow, then red when heated; decomposes at red heat to Hg, O, and SO₂, a little HgSO₄ subliming (Mohr, A. 31, 180). Decomposed by water to a yellow basic salt and H₂SO₄Aq. Hot HgSO₄ absorbs HCl and HBr gases; according to Ditte (A. Ch. [5] 17, 120) the compounds HgSO, 2HCl and HgSO4.2HBr are formed; these bodies are said to be sol. water without formation of basic salts, and HgSO, 2HCl is said to sublime unchanged when heated. Solution of HgSO, in H₂SO, Aq is said to react with PH, to form the compound 2(3HgO.SO₂).Hg₂P (H. Rose, P. 40, 75). Monohydrated mercuric sulphate HgSO₄ aq was obtained by Eisfeldt (Ar. Ph. [2] 76, 16) by covering a thin layer of HgSO, with the calculated quantity of water. The basic salt HgSO, 2HgO

(=3HgO.SO₃) is obtained by the action of hot | water on HgSO4, also by adding Na2SO4Aq to hot $Hg(NO_3)_2Aq$; a citron-yellow powder, S.G. 6.44; nearly insol. water (v. Cameron, An. 1880. 144). Decomposed by Cl at the ordinary temperature. Absorbs HCl and HBr, forming HgSO. 2HgO.6HCl(or 6HBr) (Ditte, I.c.). This basic salt was formerly known as minerale turpethum or turbith mineral (from its resemblance to a coloured gum obtained from an Oriental creeping plant [Ipomæa turpethum]). Hopkins (Am. S. 18, 364) described another basic salt 8HgSO, HgO = 4HgO.8SO, Double salts. — (1) HgSO, (NH_d)₂SO, xH₂O and HgSO, K₂SO, xH₂O (v. Hirzel, J. 1850. 322). (2) HgSO, HgI (v. Souville, J. Ph. 26, 474). (3) 2HgSO, HgS (v. Jacobsen, P. 68, 411).

Molybdenum sulphates. By evaporating a solution of MoO, in conc. H2SO, Schultz-Sellack (B. 4, 14) obtained white lustrous crystals of MoO₂.SO₃; the compound MoO₃.SO₃, 2aq, described by Anderson (\hat{B} . J. 22, 161), could not be

obtained by S-S.

The normal salt NiSO, Nickel sulphates. is formed, as a clear yellow powder, by heating powdered NiSO₄. xaq (v. infra) to c. 800°. The salt is obtained in crystals, regular octahedra, by heating NiSO₄, NiO, or NiCO₈ with five or six times its weight of fused (NH₄)₂SO₄, separating the yellow crystals of the double salt $3NiSO_4.2(NH_4)_2SO_4$ that form, and heating these in presence of some $(NH_4)_2SO_4$ in a partly closed crucible out of contact with the gases of the flame till all the ammonium salt is removed; S.G. of crystals 3.67 at 20° (Lepierre a. Lachaud, C. R. 115, 115; Klobb, C. R. 114, 836). Tobler (A. 95, 193) gives solubilities of NiSO₄ as follows :-

Temp.	S.	Temp.	s.
20	30.4	410	49.1
16	37.4	50	52.0
20	39.4	53	54· 4
23	41.0	60	57.2
31	45.3	j 70	61.9

The hexahydrate NiSO4. 6aq is obtained by dissolving Ni, NiO, or NiCO, in excess of H2SO, Aq, evaporating, and crystallising. At the ordinary temperature, bluish tetragonal pyramids separate (Brooke a. Phillips, P. 6, 193); greenish monoclinic crystals separate from warm solutions (Pierre, A. Ch. [3] 16, 252). The salt is, therefore, dimorphous (Marignac, A. 97, 294; cf. Mitscherlich, P. 12, 144; also Lecoq de Boisbaudran, A. Ch. [4] 9, 173). By dissolving this hydrate in water, and crystallising at 15° 20°, emerald-green rhombic prisms of the heptahydrate NiSO₄. 7aq are obtained, isomorphous with MgSO₄.7aq; melts at 98°-100° (Tilden, C. J. 45, 267). When this hydrate is heated to 103° it loses 6H₂O and leaves the monohydrate NiSO4. aq. By passing NH, into NiSO4Aq, compounds of the form NiSO ... xNH .. yH2O are obtained (x-4, 5, and 6; v. F. Rose, Ammon.Kobaltverbind. [Heidelberg, 1871] 27; also P. 20, 156).

Double salts. (1) With (NH₄)₂SO₄, K₂SO₄, and Tl₂SO₄ to form NiSO₄.M₂SO₄. 6aq (v. F. Rose, l.c.; Link, Crell's Ann. 1796 [1] 82; Werther, J. pr. 92, 182). (2) With BeSO, to form various compounds (Klatzo, Über die Constant. der

Beryllerde, Dorpat, 1868). (3) With OuSO,, CoSO, FeSO, MgSO, MnSO, and with some of these and also alkali sulphates (v. Pierre, A. Ch. [8] 16, 253; Lecoq de Boisbaudran, C. R. 66, 497).

Basic salts are formed by the gradual reaction of NiSO, Aq with NiCO,, and by adding a little NH, Aq to boiling NiSO, Aq (Habermann, M. 5, 440)

Palladium sulphates. The normal palladous salt PdSO. 2aq is obtained in brown crystals by evaporating a solution of Pd in H2SO, Aq containing HNO₃. Addition of a little KOHAq to a conc. solution in water of the normal salt ppts. a basic salt PdSO₄.7PdO.6aq (v. Kane, B. J.

24, 236).

Platinum sulphates. By dissolving PtO H₂ in dilute H₂SO₄Aq, Berzelius (*Lehrbuch* [5th ed.] 8, 987) obtained a dark syrup which probably contained platinous sulphate PtSO₄. Platinic sulphate Pt(SO₄)₂ was obtained by Berzelius (L.c., p. 989) by evaporating PtCl₄ with conc. H2SO4 (1 part acid for 1 part Pt), as an almost black solid. E. Davy (T. 1820) obtained the salt by oxidising PtS, by fuming HNO. Prost (Bl. [2] 46, 156) obtained a basic salt by allowing an acid solution of Pt(SO₄)₂ to stand for some days; Prost also describes several double salts of

Pt(SO₄)₂ with alkali sulphates.

Potassium sulphates. The normal salt K_2SO_4 is found in the lava from Vesuvius, in small quantities in sea water, and in some mineral springs, in combination with MgSO, and MgCl₂ as kainite (K₂SO₄.MgSO₄.MgCl₂.6aq), in combination with MgSO₄ and CaSO₄ as polyhalite (K₂SO₄.MgSO₄.2CaSO₄.2aq), and as alum. K₂SO₄ is prepared by decomposing KCl by H₂SO₄. by neutralising H2SO4Aq by KOH or K2OO3, and evaporating (for the preparation of K2SO, on the large scale v. Dictionary of Applied CHEMISTRY, vol. iii. p. 279). K₂SO, forms in white, hard, four-sided, trimetric prisms, unchanged in air. S.G. 2.656 at 4° (Playfair a. Joule, C. J. 1, 182); for S.G. at various temperatures from 0° to 100° v. Spring, B. 15, 1940. Melts above 861° (Carnelley, Melting- and Boiling-Point Tables, 1, 33); melts at 1073° according to V. Meyer a. Riddle (B. 26, 2443). H.F. $[K^2,S,O^4] = 344,600$ (Th. 3, 516); [K°SO',Aq] = -6167 (Pickering, C. J. 47, 98). Mulder (Scheikund. Verhandg. 1864. 49) gives solubilities in water as follows:-

Temp.	s.	Temp.	s.
5°	9.1	550	16.8
10	9.7	60	17.8
15	10.3	65	18.8
20	10.9	70	19.8
25	11.6	75	20.8
80	12.3	80	21.8
85	13.1	85	22.8
40	14.0	90	23.9
45	14.9	95	25.0
50	15.8	100	26·2

Gerlach (Fr. 8, 827) gives S.G. of K2SO4Aq as follows :-

P	ctge.K.SO.	S.G. K. 80, Aq	Pctge.K.SO.	S.G. K.SO.Aq
	1	1.00829	6	1.04947
	2	1.01685	7	1.05790
	B	1.02450	8	1.06644
	4	1.03277	9	1.07499
	5	1.04105	9.92 (satd	1.) 1.08805

Insol. absolute alcohol; sol. aqueous alcohol in ! proportion to amount of water present (v. Schiff, A. 118, 362). Insol. KOHAq 1.32 S.G.; sl. sol. KOHAq 20 p.c. S. in saturated NH,Aq = 0365 (Giraud, Bl. [2] 43, 552). S. in glycerin (S.G. 1.225) at ordinary temperature = 1.32 (Vogel, N. R. P. 16, 557). More sol. in solution of CuSO₄, MgSO₄, or Na₂SO₄ than in water. Reacts with strong acids to give KHSO, (for thermal measurements v. Berthelot, Essai de méc. heasterness v. Bernston, Essay at the mec. chimique, 1, 389). Dissolves in hot conc. HNO₂Aq, KHSO₄ separates on cooling, then KNO₃, and then a compound of K₂SO₄ with HNO₂; H₂PO₄Aq reacts similarly (v. Jacquelain, Biedermann's Techn. Chem. J. 4, 62). Absorbs HCl gas when heated, forming KCl and KHSO, (Thomas, C. J. 33, 372; Hensgen, B. 9, 1671; 10, 259). Decomposed by heating with NH₄Cl, giving KCl (H. Rose, P. 64, 568). Heated to redness with iron filings gives KOH mixed with Fe₂O₈ and FeS (d'Heureuse, P. 75, 255). Reduced to sulphide (or KSH) by heating in CO; to KSH and KOH by heating in H (v. Berthelot, C. R. 110, 1106). Sulphur reacts at full red heat, giving SO₂ and polysulphides of K (Berthelot, C. R. 96, 303). SO2 is without action at a red heat.

Double salts. (1) With Al₂(SO₄), Or₃(SO₄), and Fe₂(SO₄), to form K₂SO₄.M₂(SO₄), 24aq; v. potash alum, potash chrome-alum and potash iron-alum, under Alumnium sulphates (p. 568), Chromium sulphates (p. 570), and Iron sulphates (p. 572). (2) With sulphates of Ce and Di respectively, to form 3K₂SO₄.M₂(SO₄), (v. Cleve, Bl. [2] 43, 359; Czudnowitz, J. pr. 82, 129; Hermann, J. pr. 82, 385); also the salt 2K₂SO₄.Ce(SO₄), 2aq (Marignac, Ann. M. [5] 15, 275). (3) With sulphates of Cd, Co, Cu, Fe, Mg, Mn, Ni, Zn to form K₂SO₄.MSO₄.6aq (v. Graham, P. M. 1836. 327, 417; Brunner, P. 15, 476; 32, 221; Pickering, C. J. 49, 1; Tobler, A. 95, 193; Schiff, A. 108, 326; 113, 183; Gerlach, Fr. 8, 287; Marignac, Ann. M. [5] 9, 19; Wilm, A. Ch. [4] 5, 56; Pierre, A. Ch. [3] 16, 239; von Hauer, J. pr. 74, 431; Werther, J. pr. 92, 132). (4) K₂SO₄.BeSO₄.2aq (Awdejeff, P. 56, 101). (5) K₂SO₄.Na₅SO₄ (Penny, P. M. [4] 10, 401; Mischerlich, P. 58, 468; von Hauer, J. pr. 83, 856).

The acid salt KHSO, occurs in some volcanic districts as misenite; it is formed by heating 13 parts K₂SO₄ with 8 parts conc. H₂SO₄, or by heating KNO₂ and conc. H₂SO₄ in the ratio KNO₂:H₂SO₄. Crystallises in white rhombic forms, S.G. 2:273; also in monoclinic forms, S.G. 2:245 (v. Wyrouboff; abstract in C. J. 50, 665). Melts at c. 200°. Easily sol. water; Kremers gives S. 33·9 at 0°, 48 at 20°, 62·9 at 40°, 114 at 100°. An aqueous solution is said to deposit, first K₂SO₄, then KHSO₄, K₂SO₄, and then KHSO₄; by evaporating KHSO₄Aq at a low temperature until a solid mass formed on removing the flame, arborescent and very deliquescent crystals formed on the surface of the solid, which were 2KHSO₄.11H₂O according to Senderens (Bl. [3] 2, 728).

Other acid salts. (1) KHSO, K,SO, obtained by evaporating KHSO, Aq. (2) 2KHSO, K,SO, erystallising from K,SO, in H,SO,Aq (Phillips, J. pr. [2] 1, 429; Marignac, Ann. M. [5] 9, 7). (3) KHSO, H,SO, formed by heating K,SO, with Vox. IV.

less than 5 parts conc. H₂SO₄ (Schultz, P. 183,

Rhodium sulphates. The normal rhodic sulphate Rh₂(SO₄)_a is obtained by dissolving Rh₂O₄.xaq in H₂SO₄Aq, evaporating, and heating to 400° till excess of acid is removed; a red powder, decomposed by hot water to the yellow basic salt Rh₂(SO₄)_a.Rh₂O₃ (Leidié, C. R. 107, 234). Claus (Beiträge zur Chemie der Platinmetalle [Dorpat, 1854]) obtained the hydrate Rh₂(SO₄)_a.12aq, as a pale-yellow crystalline salt, by crystallising a solution of Rh₂O₃.xaq in H₂SO₄Aq, and washing with alcohol. The double salt Rh₂(SO₄)_a.3K₂SO₄ was obtained by Claus (l.c.) by evaporating a solution of the constituent salts in the ratio Rh₂(SO₄)_a:5K₂SO₄. The double salt Rh₂(SO₄)_a.Na₂SO₄ was obtained by Bunsen (A.146, 265; v. also Seubert a. Kobbé, B. 23, 2556) by the action of hot cone. H₂SO₄ on RhSO₂:6Na₂SO₂. 9aq (formed by heating RhCl₂Aq for some time with NaHSO₄).

Rubidium sulphates. The normal salt Rb₂SO₄ is obtained by neutralising H₂SO₄Aq by Rb₂CO₅ and evaporating; it crystallises in transparent, hexagonal forms, unchanged in air. S. 42.4 at 70°. S.G. 3.6438 at 0°; 3.6256 at 50°; 3.6036 at 100° (Spring, B. 15, 1940). If excess of H₂SO₄Aq is used, the acid salt RbHSO₄ separates in rhombic prisms (Kirchoff a. Bunsen, P. 115, 584).

Double salts. (1) With Li₂SO₄ to form RbLiSO₄ (Wyrouboff, Bl. [2] 48, 184, 630). (2) With CaSO₄ and PbSO₄ to form the compounds Rb₂SO₄.2MSO₄, xaq (Ditte, C. R. 89, 641). (3) With Al₂(SO₄), to form rubidium alum Al₂(SO₄), Rb₂SO₄. 24aq (Bunsen, P. 119, 1; Godeffroy, A. 181, 176; Redtenbacher, J. pr. 95. 148).

Ruthenium sulphate. Normal ruthenic sulphate Ru(SO₄)₂ is obtained by oxidising RuS₂ (formed by prolonged action of H₂S on RuCl₂Aq) by conc. HNO₂Aq, and evaporating off excess of acid. An amorphous, brownish-yellow, very hygroscopic solid. Strongly heated gives RuO₂ (Claus. J. vs. 42, 364)

(Claus, J. pr. 42, 364).

Silver sulphates. The normal salt Ag₂SO₄ is obtained by heating finely divided Ag with excess of conc. H₂SO₄, by neutralising H₂SO₄Aq by Ag₂O or Ag₂CO₅ and evaporating, by evaporating AgNO, with H₄SO₄Aq, or by ppg. AgNO₅Aq by Na₂SO₄Aq. Small, white, lustrous, rhombic crystals, isomorphous with Na₂SO₄ (Mitscherlich, P. 12, 138; 25, 301). S. 1·15 cold water, 1·45 at 100° (Wenzel, P. 82, 136). S.G. 5·322 (Playfair a. Joule, G. S. Mem. 2, 401), 5·425 when fused (Schröder, P. 106, 226). Decrepitates when heated to 300°; melts at 645° (Carnelley, C. J. 33, 279). Decomposed at very high temperature, giving Ag, SO₂, and O. Reduced to Ag by heating in H, with O, or in CO (Stammer, J. pr. 11, 70). Absorbs 2NH₂ (H. Rose, P. 20, 153).

Acid salts are obtained by dissolving Ag₂SO₄ in different quantities of H₂SO₄Aq and crystallising; Schultz (P. 133, 137) describes (1) AgHSO₄, (2) 2Ag₂SO₄,3H₂SO₄,2aq, (3) Ag₂SO₄,3H₂SO₄,2aq. Carey Lea (Am. S. [8] 44, 322) describes a hemisulphate Ag₂SO₄,Ag₄SO₄, aq, but this salt was not obtained free from Ag₂PO₄.

Forms a double salt Ag.SO₄.Al₂(SO₄), 24aq; this salt is an alum (v. Church a. Northcote, PP

C. N. 9, 155). Also combines with Ag₂S and with K₂SO₄ (Berzelius; H. Rose, P. 53,

Sodium sulphates. The normal salt Na2SO4 occurs native as thenardite, in combination with CaSO, as glauberite, and with MgSO, as loweite; the salt also occurs in sea water and in many mineral springs. It is prepared by adding H2SO4 to NaCl in the ratio NaCl:H2SO4, and strongly heating the NaHSO₄ thus produced; also by the reaction between NaCl, SO₂, steam and O at 400°-450°; 2NaCl+SO₂+O+H₂O = Na₂SO₄+2HCl (for the application of these processes on the large scale v. DICTIONARY OF APPLIED CHEMISTRY, vol. iii. p. 439). Na2SO4 is also obtained in the decomposition of NH Cl, or NaNO, by H₂SO₄; it is formed by neutralising H₂SO₄Aq by NaOHAq or Na₂CO₃, evaporating, and heating to redness the crystals of Na₂SO₄. xaq that separate. Na₂SO₄ is a white amorphous powder; S.G. 2.597 (Playfair a. Joule, C. S. Mem. 2, 401); 2.631 (Karsten, S. 65, 894). Obtained in rhombic crystals, isomorphous with Ag₂SO₄ (Mitscherlich, P. 12, 138; 25, 301), by heating to 40° a solution saturated at 33°: S.G. 2.6618, or 2.6637 crystallised at 110° (Nicol, P. M. [5] 15, 94). Dissolves in water with production of heat; according to Berthelot (C. R. 78, 1722) [Na²SO⁴,Aq] = 780 at 21·2°, but -100 at 3°; Pickering (C. J. 45, 686) thinks that two modifications of Na2SO, exist; the salt dried at 100° or 150° gave [Na^2SO^4 , Aq] = 57 at 20.4°, but after heating to temperatures varying from 250° to the melting point the heat of solution was $[Na^2SO^4,Aq] = 760$. Thomsen (Th. 3, 516) gives H.F. $[Na^2,S,O^4] = 328,600$. Tilden (Pr. 35, 345)gives solubilities as follows: 5 at 0°, 78.8 at 34° 42.7 at 100°, 41.95 at 120°, 42 at 140°, 42.9 at 160°, 44.25 at 180°, 46.4 at 230°. Étard (C. R. 113, 854) says that S. increases from 33° to 80°, remains constant to 230°, and then decreases to 320° (v. also Löwel, A. Ch. [3] 33, 334; 49, 32; Gernez, C. R. 60, 833; de Coppet, Bl. [2] 17, 146). For S.H. and expansion of Na, SO, Aq v. Thomsen (P. 142, 337) and Marignac (A. Ch. [4] 22, 415). Na, SO, melts at 861° (Carnelley, C. J. 33, 280); at 843° (V. Meyer a. Riddle, B. 26, 2443). S.G. at melting-point = 2.104 (Braun, B. 7, 958).

Heated with charcoal out of air Na, SO, is reduced to Na₂S; Gay-Lussac (A. Ch. [2] 11, 312) says polysulphides are also formed; CO2 is evolved, but no CO (Unger, A. 63, 240; Scheurer-Kestner, A. Ch. [4] 1, 412; Stromeyer, A. 107, 361); heated to bright redness with C in a vessel containing air, Na2S is formed and CO evolved (Berthelot, C. R. 110, 1106). Reduced to sulphide by heating to bright redness in carbon monoxide (B., l.c.); reduction by hydrogen begins at c. 500°, the products are Na₂S, NaSH, NaOH, H₂O, and H₂S (B., *l.c.*). Heated with hydrogen chloride, NaCl is formed (Bousingault, C. R. 78, 593); boiling HClAq produces NaCl and NaHSO, (Thomas, C. J. 33, 373). A little NaCl is formed by heating with ammonium chloride (Nicholson, C. N. 26, 47). Boiled with barium carbonate some Na₂CO₂ and BaSO₄ are formed; after a time a condition of equilibrium is attained. A very little NaOHAq is produced by boiling for some time with milk of lime; under increased pressure NaOHAq is formed

(v. Scheurer-Kestner, A. Ch. [4] 1, 412). Not decomposed by heating strongly with calcium carbonate (S.-K., l.c.). When a mixture of Na₂SO₄ and alumina is heated in steam, SO₂ is given off and sodium aluminate remains; presence of charcoal hastens the reaction (Wagner). According to Stromeyer (A. 107, 361) fusion with iron produces Fe₂O₄, Na₂O, and Na₂S.2FeS. The final products of heating with silica and carbon in a closed crucible are Na₂O.2SiO₂, S, CO₂ and CO (S.-K., C. R. 114, 117).

Hydrates of Na, SO₄. (1) The hepta-hydrate Na, SO₄. 7aq is obtained by melting Glauber's salt Na, SO, 10aq in its water of crystallisation, heating to boiling, at once closing the vessel, and shaking repeatedly as the liquid cools till 18° is reached; Na, SO, separates at first but dissolves again, and below 18° Na, SO, 7aq separates (Löwel, A. Ch. [3] 33, 334; 49, 32). The following method gives better results (L., l.c.): 22 parts Na, SO, 10aq are dissolved in 10 parts water, the solution is heated to boiling, and the flask is then closed with a cork carrying two narrow tubes bent at right angles; when the liquid is cold the outer end of one of the tubes is dipped into 30 p.c. alcohol at 40°. and the alcohol is caused to run into and fill the flask by sucking at the outer end of the other As the alcohol withdraws water, large, translucent, rhombic or tetragonal crystals of Na2SO. 7aq gradually separate. The crystals must be quickly pressed between filter paper. They rapidly absorb water from the air, changing to Na,SO, 10aq; exposed to air, even under dilute alcohol, they soon change to the decahydrate. According to Löwel (l.c.) the heptahydrate is much more soluble in water between 0° and 26° than the decahydrate. The temperature of maximum solubility is 27°; the solution then contains 56 p.c. Na₂SO₄, which is the same quantity as is present in solution of Na2SO4. 10aq saturated at 34°-36°. If a crystal of Na₂SO₄.7aq is added to the saturated solution of this salt at 27° it does not dissolve, but crystals of Na₂SO₄ separate until a concentration is reached corresponding with a saturated solution of Na2SO, at the temperature. The heptahydrate is able to form supersaturated solutions, but less readily than the decahydrate; addition of a crystal of Na₂SO₄, 7aq causes separation of the hepta-hydrate. These results were confirmed by de Coppet (Bl. [2] 17, 146).

(2) The decahydrate Na SO₄. 10aq crystallises from a hot saturated solution of Na₂SO₄ cooled to at least 32°; a supersaturated solution may be cooled in a closed vessel to 0°. If a crystal of Na₂SO₄. 10aq is dropped in, a mass of crystals of the decahydrate is formed, and the temperature rises to c. 18°. Large, white, striated, monoclinic prisms; isomorphous with Na₂CrO₄. 10aq and Na₂SeO₄. 10aq. S.G. 1.469 (Playfair a. Joule, C. S. Mem. 2, 401); 1.485 at 19°, 1.492 at 20° (Pettersson, Upsala, Nova Acta, 1874). [Na₂SO₄.10H²O] = 347,800 (Th. 3, 516). Melts at 34° (Tilden, C. J. 45, 267). Effloresces in air. When melted in the water of crystallisation, boiled, and allowed to cool in a closed vessel, Na₂SO₄ separates, but on shaking, this salt dissolves till 18° is reached, after which crystals of Na₂SO₄. 7aq separate (v. supra). Na₂SO₄. 10aq is insol. absolute alcohol; alcohol of S.G. 976 dissolves book at the superal of S.G. 976 dissolves in the salt of the superal of S.G. 976 dissolves the salt of the superal of S.G. 976 dissolves the salt alcohol of S.G. 976 dissolves the salt of the salt of S.G. 976 dissolves the salt of the salt of S.G. 976 dissolves th

solves 14.85 p.c., of S.G. 972 dissolves 5.6 p.c., and of S.G. 939 dissolves 1.3 p.c. Solubility in water increases with temperature to 33°-34°. whereat the solution contains 55 parts Na₂SO₄ in 100 of water; a crystal of Na₂SO₄, 10aq dropped into the solution dissolves, and Na2SO4 separates till 49.53 parts Na, SO, are in solution in 100 parts water, which is the same quantity of Na₂SO₄ as is present in a solution made by saturating water with Na₂SO₄ at 34° (Löwel, A. Ch. [3] 83, 334; 49, 32). As temperature is increased above 34° Na₂SO₄ seems to be formed, and the solubility of this salt decreases as temperature rises to the b.p. of saturated Na, SO, Aq. There appear to be three temperatures of maximum solubility for Na₂SO₄ and its hydrates: (1) at 34°, which is the maximum for Na₂SO₄. 10aq, (2) at 26°-27° for Na₂SO₄. 7aq, (3) at 17°-18° for Na2SO4. The quantity of salt in solution, calculated as Na. SO., is about the same at these temperatures. Löwel (l.c.) gives the following table:

S.G.	P.o. Na 80	P.c. Na.SO., 10aq
1.1015	11.025	25
1.1057	11.466	26
1.1100	11.907	27
1.1142	12.348	28
1.1184	12.789	29
1.1226	13.230	30

Acid salts. (1) NaHSO₄; by adding 7 parts H_2SO_4Aq S.G. 1.85, to 10 parts Na_2SO_4 , and heating gently till the mass melts and water-vapour ceases to be given off. By dissolving in water, or by evaporating Na2SO, in H2SO4Aq at 50°, crystals of NaHSO, aq are obtained (Mitscherlich, P. 12, 138; H. Rose, P. 82, 545). (2) Na, SO, NaHSO, (= Na, H(SO,),); by crystallising Na, SO, from half as much H, SO, in water as is required to form NaHSO, (M., L.c.; R., L.c.). (3) NaHSO₄.H₂SO₄ (NaH₃(SO₄)₂); by crystallising Na₂SO₄ from solution in almost 7 parts H₂SO₄ (Schultz-Sellack, J. pr. [2] 2, 459; Lescœur, C. R. 78, 1044).

Double salts. (1) Na2SO4.K2SO4 (Penny, P. M.

100 parts water contain, when solution is saturated,

Temp.	Na ₂ SO ₄ Sol. made v	Na SO. 10aq with Na SO.	Na ₂ 80 ₄ Sol. made w	Na ₂ SO ₄ . 10aq ith Na ₂ SO ₄ . 10aq	Na ₂ SO ₄ Sol.	Na ₂ SO ₄ . 7aq made with N	Na ₂ SO ₄ . 10aq a ₂ SO ₄ . 7aq
0°			5.02	12.16	19.62	44.84	59.23
10			9.00	23.04	80.49	78.90	112.73
15			13.20	35.96	37.43	105.79	161.57
18	53.25	871.97	16.80	48.41	41.63	124.59	200
20	52.76	361.51	19.40	58.35	44.78	140.01	234.4
25	51.53	337.16	28	98.48	52.94	188.46	365.28
26	51.31	333.06	30	109.81	54.97	202.61	411.45
30	50.37	316.19	40	184·09			
33	49.71	805.06	50.76	323.13			
34	49.53	302.07	55	412.22			
40.15	48.78	290.00					
45.04	47.81	275.34					
50.4	46.82	261.36					
59.79	45.42	242.89					
70.61	44.35	229.87					
84.42	42.96	213.98					
103.17	42.65	210.67					

at 19°:--

-;		
B.G.	P.o. Na ₂ SO ₄	P.o. Na SO. 10aq
1.0040	1.441	1
1.0079	1.881	2
1.0118	1.323	3
1.0158	1.764	4
1.0198	2.205	5
1.0238	2.646	6
1.0278	8.087	7
1.0318	8.528	8
1.0358	8.969	9
1.0398	4.410	10
1.0439	4.851	11
1.0479	5.292	12
1.0520	5.378	13
1.0560	6.174	14
1.0601	6.615	15
1.0642	7.056	16
1.0683	7.497	17
1.0725	7 ·938	18
1.0766	8.379	19
1.0807	8.820	20
1.0849	9.261	21
1.0890	9.702	22
1.0931	10.143	23
1-0978	10.584	24

with Li₂SO₄ (Rammelsberg, P. 128, 311; Wyrouboff, Bl. [2] 9, 35. (4) With sulphates of Cd, Ca, Cu, Di, In, Fe, La, Mn, and Mg; references under CADMIUM SULPHATES, CALCIUM SULPHATES, &c. (5) The compounds Na₂SO₄.M₂(SO₄), 24aq, where M = Al or Cr, are alums (references under ALUMINIUM SULPHATES and CHROMIUM SULPHATES).

Strontium sulphates. The normal salt SrSO, occurs native as celestins. It is obtained Strontium sulphates. by adding H2SO4Aq or solution of an alkali sulphate to solution of a strontium salt; large crystals are formed by dissolving ppd. SrSO, in conc. H₂SO₄, adding dilute HClAq, heating to 150°, allowing to cool, again heating to 150° with HClAq, and repeating this treatment three or four times (Bourgeois, C. R. 105, 1072). SrSO. is also obtained crystalline by fusing K2SO4 with excess of SrCl₂ (Manross, A. 82, 350). white solid; S.G. 3.927 to 3.955 crystalline (M., i.c.; Neumann, P. 23, 1); 8.7 ppd. (Schröder,
 P. 106, 226; v. also Clarke's Table of Spec.
 Gravs. [New Ed.] 82). H.F. [Sr,S,O'] = 330,909 (Th. 8, 516). Almost insol. water; S. 0145

ordinary temperature, '0104 at 100° (Fresenius); insol. dilute H₂SO₄Aq; S. conc. H₂SO₄ at 70° = 14 (Garside, C. N. 31, 245); for S. in solutions of NaCl, KCl, MgCl₂, and CaCl₂ v. Virck (C. C. 1862. 402). Melts when strongly heated; is decomposed to SrO, SO₂, and O at full white heat (Bousingault, J. 1867. 161). Reduced by heating strongly with C, Fe, or Zn (d'Heureuse, P. 75, 277), or in a steam of H or moist CO (Jacquemin, C. R. 46, 1164). Loses SO₄ when heated with As₂O₅ (Moretti, S. 9, 169), or when heated continuously to redness in HCl (Bousingault, C. R. 78, 593). Decomposed by heating with solutions of alkali carbonates. Forms double salts, SrSO₄,M₂SO₄, with alkali sulphates (H. Rose, P. 93, 604; 110, 296). An acid salt SrH₂(SO₄)₂ (= SrSO₄,H₂SO₄) is formed by digesting SrSO₄ in conc. H₂SO₄ with excess of SrSO₄

(Schultz, P. 113, 147).

Thallium sulphates. The normal thallous salt Tl2SO4 is obtained by dissolving Tl in hot H₂SO₄, or by neutralising H₂SO₄Aq by TlOH or Tl₂CO₃, and evaporating (Crookes, C. N. 24, 38). Forms white rhombic prisms, isomorphous with K₂SO₄ (von Lang, P. M. [4] 25, 348); S.G. 6.77 (Lamy, Bl. [2] 11, 210; v. also Clarke's Table of Spec. Gravs. [new ed.] 79). H.F. [Tl²,S,O⁴] = 221,000 (Th. 8, 516). S. 4.74 at 15°, 18.5 at 100° (Lamy, l.c.). Not decomposed by heating to full redness in absence of air; heated more strongly in air gives Tl₂O₂, SO₂, and O (Carstanjen, J. 1867. 279). Reduced to Tl₂S by heating with KCN. The acid thallous salt TIHSO, 3aq separates from a solution of the normal salt in H₂SO₂Aq after some months. When strongly heated yields Tl₂SO₄ and H₂SO₄ (Carstanjen, l.c.). Double salts, Tl₂SO₄MSO₄, 6aq, are formed when M = Cu, Fe, Mg, Ni, or Zn (Willm, A. Ch. [4] 9, 5; Werther, Bl. [2] 2, 272). The salt Tl. SO, Al. (SO.) 3. 24aq is an alum (v. Lamy, Bl. [2] 11, 210). Normal thallic sulphate Bl. [2] 11, 210). Normal thallic sulphate TL (SO.) 7aq is obtained as thin, white leaflets by dissolving Tl₂O₃.H₂O in fairly conc. H₂SO₄Aq and evaporating. Loses 6aq when heated to 200°, at higher temperatures SO, and O are given off and Tl₂SO₄ remains (Strecker, A. 135, 207; Crookes, C. N. 8, 243). The double salts Tl₂(SO₄)₂, M₂SO₄ [=TlM(SO₄)₂], where M=K or Na, are described by Strecker (l.c.).

Thorium sulphates. The hydrated normal

Thorium sulphates. The hydrated normal sulphate Th(SO₄)_x xaq is obtained by dissolving ThO₂ in slight excess of H_2SO_4Aq , and crystallising at $10^\circ-15^\circ$. The anhydrous salt is obtained by heating the hydrated salt to c. 400° (v. Chydenius, P. 119, 43; Delafontaine, A. 131, 100; Cleve, Bl. [2] 21, 115; Demarçay, C. R. 96, 1860). For a full discussion of the conditions of formation, relations, and solubilities of the various hydrates of $Th(SO_4)_2$ v. Roozeboom (Z. P. C. 5, 198). $Th(SO_4)_2$ has S.G. 4. 058 at 22·8° (Clarke, Am. 2, 175); 4·2252 at 17° (Krüss a. Nilson, B. 20, 1675). The S.G. of $Th(SO_4)_x$, 9aq is 2·767 according to Topsõe (Bl. [2] 21, 120). The hydrate $Th(SO_4)_x$, 9aq is isomorphous (monoclinic) with $U(SO_4)_x$, 9aq (Rammelsberg, B. B. 1886. 603). Double salts $Th(SO_4)_x$. M₂SO₄. xaq are known, where M = an alkali metal (Cleve, l.c.; Chydenius, l.c.).

Tin sulphates. The normal stannous salt SnSO, is obtained, in very small white crystals, by dissolving Sn in warm fairly cone. H₂SO₄Aq,

taking care that Sn is in excess, dissolving the white solid so formed in water, and allowing to crystallise out of contact with air (Marignac, Ann. M. [5] 15, 221). SnSO₄ is very sol. water; the solution soon deposits basic salts. The normal stannic salt Sn(SO₄)₂. 2aq was obtained by Ditte (C. R. 104, 178) by dissolving SnO₂. xaq (ppd. from SnCl₄Aq by alkali) in dilute H₂SO₄Aq and evaporating (v. also Kraskowitz, P. 35, 518). It forms white rhombic leaflets; very sol. water; decomposed by much water with separation of SnO₂. A basic salt SnO(SO₄). aq is said to be formed (Ditte, l.c.) by dissolving the normal salt in H₂SO₄Aq, warming, adding excess of Sn, and then ether.

Titanium sulphates. Normal titanous sulphate Ti₂(SO₄)₃, Saq is obtained, as violet crystals, by evaporating a solution of Ti in H₂SO₄Aq; sol. water, solution gives a black pp. when warmed (Glatzel, B. 9, 1833; Ebelmen, J. pr. 42, 76). Normal titanic sulphate Ti(SO₄)₂, Saq was obtained by Glatzel (l.c.) by oxidising the titanous salt by HNO₃, adding a drop or two H₂SO₄Aq, and evaporating; a white, translucent mass. A double salt Ti(SO₄)₂, K₂SO₄, Saq was obtained by Warren (P. 102, 449), by fusing TiO₂ with KHSO₄, treating with conc. H₂SO₄, evaporating, and washing the residue with cold water (v. also Glatzel, l.c.). By dissolving TiO₂ in hot H₂SO₄Aq, evaporating, drying the residue on a porous tile at 180°, and then heating to c. 350°, Merz (J. pr. 99, 157) obtained the acid salt TiO.SO₄ as a hard white solid.

Uranium sulphates. Normal uranic sulphate U(SO₂)₂. 9aq is obtained, as green monoclinic crystals isomorphous with Th(SO₂)₂. 9aq (Rammelsberg, B. B. 1886. 603), by evaporating a solution of UO₂ in excess of H₂SO₄Aq (R., l.c.; Ebelmen, J. pr. 27, 385). Unchanged in air; slowlyloses water of crystallisation when heated; at higher temperatures gives yellow (UO₂)SO₄, and when strongly heated leaves U₂O₈. Heated in H gives UO₂. Easily sol. dilute H₂SO₄Aq or HClAq. Decomposed by water, giving basic salts (v. R., l.c.; Athanasesco, C. R. 103, 271). Forms double salts with alkali sulphates (R., l.c.). The uranyl salt (UO₂)SO₄. xaq is obtained, in citron-yellow crystals, by dissolving U₂O₈ in H₂SO₄Aq with a little HNO₂; or by decomposing UO₂(NO₂)₂ by conc. H₂SO₄, evaporating nearly to dryness, dissolving in water, evaporating to a syrup, and allowing to crystallise slowly (Ebelmen, A. 56, 230). By evaporating a solution of the hydrated salt in conc. H₂SO₄, Schultz-Sellack obtained the anhydrous salt (UO₂)SO₄.M₂SO₄. 2aq (Rammelsberg, B. 5, 1005). Vanedium sulphates Uoradule sulphates Vanedium sulphates Vanedule sulphates

Vanadium sulphates. Vanadyl sulphate (VO)SO₄, xaq is obtained by heating a solution of V₂O₄ in conc. H₂SO₄. Gerlach (B. 11, 98) prepared several compounds of V₂O₅ and SO₂ by dissolving V₂O₅ in H₂SO₄. For more details and descriptions of various salts v. Vanadium tetroxide and pentoxide (infra).

Zinc sulphates. The normal salt ZnSO, is obtained by dissolving Zn, ZnO, or ZnCO, in dilute H,SO,Aq, evaporating, drying the crystals of ZnSO, 7aq, and gradually heating to c. 280° till water ceases to be given off; it is difficult to drive off every trace of water without decomposing some of the ZnSO, and forming

basic sulphates. Klobb (C. R. 114, 836) obtained ZnSO, in crystals by heating ZnSO. 7aq mixed with excess of (NH4)2SO4 in a partlyclosed crucible, protected from the gases of the flame, till all (NH₄)₂SO₄ was volatilised. A white solid; S.G. 3.435 at 16° (Pape, P. 120, 367); 3.6235 at 15° (Thorpe a. Watts, C. J. 37, 108). H.F. $[Zn,S,O^4] = 230,100$ (Th. 3, 516). Decomposed to ZnO, SO, and O at c. 400° (Bailey, C. J. 51, 681). Heated with charcoal, SO, and CO2 are evolved and ZnO remains; if the temperature is rapidly raised to a white heat, SO, and CO are given off and ZnS is formed. Heating in H produces an oxysulphide of Zn. For solubility in water v. infra. $[ZnSO_4,Aq] = 9,950$ (Th. 3, 516). For compounds of ZnSO, with NH, v. Schindler (Mag. Pharm. 31, 167; 36, 43), Kane (A. Ch. [2] 72, 290), Müller (Z. [2] 5, 250; 6, 96). Hydrated zinc sulphate. Several hydrates are known. The heptahydrate ZnSO. 7aq crystallises, at ordinary temperatures, in white, right rhombic, prisms, isomorphous with MgSO4. 7aq. S.G. 1-964 (Thorpe a. Watts, C. J. 37, 110). H.F. [Zn,S,O',7H'O] = 252,700 (Th. 3, 516). Melts at 50° (Tilden, C. J. 45, 267). At 100° loses 6H₂O. Poggiale (A. Ch. [3] 8, 467) gives solubility in water as follows:-

Temp,	S.	
	ZnSO. 7aq	ZnSO.
0.0	115.22	43.02
10	138-21	48·3 6
20	161.49	53.13
30	190.90	58.40
40	224.05	63.52
50	263.84	68.75
60	313.48	74.20
70	369.36	79.25
80	442.62	84.60
90	532.02	89.78
100	653.59	95.03

Gerlach (Fr. 8, 260) and Schiff (A. 110, 72) give the following data for ZnSO, Aq:-

S.G.

Gerlach, at 15°	Schiff, at 20.5°	Pctge. ZnSO. 7a
1.0288	1.0289	5
1.0593	1.0588	10
1.0905	1.0899	15
1.1236	1.1222	20
1.1574	1.1560	25
1.1933	1.1914	80
1.2315	1.2285	85
1.2709	1.2674	40
1.3100	1.3083	45
1.3532	1.3511	50
1.3986	1.3964	55
1.4451	1.4439	60

Almost insol. absolute alcohol; 100 parts of a saturated solution in alcohol of 40 p.c. contain

8'48 parts ZnSO, 7aq (Schiff, J. 1861, 87). Hexahydrate, ZnSO, 6aq; obtained by crystallising ZnSO, Aq at 40°; S.G. 2.07 (T. a. W., l.c.; v. also Marignac, J. 1855. 389). Pentahydrate, ZnSO, 5aq; obtained by boiling the finely-powdered heptahydrate with alcohol of S.G. 825; S.G. 2.206 (T. a. W., l.c.; v. also Kühn, J. 1830. 300; Schindler, Mag. Pharm. 31, 167; 36, 43; Pierre, A. Ch. [3] 16, 242).

Dihydrate, ZnSO, 2aq; formed by pouring cold gradually absorbed by conc. KOHAq of saturated ZnSO, Aq into conc. H₂SO, and wash-forming K₂(NO)₂SO, or Na₂(NO)₂SO.

ing the pp. with absolute alcohol; S.G. 2.958 (T. a. W., l.c.). Monohydrate, ZnSO, aq; obtained by heating ZnSO₄. 7aq to 100°-110° till it ceases to lose weight; S.G. 3·28 (T. a. W., *l.c.*;

v. also Graham, P. M. [3] 6, 827, 417).

An acid salt ZnSO, H.SO, 8aq is described by von Kobell (J. pr. 28, 492). Several basic salts are formed by boiling solutions of ZnSO, 7aq with ZnO or ZnO_.H₂ (v. Schindler, l.c.; Kühn, l.c.; Kane, A. Ch. [2] 72, 290; Reindel, Z. [2] 5, 508; Habermann, M. 5, 432; Athanasesco, C. R. 103, 271).

Double salts are numerous: (1) With alkali sulphates, ZnSO₄.M₂SO₄. xaq (v. Pierre, Ph. C. 1846. 410; Tobler, A. 95, 193; Graham, P. M.
[3] 6, 327, 417; Karsten, B. B. 1841). (2) With
CoSO., FeSO., MgSO., NiSO., &c., to form
ZnSO., MSO., xaq (v. Rammelsberg, P. 91, 321;
Pierre, Lc.; Etard, C. R. 86, 1399; 87, 602).

The normal salt Zirconium sulphates. Zr(SO₄)₂ is obtained by dissolving ZrO₂ in slightly diluted H2SO4, evaporating, and driving off excess of acid at a temperature below red heat. $Zr(SO_4)_2$ is decomposed by heating to redness, giving ZrO_2 . By concentrating a solution of $Zr(SO_4)_2$ containing some free H_2SO_4 , Paykull (B. 6, 1467) obtained crystals of the tetrahydrate Zr(SO₄)₂. 4aq (confirmed by Weibull, B. 20, 1394).

Basic salts are obtained by digesting a solution of a salt of Zr with saturated K₂SO₄Aq (v. Berzelius, P. 4, 117; Warren, J. pr. 75, 361).

Double salts with K₂SO₄ are formed by fusing KHSO₄ and ZrO₂ (B., l.c.; W., l.c.).

DITHIOPERSULPHATES. Under the name

of sodium dithiopersulphate, Villiers (C. R. 106, 851, 1354) described a salt Na₂S₄O₈. 5aq. This salt was said to be obtained by adding to sodium thiosulphate insufficient water to dissolve all the salt, saturating with SO₂, adding a little more water, again saturating with SO₂, keeping the yellow solution at the ordinary temperature for two or three days, passing in SO2 as long as it was absorbed, and, after standing a day or two, evaporating in vacuo over H2SO4. A mixture of Na₂S₄O₆ and Na₂S₂O₆ was thus obtained; on exposure to air the crystals of Na2S,O, effloresced, and the Na, S,O, was then picked out. The salt Na S.O. was described as white, lustrous, rhombio prisms; unchanged in air; melting at c. 125°. and giving off SO₂ at c. 140°, leaving Na₂SO₄ mixed with S. In a later paper (C. R. 108, 402) V. announced that the salt supposed by him to be Na₂S₄O₈ was only hydrated sodium tetra-

thionate Na.S.O. H.O.
NITROSOSULPHATES. In 1800 Davy (Researches, chiefly concerning Nitrous Oxyde) noticed that NO was absorbed by a mixture of Na₂SO₂Aq and NaOHAq, a compound being formed which, on addition of an acid, gave off N₂O while Na₂SO₄ remained in solution; he supposed that the NO was reduced to N₂O, the Na.SO, being oxidised to Na.SO,, and that the N2O combined with the NaOH. Pelouze (A. Ch. [2] 60, 151) found that a salt containing N, S. and O was formed in the reaction examined by Davy; to this salt he gave the composition

NazŠO,(NO),

A mixture of 1 vol. SO, and 2 vols. NO is gradually absorbed by conc. KOHAq or NaOHAq,

Ammonium nitrososulphate (NH₄)₂(NO)₂SO₃ is best prepared by passing NO for some hours into cold cone. (NH₄)₂SO₃Aq mixed with 5 to 6 times its volume of NH₄Aq. White crystals of the salt gradually form; they are washed with ice-cold NH,Aq, dried in vacuo, and kept in a well-stoppered bottle (Pelouze, l.c.). Potassium nitrososulphate and sodium nitrososulphate, M₂(NO)₂SO₃, are prepared similarly to the ammonium salt. Barium and lead nitrososulphates are obtained by ppg. aqueous solutions of the K salt by conc. BaOAq and solution of basic acetate of lead, respectively (Divers a. Haga, C. J. 47, 364). The nitrososulphates readily decompose; they are stable in solution only in presence of excess of alkali. Heated moist they give off N.O, leaving pure sulphate; heated dry they give NO and sulphite (v. D. a. H., C. J. 47, With acids, and also with most metallic salts in solution, also in contact with spongy Pt, charcoal, Ag.O, MnO₂, &c., they give N₂O and sulphates. Solution of an alkali nitrososulphate has no reaction with KMnO, Aq (D. a. H., l.c. p. 205). An alkaline solution of the K or Na salt is slowly reduced by Na amalgam, giving K₂SO₂Aq and K₂N₂O₂Aq, or the corresponding Na salts (D. a. H., l.c. p. 203).

PERSULPHATES. Persulphuric dride S2O, is formed at the anode during the electrolysis of fairly conc. H₂SO₄Aq (v. Sulphuric peroxide, under Sulphur oxides, p. 616). The acid corresponding with this oxide would be H₂S₂O₈ or HSO₄; this acid has not been isolated, but some of its salts have been prepared

by Marshall (C. J. 59, 771 [1891]).

Potassium persulphate K.S.O. is obtained by passing a current of 3 to 3 ampères, for some days, through saturated KHSO Aq contained in a Pt dish, wherein is suspended a porous cell containing dilute H₂SO₄Aq; the Pt dish stands in a vessel of copper, through which runs a stream of cold water, and which is connected with the battery so that the Pt dish becomes the anode; the cathode consists of a stout wire of Pt dipping into the dilute H2SO, Aq in the porous cell (for description, and diagram, of the apparatus v. Marshall, C. J. 59, 765-6). The granular salt that slowly separates during electrolysis is collected by filtering through Pt foil, dried on a porous plate, treated with hot water so as to obtain a nearly saturated solution, which is at once rapidly cooled (the mother-liquor yields $K_2S_2O_8$ more $K_2S_2O_8$ when again electrolysed). forms small, white, prismatic crystals; by spontaneous evaporation of a solution, large, flat tables are obtained, probably asymmetric. Slightly sol. cold water; S. at $0^{\circ} = 1.77$; insol. absolute alcohol, hot or cold. Measurements of the electrical conductivities of solutions of the salt indicated the formula KSO4, and not K2S2O8; but Bredig (Z. P. C. 12, 230) showed that Marshall's results were based on data, given by Ostwald, which were not quite accurate (the data were afterwards corrected in Ostwald's Lehrbuch [2nd ed.] 2, 730). Bredig's measurements established the formula K2S2O8; and this was confirmed by Löwenberg's determinations of the freezing-points and conductivities of solutions of the salt (Chem. Zeitung, 1892. 838). K,S,O,Aq very slowly decomposes at the ordinary temperature, giving KHSO Aq and O; in contact with

zine, decomposition is still very slow; with a Cu-Zn couple the rate of change is hastened; heating also hastens the reaction. After keeping for some time in a closed bottle a peculiar smell is noticed on opening the bottle, perhaps due to $H_2S_2O_8$ or S_2O_7 ; ozone is sometimes also given off. Heat decomposes K2S2O8; K2SO4 remains and SO, and O are given off; decomposition is marked at a little above 100°, but is not complete at 250°. When gently warmed with conc. HNO, Aq or H_2SO_4 , much ozone is given off; Cl is evolved when conc. HClAq is used. $K_2S_2O_8Aq$ does not yield pps. of persulphates with solutions of metallic salts; when pps. are formed they are due to reactions with K₂SO₄Aq formed by the decomposition of the K₂S₂O₄Aq. Solution of a salt of Ba slowly throws down BaSO4; the reaction is very slow, even when the solution is boiled. Solution of a salt of Pb also slowly forms PbSO. when heated with $K_2S_2O_8Aq$; if alkali is added, PbO₂ is ppd. AgNO₃Aq produces no pp. at once, but after a time black AgO, the solution becoming acid; several other metallic salts in presence of alkali yield pps. of peroxides with K.S.O.Aq. e.q. salts of Co. Cu, Mn, and Ni. KIAq is slowly decomposed by K2S2O8Aq, with separation of I; litmus and turmeric solutions are gradually bleached; paper and cloth become rotten when dipped in K2S2O8Aq. K4FeCy6Aq is oxidised to K.FeCy.Aq, and alcohol yields aldehyde, by warming with K₂S₂O₈Aq.

Ammonium persulphate (NH₄)2S2O8. pared, similarly to K2S2O8, by electrolysing dilute H_2SO_4Aq (c. 1 to 6 by volume) with $(NH_4)_2SO_4$; the salt is purified by cooling by ice a solution saturated at a little above the ordinary temperature. Crystallises in transparent lozengeshaped, apparently mono-symmetric tables. Very

sol. water; S. at $0^{\circ}=58\cdot2$. Reacts similarly to $K_2S_2O_8$ (cf. Elbs, J. pr. [2] 48, 185).

Barium persulphate $Ba_2(SO_4)_4$, 8aq. Prepared by rubbing in a mortar saturated (NH₄)₂S₂O₈Aq with excess of pure BaO₂H₂—Ba₂(SO₄)₄ goes into solution, and BaSO, also forms—passing a rapid stream of air till most of the NH, set free in the reaction is removed, placing in vacuo over H2SO4 till no free NH3 remains, passing in CO2 to remove excess of BaO,H, keeping in vacuo for a short time (to decompose Ba bicarbonate to BaCO₃), filtering from BaSO₄ and BaSO₅, evaporating in vacuo till crystallisation begins (with addition from time to time of a little BaOAq to neutralise H₂S₂O₈ that is set free during evaporation), dissolving the crystals that first separate in as little water as possible, filtering, cooling by ice, and drying the small prismatic crystals that form on a porous plate. crystals of Ba₂(SO₄), 8aq gradually become milky from formation of BaSO4, the change soon spreads, and the crystals crumble to a moist, powdery mass of BaSO₄. Ba₂(SO₄), 8aq is very sol. cold water; S. at 0°=52.2. Sol. absolute alcohol; on standing white crystals are deposited, probably Ba₂(SO₄)₄. 2aq.

Lead persulphate $Pb_2(SO_4)_4$. xaq; x=4 or 6, probably 6. Obtained by adding a slight excess of H₂SO₄Aq to conc. Ba₂(SO₄)₄Aq, neutralising with PbCO, filtering after some time, evaporating in vacuo over H₂SO₄ (removing PbSO₄ from time to time by filtration) till a solid mass is obtained, which is at once dried between filter paper and

placed in vacuo. Pb.(SO₄), was not obtained quite free from PbSO₄; it decomposes very readily, giving off pungent fumes, that separate

I from KIAq on paper.

Zinc persulphate was obtained, but not pure, by adding ZnSO, Aq to Ba₂(SO,), Aq in the proper proportions, filtering, and evaporating in vacuo over H2SO4. The copper salt was also obtained, but not free from CuSO. For thermal data concerning the formation and solution of the persulphates of NH,, Ba, K, and Na v. Berthelot (C. R. 114, 875; abstract in C. J. 62, 931).

PYROSULPHATES. (Disulphates. Anhydrosulphates.) Salts of the acid H₂S₂O₇ (v. Pyrosulphuric acid, under SULPHURIC ACID, p. 620). These salts bear to the sulphates a relation similar to that of the dichromates to the chromates. The sulphates may be written SO₂(OM)₂, and the pyrosulphates OM.SO₂.O.SO₂.OM. The sulphates and pyrosulphates may also be regarded as compounds of the acidic radicle SO₃ with basic radicles MO; thus sulphates MO.SO, pyrosulphates MO.2SO₃. Inasmuch as H₂S₂O₂ may be regarded as a partial anhydride of $H_2SO_4(2H_2SO_4 - H_2O = H_2S_2O_7)$, the acid $H_2S_2O_7$ is sometimes called anhydrosulphuric (cf. Acids, **v**ol. i. p. 50).

Potassium pyrosulphate K₂S₂O, is formed by heating K₂SO, with half its weight of H₂SO till acid ceases to come off at an incipient red heat. Prismatic needles; S.G. 2.277 (Jacquelain, A. Ch. [2] 70, 311). Melts at 300°, according to Schultz-Sellack (B.4, 109). Cannot be crystallised from water. Solution in fuming H₂SO, gives crystals of the acid salt KHS₂O₇, melting at 168° (S.-S., *l.c.*). Heated with alcoholic solution of KSH gives K₂SO₄, K₂S₂O₃, and H₂S; boiled with C₂H,ONa in alcohol gives KNaSO₄ and KEtSO, (Drechsel, J. pr. [2] 5, 367). The pyrosulphates of NH4, Ba, Ag, and Na are prepared similarly to $K_2S_2O_7$ (v. Schulze, B. 17, 2707; J., l.c.; S.-S., l.c.) M. M. P. M.

SULPHAZIDES X.NH.NH.SO2.Y.

Formation .- 1. By the action of alcoholic SO₂ upon diazo-compounds.—2. By the action of hydrazines upon sulphonic chlorides Y.SO₂Cl. 3. By the action of hydrazine hydrochlorides upon sulphinic acids Y.SO.H.—4. By reduction of the compounds X.N₂.SO₂.Y.

Reaction .- By heating with aqueous alkalis (c.g. baryta-water) they are decomposed into a sulphinic acid, a hydrocarbon, and nitrogen: X.N.H.2.SO₂.Y = XH + Y.SO₂H + N₂.

References: Koenigs, B. 10, 1531; Wiesinger, B. 10, 1715; Fischer, A. 190, 132; Escales, B. 18, 893; Limpricht, B. 20, 1238.

SULPHAZOTISED ACIDS v. SULPHUB OXY-

ACIDS, NITROGEN DERIVATIVES OF (p. 619). SULPHIDES. Binary compounds of sulphur.

The name is generally applied only to binary compounds of S with elements less negative than itself—that is, to compounds with elements other than Br, Cl, F, I, or O; thus SO₂ and SO₃ are called oxides of sulphur rather than sulphides of oxygen. Compounds of S with organic radicles, which compounds react similarly to sulphides of metals, must be regarded as binary compounds if the definition of the term sulphide is to be made as wide as possible. In the present article, however, only binary compounds of S with less negative elements are included. Sulphides of

all the metals, and also of the non-metals H, B, C, N, Si, P, Se, and Te, are known.

Many sulphides of metals occur native. Sulphides are frequently prepared by the direct union of the elements; sometimes by heating metallic oxides with excess of S, e.g. As₂S₃—in many cases a mixture of sulphide and oxide is formed in this way; addition of alkali carbonate to the mixture of metallic oxide and S sometimes brings about formation of sulphides, e.g. formation of sulphides of Cr and U; polysulphides mixed with sulphates are produced by heating the hydroxides or carbonates of alkali metals with S. Most heavy metals give sulphides when their oxides are heated in H2S gas, also when H.S is passed into solutions of their salts. Heating metals in H₂S gas often produces sulphides. Sulphides are also formed by heating one metal with the sulphide of another, more negative, metal. Reduction of sulphates, generally by heating in H or with C, frequently yields sulphides. Some metallic oxides yield sulphides when strongly heated in vapour of CS2. Sulphides of many heavy metals are formed by immersing the metals in $(NH_4)_2SAq$ containing excess of S (v. Priwoznik, A. 164, 46). Many metallic oxides yield sulphides by heating with dry $Na_2S_2O_3$ (v. Landauer, Fr. 1872, 427).

The sulphides of the alkali and alkaline earth metals are soluble in water; other sulphides are insoluble, or only very slightly soluble, in water.

Very dilute aqueous solutions of several metallic sulphides, in the colloidal form, were obtained by Winssinger (Bl. [2] 49, 452) by the following methods: (1) ppg. the sulphide by H₂S from an extremely dilute solution, and dialysing; (2) washing the ppd. sulphide for a long time with cold water, or with dilute H.SAq; (3) forming the sulphide in a solution free from all substances capable of causing coagulation. In these ways W. obtained colloidal soluble sulphides of Bi, Co, Au, Fe, In, Pb, Hg, Mo, Ni, Pd, Pt, Ag, Tl, W, and Zn. For preparation of colloidal soluble CdS v. Prost (C. C. 1888. 82), and of colloidal soluble CuS v. Spring a. de Boeck (Bl. [2] 48, 165).

Metallic sulphides are decomposed by strong acids, giving salts and H,S; secondary reactions frequently occur, S almost invariably separating. Several sulphides, e.g. As2S2 and HgS, are sublimable out of contact with air; all sulphides are oxidised by heating to redness in The sulphides of the very positive metals yield sulphates, and the sulphides of the heavy metals generally form oxides (the sulphides of Au, Hg, Pt, and Ag form metal) and give off SO. All sulphides of metals yield metallic chlorides and S₂CL₂ when heated in a stream of Cl. Many metallic sulphides are decomposed by water, at different temperatures, giving oxides or hydroxides and H2S (cf. de Clermont a. Frommel, C. R. 87, 330). Sulphides of non-metals are decomposed by water, generally to oxyacids and H₂S; NS gives NH₄ salts of H₂S₂O₂ and H₂S₃O₃. Many sulphides of heavy metals are decomposed by heating with water and I, giving iodides, S, and a little sulphate (v. Filhol a. Mellies, A. Ch. [4] 22, 58).

The double sulphides, formed by the union of sulphides of more positive elements with sulphides of less positive elements, are comparable

with the double oxides or oxysalts; most of these double sulphides are probably best regarded as salts of thio-acids corresponding with salts of oxyacids, but it is customary to apply this conception generally only to those compounds which contain sulphides of non-metals or of Sb, As, or Sn. The double sulphides, or thio- salts, are generally decomposed by water, forming oxides and H₂S. Many double sulphides, containing K₂S or Na₂S as one constituent, are formed by fusing metals or metallic sulphates with K₂CO₂, or Na₂CO₃, and S (v. Schneider, J. pr. [2] 7, 214; 9, 209; 10, 55). The sulphides may be classified, like the oxides, as monosulphides, disulphides, &c. It is better, however, to divide them into basic, acidic, indifferent, and persulphides; but this classification has been less developed and applied than in the case of the oxides. Basic sulphides react with acids to form salts and H2S, just as basic oxides react with acids to form salts and H2O. Acidic sulphides react with the basic sulphides to form thio-salts, as acidic oxides react with basic oxides to form oxy- salts; but few compounds have been isolated of acidic sulphides with H2S, corresponding with the compounds of acidic oxides with water which are oxyacids. The more strongly acidic sulphides—that is, the sulphides of non-metals—are decomposed by water, generally giving H2S and oxyacids. The persulphides have not been much studied; any sulphide of an element containing more S than the basic or acidic sulphide of the same element may be placed provisionally among the persulphides. A sulphide which does not react as basic or acidic, and which from its composition cannot well be called a persulphide, may be classed as an indifferent sulphide. The sulphides of C and P combine with several metallic sulphides to form thio-salts; double sulphides (or thio-salts) have not been isolated containing sulphides of B, N, Si, Se, or Te; the sulphides of Sb, As, and Sn combine with the sulphides of the alkali metals, and with some other sulphides of positive metals, to form thio- salts. The greater number of the double sulphides are formed by the union of two sulphides, both of which are metallic. The sulphide of that metal which, on the whole, is the more positive is regarded as the basic radicle; and the sulphide of the metal which, on the whole, is the less positive is regarded as the acidic radicle of the double sulphide. Some metallic sulphides must be classed both as basic and acidic; Cu,S, for example, combines with As2S3, and is therefore basic towards the distinctly acidic sulphide of As, but it also combines with K2S, and is therefore acidic towards the distinctly basic sulphide of K. It is impossible to divide the metallic sulphides into two dis-tinctly-marked classes; the relativity of the terms' basic' and 'acidic' becomes even more apparent in dealing with sulphides than in dealing with oxides. The sulphides BaS, BaS, CaS, and CaS, may be taken as examples of those which are loosely put together under the name persulphides. As examples of indifferent sulphides NS and Cr.S. may be mentioned. Many metallic sulphides which have been little studied—and which must, therefore, for a time be called indifferentare very probably basic in their reactions; such are Al₂S₂ and CdS. M. M. P. M.

SULPHIDO-DIACETIC ACID C.H.SO. (A. S(CH.CO.H). Thiodiglycollic acid. [129°]. S. 42 at 18°. Formed from chloro-acetic acid and alcoholic ammonium sulphide, aqueous Ca(SH)20 or Na₂S (Schulze, Z. 1865, 73; 1866, 184; Schreiber, J. pr. [2] 13, 472; Lovén, B. 17, 2818). Formed also by heating bromo-acetic acid with benzyl sulphide, allyl sulphide, or ethylene sulphide (Letts, Tr. E. 28, 612). Trimetric tables, v. sol. alcohol. Oxidised by KMnO₄ to SO₂(CH, SO₂H)₂. The Na salt is converted by treatment with sodium chloro-acetate into CO2H.CH2SMe CO [150°] (Delisle, The analogous compound B. 25, 2450). (CO₂H.CH₂)₂S CO_C CO_C SO [158°] is formed from chloro-acetic acid, Na₂CO₂, and Na₂S (Delisle),

and yields Na,A" 3aq.
Salts.—K₂A" aq: deliquescent prisms.—
KHA".—CaA". S. 2 at 21°.—BaA".—BaA" 5aq. - ZnA" 4aq. — PbA". — Pb₂A"O. — CuA" **a**q.-

Ag₂A".

Methyl ether Me.A". (135° at 11 mm.). Ethyl ether Et.A". (268° cor.). Formed from chloroacetic ether and alcoholic KSH Wislicenus, A. 146, 153).

Amids S(CH₂.CO.NH₂)₂. Formed chloro-acetic acid and alcoholic (NH₄)₂S. Formed from

acid S(CH₂.CO₂H).CH₂.CO.NH₂. Amic[125°]. Prisms, m. sol. cold water.—BaA'2 aq.-CaA'2 aq.—AgA': needles (from hot water).

Imide $S < CH_2 CO > NH$. [128°]. by heating the ammonium salt at 200°. Crystals.

Anhydride $S < \frac{CH_2 \cdot CO}{CH_2 \cdot CO} > 0$. [102°]. (158° at 10 mm.). Formed by boiling the acid with AcCl (Anschütz, A. 273, 68). Needles (from CHCl.).

Chloride S(CH...COCl)₂. Mono-anilide S(CH...CO₂H).CH...CONHPh.

Di-anilide S(CH, CO.NHPh), [168°], p-Toluide S(CH, CO, H).CH, CO.NHC, H,. [95°]. Needles.

Di-sulphido-di-acetic acid (di-thio-di-glycollic acid) S₂(CH₂.CO₂H)₂ [100° uncor.]. Formed from thio-glycollic acid CH₂(SH).CO₂H by Fe₂Cl₂ or by atmospheric oxidation of the alkaline solution (Claesson, B. 14, 410; Ginsburg a. Bondzynski, B. 19, 114). White plates or prisms. V. sol. water, alcohol, and ether, al. sol. benzene. Gives with AgNO, a white pp. AgHA". By tin and HCl it is reduced to thio-glycollic acid.—
K_A" 1½aq: easily soluble crystals.—KHA" aq.—

BaA" 4aq: amorphous pp.

Ethyl ether Et,A". (c. 280°).

Amide [155°]. Crystalline.

SULPHIDO-DI-ACETOACETIC ETHER

C₁₂H₁₈SO₈ i.e. S(CHAc.CO₂Et)₂, [90°] (S.); [84°] (D.); [76°] (B.). Formed by the action of S₂Cl₂(2 mols.) or of SCl₂ on sodium acetoacetic ether (1 mol.) suspended in benzene (Buchka, B. 18, 2092; 22, 2545, 2555; Delisle, B. 22, 306). Formed also from cupric acetoacetic ether and S in benzene (Schönbrodt, A. 253, 197). Needles, v. e. sol. benzene. Converted by KOH into sulphido-diacetic acid. Yields a crystalline phenyl-hydrazide C., H., N., SO., converted by warming with an alcoholic solution of phenylhydrazine into oxy-phenyl-methyl-pyrazole-azobenzene NPh.CO C:N.NHPh (Michaelis a. Phillips, B. 23, 560). Phenyl-hydrazine (2 mols.) added to sulphido-di-acetoacetic ether dissolved in cooled HOAc forms C₂₀H₁₈N₄SO₂, which yields B'HOAc [161°] and B'HOEt (Michaelis, B. 23, 2476; Sprague, C. J. 59, 332).

SULPHIDO-ANILINE v. DI-AMIDO-DI-PHENYL

BULPHIDE.

DISULPHIDO-DI-BENZOIC ACID

S₂(O₂H₁.CO₂H)₂. [244°]. Formed by oxidation of O₆H₄(SH).CO₂H by moist air or by bromine water (Frerichs, B. 7, 794; Hübner a. Upmann, Z. 1870, 294). Formed also from m-diazo-benzoic acid and H.S (Griess, J. pr. [2] 1, 102) and, in small quantity, by fusing sodium sulphobenzoate with sodium formate (Ador, B. 4, 622; Meyer, B. 6, 1150). Needles, nearly insol. hot water, sl. sol. alcohol.—(NH₄)₂A" 2aq.—CaA" 3aq.—BaA" 3aq.—PbA" aq.—Cu₂(OH)₂A" 5aq.— Ag₂A" 1 aq: yellowish-white pp.

γ-SULPHIDO-DIBUTYRIC ACID S(CH₂CH₂CH₂CO₂H)₂. [99°]. Formed by warming its nitrile with fuming HClAq (Gabriel, B.

23, 2492). Flat crystals.

Nitrile S(CH2.CH2.CH2.CN)2. (above 300°). Formed by boiling γ -chloro-butyronitrile with alcoholic K.S. Thick liquid.

a-Sulphido-dibutyric acid S(CHEt.CO₂H)₂. [105°]. Formed by heating a-bromo-butyric ether (2 mols.) with an alcoholic solution of KSH (1 mol.) and KOH (1 mol.), and saponifying the product (Loven, J. pr. [2] 33, 102). Tufts

of needles (from water).—BaA". a-Sulphido-di-isobutyric acid S(CMe₂.CO₂H)₂. Formed from a-bromo-isobutyric ether and alcoholic K,S, the product being saponified. Tablets (from water), insol. dilute H2SO4.-BaA" 2aq.

Di-γ-sulphido-dibutyric acid S₂(CH₂,CH₂,CO₂H)₂. [109°]. Formed by boiling the amide with conc. HClAq (Gabriel). Flat plates, v. sol. alkalis.

Amide S₂(C₃H₆.CO.NH₂)₂. [167°]. Formed by dissolving Cy.C₃H_a.S.Cy in cold conc. H₂SO₄.

SULPHIDO-DICINNAMIC ACID C18H14SO4 i.e. (PhCH:C(CO₂H))₂S. Formed by boiling sulphido-diacetic soid with Ac₂O and NaOAo (Loven, B. 18, 3242). Small needles (from alco-Yields C18H14Br2SO4 hol), insol. water. Na,A"2 aq: silvery plates.

Di-a-sulphido-di-cinnamic acid C22H22S2O4 i.e. (Ph.CH:C.CO₂H)₂S₂ [179°]. Formed by the oxidation of sulphydro-cinnamic acid Ph.CH:C(SH).CO₂H by iodine in alcoholic solution. Long yellowish needles, v. sol. alcohol, less in benzene (Ginsburg a. Bondzynski, B. 19, 123).

DI-SULPHIDO-DI-METHENYL-DI-AMIDO-DI-NAPHTHYL MERCAPTAN

 $S_2(C \stackrel{N}{\leqslant}_S > C_{10}H_6)_2$. Formed by oxidation of C₁₆H₆<NSC.SH with alkaline K₂FeCy₆ (Jacobson a. Frankenbacher, B. 24, 1408). The (α)-compound melts at 194°, the (β) - isomeride at 180°. DISULPHIDO - DI - METHENYL-DI-AMIDO-

DI-PHENYL MERCAPTAN $S_2(O < N > O_4H_4)_{2^*}$ [186°]. Formed by oxidising sulphydro-methenylamido phenyl mercaptan with K, Cr,O, and HOAc (Jacobson a. Frankenbacher, B. 24, 1404; cf. Hofmann, B. 20, 1789). Plates (from benzene).

DI - SULPHIDO - DI - METHENYL - DI -BENZENYL - DI - AMIDOSULPHIM

 $S_2(C \leqslant_{S,N}^N \geqslant_{CPh})_{s^*}$ [120°]. Formed by oxidation of C_eH_s . $C \leq_{N.S}^{N} > C.SH$ (Crayen, B. 24, 889). Stellate groups of needles, insol. water, sl. sol.

alcohol. Reduced by sodium-amalgam to the parent substance.

DI - SULPHIDO - DI - METHENYL - DI - p -TOLENYL-DI-AMIDOSULPHIM

 $S_2(C < N > C.C_6H_4Me)_2$. [169°]. Formed by oxidation of C,H,.C NN.S C.SH with HNO. free from nitrous acid (Crayen, B. 24, 392). Needles, insol. alcohol and ether, sol. hot C,H,.

DI-SULPHIDO-DI-METHYL-ANILINE v. Tetra-methyl-di-amido-di-phenyl-di-sulphide.

DI - SULPHIDO-DI-NAPHTHALENE-DI-(B)-SULPHONIC ACID S₂(C₁₀H_e.SO₂H)₂. Formed from the product of the action of potassium xanthate on diazotised (β) -naphthylamine (β) sulphonic acid by treatment with alcoholic potash (Leuckart, J. pr. [2] 41, 223). Colourless plates, v. sol. water and alcohol. Reduced by zinc-dust and dilute H_2SO_4 to $C_{10}H_6(SH).SO_8H$. K2A".—PbA": amorphous pp., sl. sol. water.

 $SULPHIDO-DI-(\beta)-NAPHTHYL-CARBAMIC$

ACID. Phenyl ether $S < \frac{C_{10}H_s}{C_{10}H_s} > N.CO.OPh.$ [215°]. S. (96 p.c. alcohol) '20 at 16°; S. (benzene) 1'20 at 16°. Formed from the chloride and alcoholic NaOPh (Paschkowezky, B. 24, 2916). Needles.

Chloride S: $(C_{10}H_6)_2$:N.COCl. [255°] Formed by heating β -imido-di-naphthyl sulphide with COCl₂ in toluene at 170°. Needles (from benzene-alcohol), v. sl. sol. ether. Converted by aniline into S:(C10H6)2:N.CO.NHPh, which is crystalline, S. (96 p.c. alcohol) 04; S. (benzene) ·18 at 17.5°, converted by boiling aniline into $S(C_{10}H_aNH_2)_2$ and $CO(NHPh)_2$. The chloride is converted by alcoholic NH_3 at 145° into S<C₁₀H₆ N.CO.NH₂, crystallising in needles, S (96 p.c. alcohol) '06; S. (benzene) '10 at 17.5°. The chloride $S < \stackrel{C_{10}H_6}{C_{10}H_6} > N.COCl$ is converted by β-imido-di-naphthyl sulphide in xylene at 280° into $(S < C_{10}H_{\bullet} > N)_2CO$, crystallising in yellowish plates and tables [over 350°].

SULPHIDO-DI-PHENYL-AMINE v. IMIDO-DI-PHENYL BULPHIDE.

SULPHIDO-DI-PHENYL-CARBAMIC ACID. Ethylether $S < \stackrel{C_0H_4}{\underset{C_0H_4}{}} > N.CO_2Et$. [110°]. Formed from imido-di-phenyl-sulphide and

ClCO₂Et (Fränckel, B. 18, 1845). Silky plates. Phenyl ether S:(C₆H₄)₂:N.CO₂Ph. [164°]. S. (96 p.c. alcohol) 24 at 16°; S. (benzene) 1.9 at 16°. Formed by heating the chloride

with NaOPh in alcohol on the water-bath (Paschkowezky, B. 24, 2908). Needles.

Chloride S:(C₆H₄)₂:N.COCl. [172°]. Formed from S:(C₆H₄)₂:NH and COCl₂ at 100°. Colourless prisms, sol. chloroform, and hot HOAc.

p-SULPHIDO-DI-PHENYL-DI-HYDRAZINE S(C₆H₄.NH.NH₂)₂. [115°]. Formed from S(C₆H₄.NH₂)₂ by diazotisation, treatment with NaHSO₃, and reduction with zinc-dust (Ruhl, B. 23, 3482; A. 270, 149). Yellowish leaflets, decomposing at 130°, sl. sol. cold water and ether, v. sol. alcohol. Reduces Fehling's solution in the cold. Benzoic aldehyde yields $S(C_6H_1, N_2H:CHPh)_2$ — B"2HCl. [209°]. -B"H,SO, [219°].—B"H,C,O,

SULPHIDO - PHENYL - NAPHTHYLAMINE $S < C_{10}H_6 > NH$. Imido-phenyl-naphthylamine. Formed by heating phenyl-naphthyl-amine with S at 240° (Kym, B. 23, 2464). The (a)-compound melts at 138°, the (B)-isomeride at 178°. Both form yellow crystals, m. sol. hot alcohol,

SULPHIDO - DI - PHENYL - DI - (\beta) - NAPH -**THYL-UREA** $N(C_{10}H_{7})_{2}.CO.N < C_{6}H_{4} > S.$ [225°]. S. (96 p.c. alcohol) ·12; S. (benzene) ·55 at 16°. Formed from S:(C₆H₄)₂:N.COCl and (C₁₀H₇)₂NH at 260° (Paschkowezky, B. 24, 2914). White nodules, sl. sol. hot alcohol, sol. hot benzene.

and give a deep blue solution in conc. H₂SO₄.

Sulphido - tri - phenyl - (β) - naphthyl - urea NPh $(C_{10}H_{7})$.CO.N: $(C_{6}H_{1})_{2}$:S. [170°]. S. (96 p.c. alcohol) :52; S. (benzene) 3:38 at 15°. Formed by heating sulphido-di-phenyl-carbamic chloride with phenyl-(\$\beta\$)-naphthylamine at 250° (P.).

SULPHIDO - DI - PHENYL - DI - TOLYL-DI-GUANIDINE (NHPh.C(NH).NH.C,H,)2S. [153°]. Formed from the corresponding thio-urea, HgO, and alcoholic NH₃ (Truhlar, B. 20, 675). Needles (from ether ligroin).—C₂₈H₂₈SN₆H₂PtCl₆.

Sulphido-tetra-phenyl-di-tolyl-di-guanidine (NHPh.C(NPh).NH.C,H₆)₂S. [106°]. Formed by warming an alcoholic solution of (NHPh.CS.NHC, H₆)₂S with aniline and HgO. Grey amorphous powder, v. e. sol. alcohol.

Disulphido - di - phenyl - tetra - tolyl-di-guanidine NPh:C</br>

NH.C,He.S.C,He.NH
C.NPh.

[119°]. Formed by heating the corresponding thio-urea with excess of aniline and HgO. Resinous mass, v. e. sol. alcohol.

SULPHIDO - DI - PHENYL - DI - TOLYL - DI-THIO-DI-UREA (NHPh.CS.NH.C,H,)2S. [134°]. Formed from $S(C_1H_d, NH_2)_2$ and phenyl-thiocarbimide (Truhlar, B.20, 670). Needles.

SULPHIDO-DI-PHENYL-UREA C19H10N2SO i.e. $S < {}^{C_0H_1}_{C_0H_2} > N.CO.NH_2$. [202°]. S. (96 p.c. alcohol at 17·5°) 3; S. (benzene) 2·7 at 17·5°. Formed from S:(C,H,)2:N.COCl and alcoholic NH, (Paschkowezky, B. 24, 2908). Plates.

Sulphido-tri-phenyl-urea

S<C.H. N.CO.NHPh. [169°]. S. (96 p.c. alcohol) 26 at 17.5°; S. (benzene) 3.74 at 17.5°. Formed from S(C₆H₄)₂NCOCl and aniline. Bluish needles, sol. alcohol.

Sulphido-tetra-phenyl-urea

S(C₆H₄)₂N.CO.NPh₂. Formed in like manner, using NPh.H. Hexagonal plates.

Di-sulphido-tetra-phenyl-urea

 $CO(N < \hat{C}_0H_1>S)$. [225°] (F.); [231°] (P.). S. (96 p.c. alcohol) 043 at 17°; 4.16 at 78°. by heating NH:(CoH.):S with di-tolyl sulphide with CS, (Truhlar, B. 20, 672).

S:(C_aH₄)₂:N.COCl (Fränckel, B. 18, 1848). Plates (from HOAc), v. sl. sol. ether and hot alcohol.

DI-8ULPHIDO-DI-PROPYL-DI-PHTHAL -AMIC ACID $C_{22}H_{24}N_2S_2O_6$ i.e. $S_2(C_8H_6.NH.CO.C_0H_4.CO_2H)_2$. [136°]. Formed by boiling (γ) -phthalimido-propyl sulphocyanide with a 10 p.c. solution of KOH (Gabriel a. Lauer, B. 23, 89). Plates (from HOAc), v. sol. alkalis.—K₂A": crystalline pp.

(a)-SULPHIDO-DIPROPIONIC ACID S(CHMe.CO₂H)₂. Thiodilactylic acid. [125°]. Formed, together with CH₃.CH(SH).CO₂H, by boiling potassium a-chloro-propionate with KSH (Schacht, A. 129, 4; Böttinger, A. 196, 106). Formed also by passing H,S into a solution of potassium pyruvate (Böttinger, B. 12, 1425) and by the action of CH₃.CH(SK).CO₂K CH_3 .CHCl.CO₂K (Lovén, J. pr. [2] 29, 373). Monoclinic prisms, v. e. sol. water, alcohol, and ether. Not affected by nascent hydrogen. Oxidised by dilute HNO, to SO₂(CHMe.CO₂H)₂. -K,A": deliquescent.—BaA": amorphous, v. sol. water (S.). Its solution on boiling deposits a crystalline salt, S. ·1 (L.).—Ag₂A": amorphous

Di-(a)-sulphido-dipropionic acid S₂(CHMe.CO₂H)₂. [142°]. Formed by oxidation of CH₃.CH(SH).CO₂H by I or FeCl₃ (S.; Böttinger, A. 196, 103; B. 16, 1047; Lovén, J. pr. [2] 29, 372). Needles, sl. sol. cold water, v. sol. alcohol Reduced by zinc and HClAq to $CH_3.CH(SH).CO_2H.$ — $(NH_4)_2A''.$ — $K_2A'' 2aq.$ — Ag₂A": amorphous pp.

Di-8-sulphido-dipropionic acid

S₂(CH₂,CH₂,CO₂H)₂. Formed by oxidation of CH₂(SH),CH₂,CO₂H (L.). Thin silvery plates. Formed by oxidation of

Tri-sulphido-di-propionic acid S₂(CHMe.CO₂H)₂. [95°]. Formed $\mathring{\text{CH}}_3.\text{CO.CO}_2\mathring{\text{H}}$ and $\mathring{\text{H}}_2\mathring{\text{S}}$ (Lovén, J. pr. [2] 47, 173). Plates, v. sol. hot water. Yields α-sulphopropionic acid on oxidation.

SULPHIDO-TOLUIDINE v. DI-AMIDO-DI-TOLYL SULPHIDE.

SULPHIDO - DI - TOLYL - DI - CARBAMIC ETHER S(C_eH₃Me.NH.CO₂Et)₂. [113°]. Formed from imido-di-tolyl sulphide and ClCO2Et (Truhlar, B. 20, 668). Needles, v. sol. alcohol.

DI-SULPHIDO-TETRA-TOLYL-DI-GUANI- $\begin{array}{ll} \textbf{DINE} & C(NH) {<} \begin{matrix} NH.C, H_{\theta}.S.C, H_{\theta}.NH \\ NH.C, H_{\theta}.S.C, H_{\theta}.NH \end{matrix} {>} C(NH). \end{array}$

[196°]. Formed by boiling the corresponding thio-urea with alcoholic NH, and HgO (Truhlar, B. 20, 673). White amorphous powder, v. sol. benzene and hot alcohol. -B'2H,PtCla: brown ${f amorphous}$ powder.

SULPHIDO-DI-TOLYL-THIO-UREA

 $S<_{C,H_s,NH}^{C,H_s,NH}>CS.$ [235°]. Formed from

S(C,H_e,NH₂)₂ and alcoholic CS

Sulphido-di-tolyl-di-thio-di-urea S(C,H,NH.CS.NH2)2. [121°]. Formed by evaporating the hydrochloride of di-amido-di-tolylsulphocyanide sulphide with ammonium (Truhlar, B. 20, 669). Amorphous powder, sl. sol. ether.

Di-sulphido-tetra-tolyl-di-thio-di-urea CS NH.C,H.S.C,H.NH CS. [231°]. Formed by boiling an alcoholic solution of di-p-amido-

White amorphous powder, nearly insol. ordinary

SULPHIDO-DI-TOLYL-DI-UREA

 $S(C_nH_2Me.NH.CO.NH_2)_2$. di-Formed from amido-di-tolyl sulphide, hydrochloride and potassium cyanate (Truhlar, B. 20, 669). Crystallises from benzene in needles [151°] containing C.H.

Di-sulphido-tetra-tolyl-di-urea

CO NH.C,H_s.S.C,H_s.NH CO. Formed from NH.C,H_s.S.C,H_s.NH S(C.H_s.NH₂)₂ and COCl₂ (Truhlar, B. 20, 671). White amorphous powder, sol. hot alcohol.

SULPHIDO-DI-ÎSOVALERIC ACID

S(C₄H₈.CO₂H)₂. Formed from bromo-isovaleric acid and K₂S (Lovén, *J. pr.* [2] 33, 102). Crystals.

-BaA": in soluble powder.

SULPHIMIDE SO₂.NHAq. Obtained only in aqueous solution, which is prepared by decomposing the Ag derivative, SO₂.NAg, with the proper quantity of dilute HClAq, and filtering. The compound SO₂.NAg is prepared by saturating well-cooled SO₂Cl₂ in 15-20 vols. CHCl₃, with dry NH₃, dissolving the pp. in water, acidifying with HNO₃, ppg. all Cl by AgNO₃Aq, filtering, neutralising with KOHAq, and adding more AgNO, Aq; the pp. thus obtained is purified by recrystallisation from water. The compound SO2.NAg is also obtained by heating SO₂(NH₂)₂ (v. Sulphamide, p. 567) to 200°-210°, dissolving in water, ppg. by AgNO₃Aq, and recrystallising the pp. from water. The solution in water of SO. NH is strongly acid; when dilute it may be boiled for a short time without decomposition; NH4.HSO4 is formed on evaporation, even below 40°; the solution is decomposed by warming with acids, giving H2SO4Aq and NH, Aq; excess of alkali reacts slowly even on boiling (W. Traube, B. 25, 2472; 26, 607). Metallic derivatives of SO₂.NH wherein H is replaced by NH., Ba, Ca, Pb, K, Ag, and Na are described by T. (cf. Imido-sulphonic acid and salts, under Sulphonic Acids and Derivatives, p. 600). M. M. P. M.

SULPHIMIDO-AMIDE SO2NH2-NH.NH2SO2. (Imido-sulphamide. Imido-sulphonamide. Prepared by adding Imido-sulphurylamide). ammonium carbamate very slowly to S2O5Cl2 well cooled in a small closed flask, allowing to stand for some time with a CaCl, tube fitted through the cork, then heating to 60° for some hours, washing out NH₄Cl by repeated treatment with dry alcohol saturated with NH3, dissolving the residue in a little NH3Aq, and placing the solution in ice. Forms white lustrous crystals; boiling NH, Aq transforms it wholly to ammonium imidosulphonate; with boiling conc. HClAq is completely changed to (NH,)₂SO₄; when boiled with NaOHAq gives off two-thirds of its Nas NH₃ and forms sodium imidosulphonate (Mente, A. 248, 263). M. M. P. M.

SULPHINDIGOTIC ACID v. INDIGO DI-SUL-

PHONIC ACID.

Alkyl iodides combine with SULPHINES. di-alkyl sulphides forming compounds R₂SR'I, from which moist Ag₂O produces strong bases R₂SR'.OH. In these bases the group (SR₂R') may be looked upon as a monovalent basylous radicle derived from the hypothetical sulphine (SH_a). Thus Me₂SEtI may be called di-methylethyl-sulphine iodide, but in this dictionary it

is described as the ethylo-iodide of DI-METHYL SULPHIDE. In the same manner Me, SCI may be called tri-methyl-sulphine chloride, but in this dictionary it is described as the methylo-chloride of DI-METHYL-SULPHIDE.

SULPHINIC ACIDS. Organic acids containing the group SO(OH) where S is united to carbon. They may be obtained by reducing the chlorides of the sulphonic acids, in alcoholic or ethereal solution, with zinc-dust (Otto, B. 9, 1584). Fatty sulphinic acids are formed by the action of SO_2 or of SO_3 on zinc alkyls (Hofmann, A. 102, 72; 106, 287). They are readily oxidised to sulphonic acids. The aromatic sulphinic ethers are readily oxidised by KMnO, and HOAo to sulphonic ethers (Otto a. Rossing, B. 19, 1224). By zinc and dilute H₂SO₄ sulphinic acids are reduced to mercaptans. The aromatic sulphinic duced to mercaptans. The aromatic sulphinic acids yield K₂SO₃ and hydrocarbons when fused with potash. The alkyl ethers of sulphinic acids X.SO.OR are formed by passing HCl into a solution of the acid in the corresponding alcohol, and also by the action of chloro-carbonic ethers Cl.CO.R upon the sodium sulphinate X.SO.ONa in cold alcohol, CO2 being evolved. The ethers are not formed by the action of alkyl halogenides upon sulphinates, for by this reaction the isomeric sulphones X.SO₂.R are produced instead (Otto a. Rossing, B. 18, 2493). H₂S passing through benzene sulphinic ether at 50° forms mercaptan, benzene sulphinic acid, benzene sulphonic acid, and other products (Otto a. Rossing, B. 20, 2275). COCl₂ converts sodium benzene sulphinate into the anhydride (Ph.SO),O, which is v. sol. ether and benzene, and decomposed by water and alcohol into benzene sulphinic acid and ether respectively (Otto, B. 20, 3337)

SULPHITES AND HYPOSULPHITES. SULPHITES are salts of sulphurous acid formed by replacing H in H.SO, by metals. All sulphites may be represented by the formula xMO.ySO, where MO stands for an equivalent of a metallic oxide; the normal sulphites belong to the forms $M_{2}^{1}SO_{3}$, $M_{2}^{11}SO_{3}$, $M_{3}^{111}SO_{3}$, and $M_{3}^{11}(SO_{3})_{2}$; . the greater number of the acidic sulphites are of the forms M¹HSO₃ and M¹¹(HSO₃)₂, a few which contain more than one equivalent of acidic radicle to one of basic radicle may be represented as M₂SO₃.xSO₂. Double sulphites are fairly numerous.

Many sulphites are prepared by passing SO. into water wherein metallic hydroxides or carbonates are dissolved or suspended; some are formed by double decomposition from the alkali sulphites. Several metallic sulphites that are insoluble in water can be prepared by the reaction between solutions of the sulphates and solution of Na2SO3; two main reactions occur, approximately in accordance with the equations (1) $M^{II}SO_4Aq + Na_2SO_3Aq = M^{II}SO_3 + Na_2SO_4Aq$, $(2)M^{11}SO_{2}Aq + 2H_{2}O = MO_{2}H_{2} + H_{2}SO_{3}Aq$, so that normal sulphites are formed in some cases and basic sulphites in other cases. This method of forming sulphites has been examined by Seubert a. Elten (Zeit. f. anorg. Chem. 4, 44): they find that normal sulphites are always formed when M=Ba, Cd, Ca, Fe¹¹, Pb, Ag, Sr, or Tl (with HgII a double sulphite HgSO, Na, SO, H2O was formed); that normal sulphites are formed at low temperatures, using the salts in the ratio of equal molecules, when M = Mg, Mn, Snu, U(UO2), or Zn, but that at higher temperatures. or with more dilute solutions, basic sulphites of these metals are produced; and that basic sulphites are always obtained when M = Al, Be, Bi, Cr, Co, Cu, Fe¹¹¹, or Ni.

The sulphites of the alkali metals, and the acid sulphites of the alkaline earths, are soluble in water; most other sulphites are insoluble. Sulphites in aqueous solution, and many in moist air, readily oxidise to sulphates; salts of other S oxyacids are sometimes formed, and S is frequently separated (cf. Pierre, C. R. 62, 460; 73, 749). Oxidisers, such as ClAq, HNO, Aq, &c., quickly convert sulphites into sulphates; sulphite solutions are, therefore, energetic reducers. Solutions of sulphites are readily reduced, giving H.S or metallic sulphides, by SnCl₂Aq, Zn and HClAq, &c. Heated with C, they are reduced to sulphides, sometimes to oxides. Sulphites decompose at a red heat to sulphides and sulphates, or to oxides and SO₂. Sulphites are decomposed by almost all acids, not by CO₂Aq or boric acid, giving off SO₂. Solutions of sulphites generally form thiosulphates by reacting with S, H2S, or alkali hydrosulphides. Sulphites give sulphates and thiosulphates when heated in SO₂ (Divers, C. J. 47, 205). Sulphites scarcely react with POCl₃, according to Divers (l.c.); a little metallic chloride and phosphate are formed, but no SOCl₂. Divers (i.c.; also C. J. 49, 577) contends that the normal sulphites have the constitution SO₂OM.M-that is, that they contain the sulphonic group SO₂OM-and that they are not thionyl compounds, SO(OM)2.

Aluminium sulphites. The only salt that has been definitely isolated is the basic sulphite Al₂O₃.SO₂. 4aq; obtained by dissolving freshly ppd. AlO₃H₃ in SO₂Aq, and heating to 74°, when the salt separates as a white powder. Heated in air gives off H₂O and SO₂, and leaves sulphate (Gougginsberg, A. 45, 132; confirmed by Röhrig,

J. pr. [2] 37, 217).

Ammonium sulphites. The normal salt (NH₄)₂SO₂, aq was found in the fumes from the residues from gas-works used in the preparation of $(NH_4)_2SO_4$ (Scheitz, Ar. Ph. [3] 5, 332); it is prepared by adding absolute alcohol to SO₂Aq made alkaline by NH₂Aq, or by passing moist SO, and moist NH, into absolute alcohol (Muspratt, P. M. [3] 30, 414). White monoclinic leaflets; sol. in 1 part water at 12° with disappearance of heat. Does not deliquesce in air, but oxidises gradually to sulphate (Marignac, Ann. M. [5] 12, 25). The acid salt NH. HSO, is obtained, as very deliquescent rhombic crystals, by saturating a solution of the normal salt with SO₂ and allowing to crystal-lise in vacuo; loses SO₂ at the ordinary temperature; S separates on exposure to light; a saturated solution heated to 150° in a sealed tube decomposes to S, H2SO4Aq, and (NH₄)₂SO₄Aq (Barbaglia a. Gucci, B. 13, 2325). The basic salt 3(NH₄)₂O.2SO₂. 2aq described by Muspratt could not be obtained by Marignac. Double salts are described under the various sulphites.

Barium sulphites. The normal salt BaSO. prepared by double decomposition from BaCl, Aq and Na, SO, Aq; a white powder, insol. water: may be crystallised in hexagonal forms from warm saturated SO, Aq; heated in a closed tube decomposes to BaSO, and BaS (Rammelsberg, P. 67, 391). Easily sol. HClAq (Johnson, C. N. 58, 155; cf. Hodges, ibid. 128).

Beryllium sulphites. No salt has been isolated; a solution of a sulphite (? BeSO_s) is obtained by dissolving BeO₂H₂ in SO₂Aq, but crystals do not form on evaporation alone or over H₄SO₄; addition of alcohol causes formation of a thick syrup (Atterberg, Bl. [2] 24, 358; cf. Berthier, A. Ch. [3] 7, 77).

Bismuth sulphites. A basic salt

2Bi₂O₂.3SO₂.5aq is formed by the action of conc. SO₂Aq on BiO₂H₂ (Röhrig, J. pr. [2] 37, 217; cf. Muspratt, P. M. [3] 30, 414).

Cadmium sulphites. 'The normal salt CdSO₂

is obtained by dissolving CdCO, in SO2Aq, and crystallising; white, indistinctly crystalline salt; scarcely sol. water; heated gives SO₂, CdO, CdS, and CdSO₄ (Rammelsberg, P. 67, 256). By treating Cd with SO₂Aq, filtering from CdS, and concentrating carefully, Fordos a. Gélis obtained the dihydrate CdSO, 2aq; the same salt was obtained by Muspratt (l.c.); confirmed by Röhrig (l.c.). Deniges (Bl. [3] 7, 569) failed to obtain this hydrate, but states that a trihydrate is produced by mixing equal vols. of 10 p.c. solutions of CdSO, acidified with acetic acid and Na SO₃. A double ammonium-cadmium sulphite CdSO₃. (NH₄)₂SO₃ was obtained by Rammelsberg (l.c.)

Calcium sulphites. The normal salt CaSO, 2aq is prepared by ppg. CaCl,Aq by Na,SO,Aq, dissolving in SO,Aq, and crystallising; it forms small lustrous crystals (Muspratt, l.c.). Also obtained by passing SO2 into water with CaCO, in suspension (Röhrig, l.c.). Heated to 80° gives 2CaSO, aq, and loses all water at 100° (R., l.c.). R. failed to isolate any acid salt.

Chromium sulphites. A basic chromic salt 2Cr₂O₃.3SO₂.16aq was obtained by Muspratt (P. M. [3] 30, 414), as a pale-green pp., by adding alcohol to solution of CrO,H, in SO,Aq; composition confirmed by Röhrig (J. pr. [2] 37, 217). Alkali sulphite solutions do not ppt. solutions of chromic salts even after long boiling (Berthier, A. Ch. [3] 7,77). By adding K₂SO₂Aq to CrCl₂Aq, Moberg (J. 1847–8. 418) obtained a reddish pp. which was perhaps a chromous sul-

Cobalt sulphites. The normal cobaltous salt CoSO₂. 5aq was obtained by Muspratt (l.c.), as a red granular pp., by passing SO, into water with CoCO, in suspension, boiling out air from the solution, and letting cool in a closed vessel. By evaporating in H, Rammelsberg (P. 67, 391) obtained the *trihydrate*; the existence of both hydrates has been confirmed by Röhrig (l.c.). Double salts CoSO₃, K₃SO₂ and 2CoSO₃, Na₂SO₃, CoO were obtained by Schultzer (I. 1865, 270) by heating CoSO₃ or CoCl with (J. 1865. 270) by heating CoSO, or CoCl, with alkali sulphite solutions. Double salts of cobaltic sulphite Co.(SO.). M.SO., where M = K or Na, were obtained by Geuther (A. 128, 157) by continued heating of CoO,H, with fresh quantities of conc. M2SOAq.

Copper sulphites. Neither cuprous nor cupric sulphite has been isolated, but several double salts of cuprous sulphite are known. The green liquid obtained by dissolving CuCO, (Berthier, 4 Ch [21 7 86) or CnO. H. (Bettoar 4 39 178) in SO₂Aq probably contains Cu₂SO₃; but this solution rapidly decomposes with separation of a red pp. and formation of CuSO₄Aq. The red solution formed by passing SO₂ into water with freshly prepared Cu₂SO₃(NH₄)₂SO₃ (v. infra) in solution was supposed to contain Cu₂SO₃ by Rogojski (J. pr. 53, 403); but Péan de Saint-Gilles obtained only Cu₂SO₃. CuSO₃. 2aq from this solution (A. Ch. [8] 42, 23), and this was confirmed by Svensson (B. 4, 713).

According to Newbury (Am. 14, 232), a basic cupric sulphite 6CuO.4SO₂. 3aq is formed by passing SO₂ into cold water with CuO₂H₂ in suspension, and sending a stream of air into the green solution so formed until a bright-yellow pp. is produced. When this basic salt is boiled with water for a few minutes CuSO₄Aq is formed, and a brown pp. is thrown down. The brown pp. is thought by N. to be Cu₂SO₃; on continued boiling with water it gives off SO₂, and Cu₂O

remains.

Double cupro-cupric sulphites. The dihydrated salt Cu₂SO₃.CuSO₃. 2aq was obtained by Chevreul (A. Ch. 83, 183) by heating CuO or CuCO₃ with SO₂Aq; it is also prepared by boiling CuSO₄Aq with solution of an alkali sulphite or thiosulphate, by boiling the green double salt Cu₂SO₃.(NH₄)₂SO₃ (infra) with water (P. de St.-G., l.c.), by passing SO2 into conc. Cu(C2H3O2)2Aq at 65° till the yellow pp. that forms is dissolved, and then letting stand in air (Étard, C. R. 93, 725), and by passing a slow stream of SO₂ into 10 p.c. CuSO, Aq containing a roll of thin sheet Cu till the liquid is almost colourless (the salt deposits on the Cu) (Newbury, Am. 14, 232). Translucent, garnet-red octahedra; S.G. 3.57; gives up 2H₂O, and also SO₂, above 150°, and at a higher temperature leaves Cu₂O with some CuSO₄ (Rammelsberg, P. 67, 391). Soluble SO2Aq, HClAq, and NH2Aq; KOHAq ppts. hydrates of CuO and Cu2O; heated with water to 200° gives CuSO, Aq and Cu in crystalline leaflets (Geitner, A. 129, 850). The pentahydrate Cu₂SO₂.CuSO₂. 5aq separates, as a greenish-yellow pp., from conc. Cu(C₂H₂O₂),Aq, into which a slow stream of SO, has been passed till the liquid has become emerald-green (P. de St.-G., l.c.). Easily sol. SO2Aq or acetic acid, also in solutions of cupric salts and in NH,Aq; when boiled with water the red dihydrate is formed.

Double salts of cuprous sulphite with alkali sulphites are readily produced by the reaction of conc. solutions of alkali sulphites with solutions of cupric salts. These double salts are colourless and crystalline; their composition is xCu₂SO₃.yM₂SO₃.xaq (x generally = 1), where M = NH, K, or Na (for details of preparation, composition, and properties v. Rogojski, J. pr. 53, 403; Péan de Saint-Gilles, A. Ch. [8] 42, 23; Vohl, J. pr. 95, 218; and Svensson, B. 4, 713). Some acid salts of the form Cu₂SO₃.xM₂SO₂.ySO₂.xaq have also been prepared (S., l.c.), and some salts of alkali sulphites with both Cu₂SO₃ and CuSO₂ (P. de St.-G., l.c.; R., l.c.).

Didymium sulphites. The normal salt Di₂(SO₂)₂ was obtained, as a reddish-white powder, by passing SO₂ into water with Di₂O₂ in suspension, and boiling off excess of SO₂ (Marignac, A. Ch. [3] 58, 148; Cleve, Bl. [2] 89,

161).

Gold sulphites. No gold sulphite has Double salts of Au,80, been isolated. Am₂SO₈ and Na₂SO₂ of the form Au₂SO₂.3M₂SO₂. 3aq are known; also a Ba salt Au₂SO₃.3BaSO₃. aq. The Na salt is obtained by adding Na2SO2Aq to a boiling alkaline solution of NaAuO₂, or by saturating NaAuO₂Aq at 50° with SO₂, then carefully adding BaCl₂Aq to the solution till free H2SO, and H2SO, are ppd., filtering, adding more BaCl2Aq, quickly filtering off the purple-red Au₂SO₃.3BaSO₃. aq that separates, washing the pp. rapidly out of contact with air, decomposing it by the proper quantity of Na₂CO₂Aq, adding some alcohol to the solution, filtering off any Ba salt that ppts., and adding more alcohol, when the Na salt separates as an orange-red, very easily decomposed, pp. (v. Himly, A. 56, 252; 59, 95). When AuCl, Aq is added, drop by drop, to a warm solution of (NH4)2SO2 in conc. NH2Aq, white, lustrous, six-sided plates separate of the salt $(NH_4)_2SO_3.3(NH_3Au)_2SO_3$. 3aq; the mother-liquor from this salt yields the salt Au₂SO₃.3(NH₄)₂SO₃. 3aq, very similar to the corresponding Na salt (Haase, Z. 1869. 535).

Indium sulphites. The only salt that has been isolated is a basic salt 2In₂O₃.3SO₂.8aq. Obtained by boiling a solution of a salt of In with excess of NaHSO₃Aq; insol. water, sol. in dilute acids; loses 3H₂O at 100°, and all water at 260°; decomposed at 280°, giving off SO₂ and

leaving In₂O₃ (Bayer, A. 158, 372).

Iridium sulphites. The normal iridic salt Ir₂(SO₃)₃, 6aq is obtained, in yellow crystals, by passing SO₂ into water with IrO₄H₄ in suspension, filtering and evaporating. The salt loses all water at 160°-180°; at a higher temperature Ir₂O₃ remains; sl. sol. water, easily sol. acids; decomposed by boiling with KOHAq with separation of Ir₂O₄ (Birnbaum, A. 136, 179). The insoluble matter that remains on treating IrO₅H₅ in water with SO₂ is a basic salt Ir₂O₃.SO₂ 4aq (Birnbaum, l.c.). Double salts of iridous sulphite were obtained by Seubert (B. 11,1761) in separating Ir from Rh by means of Na₂SO₃ (by Bunsen's method, A. 146, 274). Seubert gave the formulæ IrSO₂.3Na₂SO₃. 10aq and IrH₂(SO₂)₂.3Na₂SO₃. xaq, x being = 4 and 10, to the salts he prepared.

Iron sulphites. The normal ferrous salt FeSO, 3aq is formed, along with FeS₂O₃, by dissolving iron wire in SO₂Aq out of contact with air on evaporating (out of air) the sulphite crystallises out, leaving the thiosulphate in solution (Fordos a. Gélis, J. Ph. [3] 4, 333; cf. Koene, P. 63, 245, 631; also Muspratt, P. M. [3] 30, 414). Sl. sol. water, easily sol. SO₂Aq; loses H₂O, and then (at 250°) SO₂, on heating. Basic ferric salts are formed by reacting on ferric salt solutions with alkali sulphites; on adding alcohol to the red liquids thus formed, Koene (P. 63, 245, 431) obtained the salts Fe₂O₂, SO₂, 6aq and 2Fe₂O₃, 3SO₂ (cf. Muspratt, L.c.). By passing SO₂ into water with FeO₃H₂ in suspension, and adding KOHAq, Muspratt (L.c.) obtained the double salt Fe₂O₃, SO₂, 2K₂SO₃, 5aq.

Lead sulphites. Only the normal salt PbSO₂ has been obtained; a white powder, insol. water, formed by adding Na₂SO₂Aq to-solution of a salt of Pb (confirmed by Röhrig.

J. pr. [2] 87, 217).

Lithium sulphites. The normal salt Li.SO₂, aq is prepared by passing SO₂ into water with Li₂CO₂ in suspension, till CO₂ ceases to be given off, and evaporating on a water-bath or over H₂SO₄ (Röhrig, J. pr. [2] 37, 217). Danson (C. J. 2, 205) represented the salt as a hexahydrate. By adding ether to the solution obtained from Li₂CO₃, as described, R. got the dihydrate Li₂SO₃, 2aq. Easily sol. water, somewhat sol. alcohol, very sl. sol. ether; oxidised in moist air. Röhrig failed to obtain an acid salt. By adding the proper quantity of K₂CO₃ to the acid solution of Li₂CO₃ in SO₂Aq, evaporating to a syrup, and placing in a freezing mixture of snow and salt, R. obtained the double salt 2LiKSO₃, aq; by a similar process he got monoclinic crystals of Cli₂SO₃.Na₂SO₃. Saq (J. pr. [2] 37, 217).

Magnesium sulphites. The normal salt

Magnesium sulphites. The normal salt MgSO₂, xaq is formed by suspending magnesia alba in water, passing in SO₂, and concentrating the solution (Rammelsberg, P. 52, 89); if evaporation is carried on over H₂SO₄ (Röhrig, l.c.), or in vacuo below 100° (Hartog, C. R. 104, 1793), the hexahydrate separates; if evaporation proceeds above 100° the trihydrate is obtained (Muspratt, l.c.; R., l.c.; H., l.c.). Slowly oxidises in air to sulphate; loses all water at 200°; when more strongly heated the salt loses SO₂ and

leaves MgO.

Double salts with (NH₄)₂SO₃ are formed by mixing the constituents and evaporating; also by dissolving MgO₂H₂ in cold (NH₄)₂SO₃Aq (Rammelsberg, P. 94, 507; Hartog, Lc.). Manganese sulphites. Several hydrates of

the normal manganous salt MnSO, have been described. By adding alkali sulphite solution to dilute MnCl2Aq so long as the pp. that forms dissolves again, and then letting stand, monoclinic crystals of the trihydrate MnSO₃. 3aq are obtained; this hydrate is also formed by mixing 10 p.c. of MnSO, Aq, acidified with acetic acid, and Na₂SO₈ (Deniges, Bl. [3] 7, 569); if Na₂SO₃Aq is added to hot dilute MnCl₂Aq the monohydrate is said to be produced. (Bammelsberg (P. 67, 245, 391) gives the formula 2MnSO_s. 5aq to the salt obtained by reacting on Mn acetate solution with Na, SO, Aq, and this is confirmed by Röhrig (J. pr. [2] 37, 217; cf. also Muspratt, P. M. [3] 80, 414). Double salts MnSO₂.K₂SO₃ and 2MnSO₂.K₂SO₃ were prepared by Gorgeu (C. R. 96, 376) by saturating 20 p.c. K₂SO₂Aq with SO₂, adding 4 p.c. MnSO₃. 3aq, and evaporating over an absorbent of SO2; double Mn-Na salts are also described by G. (l.c.).

Mercury sulphites. According to Péan de Saint-Gilles (C. R. 34, 905) the normal mercuric salt HgSO, is formed by adding dilute Na₂SO₂Aq to a very conc. solution of Hg(NO₂)₂ free from excess of acid; it is, however, difficult to obtain the salt free from basic sulphites. The basic mercuric sulphite 2HgO.SO₂ was obtained pure by P. de St.-G. by using solution of basic mercuric nitrate. The normal salt is easily decomposed; boiling water produces Hg₂SO₂ and Hg. Boiling water is said (P. de St.-G., l.c.) to convert the basic salt into the isomeric compound Hg₂SO₄ (2HgO.SO₂-Hg₂O.SO₃). Divers a. Shimidzu (C. J. 49, 538) say that normal mercuric sulphite cannot be isolated; the reaction of Na₂SO₂Aq with Hg(NO₂)₂Aq produces at first the basic salt 2[2HgO.SO₂]. aq, and then a

salt which is most simply represented as HgSO₂.Hg₂SO₂. 4aq, and which is called mercurosic (or mercuro-mercuric) sulphite by D. a. S., and represented by them as

 $H_g < SO_2O > H_{g_2}$ 4aq. D. a. S. represent the

basic salt as $Hg < SO_2OHgO > Hg. aq$, and call it mercuric oxysulphite.

For the best conditions of preparation of 2HgO.SO₂ v. D. a. S., l.c., p. 550, and for the reactions of this salt v. ibid. pp. 546-50; for preparation of mercurosic sulphite v. ibid. p. 564, and for the reactions of this salt, pp. 559-63.

By adding dilute Na₂SO₃Aq to moist Hg₂SO₄ or to HgNO_sAq, D. a. S. (l.c., p. 572) obtained a greyish-black amorphous solid to which they gave the formula $Hg_i(SO_s)_2$ aq, the constitution Hg SO₂O Hg₃. aq, and the name mercuric hypomercurous sulphite or hypomercurosic sulphite. For the best method of preparing this salt (action of SO₂Aq on Hg₂SO₄ or HgNO₃) v. D. a. S., l.c., p. 571; for the reactions of the salt v. pp. 567-70. It is evident that the formula given to this sait is the same as that of normal mercurous sulphite $(Hg_4(SO_3)_2 = 2Hg_2SO_3)$; the salt is produced from mercurous compounds, and in many of its reactions yields mercurous compounds; D. a. S. insist that the salt belongs to a new class of Hg compounds that contain what they call the 'hypomercurous radicle.'

By treating solid HgCl₂ with NaHSO₃Aq, Wicke (A. 95, 176) obtained a solution that deposited a white crystalline powder to which he gave the composition $Hg(HSO_s)_2$ —*i.e. acid mercuric sulphite.* According to D. a. S. (*l.c.*, p. 554) the pp. is $HgSO_s$.Na₂SO_s.aq. By reacting on HgO with SO2Aq a white solid is formed, together with a solution containing a compound of Hg. Rammelsberg thought that the white solid was a basic mercurous sulphite of varying composition, and that the solution contained Hg₂SO, (P. 67, 405; the reaction was also examined by P. de St.-G. and by Vogel). D. a. S. say that the white residue is $HgSO_3$. Hg_2SO_3 .4aq (mercurosic sulphite, according to D. a. S.); by using freshly ppd. HgO suspended in water, and adding a little of this to SO2Aq (nearly free from H₂SO₄), D. a. S. obtained a clear solution which reacted as a solution of HgSO₃ in H₂SO₃, and which they regarded as a solution of acid mercuric sulphite, Hg(HSO₃)₂ (v. l.c., pp. 554-8). Several double salts of mercuric sulphite with alkali sulphites have been isolated. The chief are $HgSO_3$, M_2SO_3 , xaq, where $M = NH_4$, K, or Na, and 2HgSO₃.Na, SO₃. aq (v. P. de St.-G., C. R. 34, 905; Hirzel, A. 84, 258; D. a. S., l.c., pp. 538-46). D. a. S. regard these double salts as alkali derivatives of the hypothetical acid Hg SO OH (they greatly doubt the existence of the salt 2HgSO, Na, SO, described by P. de St.-G.); this view is confirmed by Barth's measurements of the electrolytic conductivities, and the freezing-points, of solutions of these salts (Z. P. C. 9, 176); Barth's results point to the dissociation in dilute aqueous solution of these

salts into three ions, HgS₂O₆, R, and R'.

Nickel sulphites. The normal salt

NiSO₂, Saq separates, in green tetrahedric crys.

tals, on slowly evaporating a solution formed by passing SO₂ into water with NiO. xaq in suspension (Rammelsberg, P. 67, 391). By boiling the solution of NiO. xaq in SO₂Aq, Muspratt (P. M. [8] 30, 414) obtained a tetrahydrate NiSO₃. 4aq; this hydrate was also obtained by Röhrig (J. pr. [2] 37, 217) by quickly evaporating the solution at c. 150°. The compound with ammonia, NiSO₃.3NH₃. 3aq, separates, as a blue crystalline pp., on adding alcohol to an ammoniacal solution of NiSO₄ (Rammelsberg, Lc.).

of NiSO₃ (Rammelsberg, l.c.).

Osmium sulphites. The normal salt OsSO₃ was obtained by Claus (J. pr. 90, 65) by treating OsO₄Aq with excess of SO₂, and evaporating or adding Na₂CO₃ or Na₂SO₄; after drying the salt is a blackish-blue powder, insol. water, sol. HClAq without giving off SO₂, reppd. from this solution by KOH or K₂CO₃, decomposed to K₂SO₃ and OsO. xaq by long boiling with conc. KOHAq. When moist the salt oxidises in air to OsSO₄. The double salt OsSO₃.2K₂SO₄.2KHSO₃.4aq was obtained by Claus (A. 63, 355) as a pale rose-red crystalline powder, by heating K₂OsCl₆ with K.SO.Ag.

Palladium sulphites. Only the double pallado-sodium salt PdSO₃.8Na₂SO₃.2aq has been isolated; a white pp. obtained by dropping NaOHAqinto PdCl₂Aq, after adding SO₂ (Wöhler,

A. 174, 199).

Neither platinosul-Platinum sulphites. phite PtSO, nor platinisulphite Pt(SO₃)₂ has been isolated. Several salts which may be regarded as double compounds of PtSO₃, but are better classed as platinosulphonates or platinosulphites, have been obtained; they are shortly described under Platino-sulphonates (this vol., o. **2**85). By passing SO₂ into water with PtO,H, in suspension, and adding alkali sulphites to the dark-red liquid so formed, Birnbaum (A. 139, 172) obtained double salts of the form $PtO.SO_s.xM_2SO_s.yaq$, where M=K or Na, x varied from 1 to 2, and y also from 1 to 2. These salts gave no pp. of BaSO, with BaCl2Aq, with HClAq SO₂ was given off and PtCl, formed; hence they are better classed as double salts of platinyl sulphite, PtO.SO_s, than of platinous

sulphate PtSO.

Potassium sulphites. The normal salt K2SO3. 2aq is obtained, in large, deliquescent, monoclinic prisms, by passing SO₂ into K₂CO₃Aq as long as CO₂ is given off, and evaporating over H2SO4 at the ordinary temperature (Muspratt, P. M. [8] 30, 414). More soluble cold than hot water; decomposed by heat to K2S and K2SO4 (Rammelsberg, A. 50, 259; results of M. and R. confirmed by Röhrig, J. pr. [2] 87, 217; v. also Hartog, C. R. 109, 179, 221, 436). The acid salt KHSO, separates in white needles on adding alcohol to a solution of the normal salt saturated with SO₂ (M., l.c.; confirmed by Röhrig, l.c.). By passing SO, into hot saturated K₂CO, till CO, ceased to come off and the solution was greenish, and then allowing to cool, Muspratt (l.c.) obtained hard monoclinic crystals of the acid salt $K_2SO_3.SO_2$ (= $K_2S_2O_5$), sometimes called pyrosulphite; only sl. sol. water; decomposed by heat to K₂SO₄, SO₄, and S (confirmed by Röhrig, l.c.). Dilute K₂SO₄Aq heated in a closed tube to 100° and then let stand is very slowly decomposed to K,SO, H,S,O, and S (Saint-Pierre, C. R. 62, 460; 73, 749). K,SO,Aq with KNO,Aq gives K salts of various acids; v. Sulphur oxyacids, nitrogen derivatives of (p. 619).

Double sulphites of potassium and sodium. Schwicker (B. 22, 1728) obtained two distinct salts KNaSO_s. xaq; one by neutralising conc. KHSO, Aq by the proper quantity of Na₂CO₂ and evaporating over H₂SO₃, the other by adding K₂CO₃ to NaHSO₃Aq. Röhrig (*J. pr.* [2] 37, 217) ppts. the salts by adding alcohol, or better ether. The first salt, heated with EtI to 140°, gives 4(SO₂Et.OK).NaI, the second gives 4(SO,Et.ONa).KI; hence the first salt seems to be NaSO2.OK and the other KSO2.ONa. Schwicker assigns different quantities of crystalline water to the salts, viz. NaSO₂.OK. 2aq and KSO₂.ONa. aq; Röhrig (*l.c.*) gives 2aq to both salts. The two salts seem to be isomeric. Aqueous solutions of these salts have identical electrolytic conductivities, as would be expected (Barth, Z.P.C. 9, 176). Schwicker (l.c.) also describes the salts KNa,H(SO₃)₂. 4aq and K₂NaH(SO₃)₂. 3aq. Hartog (C. R. 109, 179, 221, 436) describes the salts $K_2SO_3.2Na_2SO_3.SO_2.9aq (=K_2O.2Na_2O.4SO_2.9aq)$ and $2K_2SO_3.(NH_1)_2SO_3.SO_2.9aq$ $(=2K_2O.(NH_4)_2O.4SO_2.9aq).$

For other double salts of potassium sulphite v. supra, cobalt sulphites, copper sulphites, iron sulphites, lithium sulphites, manganese sulphites, mercury sulphites, osmium sulphites, and platinum sulphites; and infra, ruthenium sulphites and silver sulphites.

Rhodium sulphites. The normal rhodic salt Rh.(SO₃)₃. 6aq is prepared, as a yellowish crystalline mass, by dissolving RhO₃H₂, aq in SO₂Aq and evaporating; fairly sol. water, insol. alcohol (Bunsen, A. 146, 265). The double rhodous salt 4RhSO₃.6Na₂SO₃, 9aq is obtained by treating Na₃RhCl₃. 12aq with excess of NaHSO₃Aq (Bunsen, l.c.); it has been examined by Seubert a. Kobbé (B. 23, 2556), who prepared it by heating a solution of RhCl₃. 4aq with excess of NaHSO₃Aq for some time, washing the pp. with cold water, and drying over P₂O₃. Pale-yellow solid; very sl. sol. hot water; dissolves in dilute HNO₃Aq, giving off SO₂; warmed with conc. H₂SO₄ forms Rh₂(SO₄)₃. Na₂SO₄ (B., l.c.; S. a. K., l.c.).

Ruthenium sulphites. Only the double rutheno-potassium salt RuSO₃.K₂SO₃ has been isolated; prepared by heating K₂RuCl₃Aq with K₂SO₃Aq till a dark-red solution is formed, evaporating to dryness, dissolving, again evaporating to dryness, and repeating these processes till a pure white residue is obtained (Claus, J. pr. 42, 351).

Silver sulphites. The normal salt Ag₂SO₃ forms small, white lustrous crystals; obtained by treating AgNO₂Aq with alkali sulphite solution or with SO₂Aq; if excess of SO₂Aq is allowed to remain in contact with the salt for some time Ag is formed (H. Rose, P. 33, 240). Darkens in air (Muspratt, P. M. [3] 30, 414; Sodeau, C. N. 65, 102); products depend on method of preparation of the salt (v. Stas, Stas Nouv. R.). Heated alone, or with water, to 100° gives Ag₂SO₄, Ag, and SO₂ (Berthier, A. Ch. [3] 7, 82; cf. Geitner, J. 1864. 142, who says that decomposition begins at 200°). Scarcely sol. water or SO₂Aq; sol. NH₂Aq (Berthier, Lo.), also in alkali sulphite solutions.

Double salts. AgNaSO₂, 2aq; small lustrous needles, formed by dissolving Ag₂SO₃ in hot saturated Na₂SO₃Aq, letting cool, and pressing salt that separates (cannot be washed with water) (Svensson, B. 4, 714). AgKSO, xaq, prepared like the Na salt. Svensson (l.c.) describes three double salts with (NH₄)₂SO₃, obtained by dissolving Ag2SO3, or AgOl, in (NH_{.)},SO_{.A}q; he gives the formulæ Ag(NH₄)SO_{.3}, Ag_{.5}SO_{.4}.6(NH₄)₂SO_{.3}. 19aq, and Ag_{.2}SO_{.3}.3(NH₄)₂SO_{.3}.4(NH₄)HSO_{.3}. 18aq. Sodium sulphites. The normal salt Na_{.2}SO_{.3}

is prepared by completely saturating Na2CO2Aq with SO2, warming, adding an equal quantity of the same Na₂CO₂Aq, evaporating, and crystallising above 33° (Rammelsberg, P. 56, 298; Schultz-Sellack, J. pr. [2] 2, 459). Also obtained by heating the hydrate Na₂SO₂. 7aq to 150°. Na, SO, also crystallises from solution of NaHSO, that has been made strongly alkaline. Heated above 150° melts to a yellowish-red mass containing sulphide and sulphate in the ratio Na₂S:3Na₂SO₄ (R., P. 67, 246; 94, 507). The heptahydrate Na₂SO₃. 7aq is obtained by crystallising at the ordinary temperature a solution of Na₂CO₂ saturated with SO₂, warmed, and mixed with an equal quantity of the same Na,CO,Aq; monoclinic prisms; S.G. 1.561 (Buignet, J. 14, 15). Effloresces and partially oxidises in air. Very sol. water; maximum solubility at 33°. Another hydrate, Na SO 10aq, was described by Muspratt (P. M. [3] 30, 414) as formed by evaporating Na, 80, Aq over H2SO,; according to Röhrig (J. pr. [2] 37, 217) only the heptahydrate exists.

The acid salt NaHSO, is formed by saturating Na CO Aq with SO and cooling; better crystals are obtained by treating Na CO 1. 10aq with SO₂ and evaporating the liquid so produced in SO₂. Easily sol. water, but less sol. than Na₂SO₃. 7aq. (For heat of solution v. De Forcrand, C. R. 98, 738.) Gives off SO₂ on exposure to air; SO2 is also removed from NaHSO,Aq by passage of an indifferent gas (Gernez, C. R. 64, 606). When heated gives off H₂O, SO₂, and S, leaving Na₂SO₄; heated in a closed tube forms Na₂SO₄, H₂SO₄, S, and H₂O (Barbaglia a. Gucci, B. 13, 2325). Reacts with I to give NaHSO, and NaI (Spring a. Bourgeois, Ar. Ph. 229, 707, where references are given to other experiments). Another acid salt Na₂SO₃.SO₂ (= Na₂S₂O₃), sometimes called pyrosulphite, is said to separate, in lustrous prisms, from conc. hot NaOHAq saturated with SO.

(v. Röhrig, l.c.).

withDouble salts ammonium sulphite. Marignac (Ann. M. [5] 12, 30) described a double salt obtained by passing NH, into conc. NaHSO, Aq; the salt is probably identical with that described by Schwicker (B. 22, 1728), to which he gives the formula Na₂SO₂.(NH₄)HSO₃. 4aq. By partially saturating cone. (NH₄)HSO₃ with Na₂CO₂Aq, Trauber (J. C. T. 1888, 44) obtained the salt 2Na₂SO₃.(NH₄)₂S₂O₅. 10aq; Hartog writes the formula of this salt 2Na₂O.(NH₄)₂O.4SO₃, and says it crystallises with 9aq (C. R. 109, 179, 221, 48t). Example 18 of NS Constitution 18 SO₄. 436). For other double salts of Na, SO, v. supra, cobalt sulphites, copper sulphites, gold sulphites, iridium sulphites, lithium sulphites, manganese sulphites, mercury sulphites, palladium sulphites, platinum sulphites, potassium sulphites, rhodium sulphites, and silver sulphites.

Strontium sulphites. The normal salt SrSO, is formed by passing SO, over SrO at 290° (Birnbaum a. Wittich, B. 13, 651); also by decomposing SrCl2Aq by alkali sulphite solution. Small white crystals; slowly oxidises in air to sulphate (Muspratt, P. M. [3] 30, 414); gives SrS and SrSO, on heating (Forster, P. 133,

Thallium sulphites. The normal thallous salt Tl₂SO₃ is prepared by adding Na₂SO₃Aq to Tl₂SO,Aq, and crystallising the pp. from warm water. White crystals; S. 3.34 at 15°; insol. alcohol; S.G. 6.427 at 20° (Seubert a. Elten, Zeit. f. anorg. Chem. 2, 434). The same salt was obtained by Röhrig (J. pr. [2] 37, 217) by passing SO, into solution of a thallous salt, evaporating at 100° or over H2SO4, or adding alcohol.

Tin sulphites. Basic stannous salts, xSnO.2SO₂. 20aq, x being probably 5, 8, and 11, were obtained by Röhrig (J. pr. [2] 37, 217) by treating freshly ppd. SnO. xaq with SO₂Aq.

Titanium sulphites. Berthier (A. Ch. [3] 7,

77) made some observations, but described no

definite salts.

Uranium sulphites. A basic uranous salt UO₂.SO₂. 2aq = U(OH)₂SO₃. aq is obtained, as a greyish-green pp., by adding Na, SO, Aq to UCl, Aq (Rammelsberg, P. 56, 125; confirmed by Röhrig, J.pr. [2] 37,217). The uranyl salt UO2. SO3. xaq was obtained, in yellow prisms, by Muspratt (P. M. [3] 80, 414), by passing SO₂ into water with UO, in suspension, and allowing the solution so formed to evaporate at the ordinary temperature; Muspratt gave x=8, according to Röhrig (l.c.) x=4 (cf. Girard, C. R. 34, 22; and Reinele, P. 125, 238).

Zinc sulphites. The normal salt 2ZnSO₃. 5aq is prepared by evaporating, or by adding alcohol to, a solution of ZnCO, in SO2Aq (Rammelsberg, P. 52, 90); also by mixing cold 25 p.c. ZnSO₄Aq acidified with acetic acid with cold 25 p.c. Na₂SO₃Aq, and letting stand (Denigès, Bl. [3] 7, 569). Muspratt (l.c.), Fordos a. Gélis (J. Ph. [3] 4, 333), and Koene (P. 63, 245, 431) give the formula ZnSO₃. 2aq; but Röhrig (J. pr. [2] 37, 217) confirms Rammelsberg's formula.

Zirconium sulphites. Compositions of salts obtained by ppg. Zr salt solutions by alkali sulphites are doubtful (v. Berthier, A. Ch. [3] 7,

77; Hermann, J. pr. 31, 77).

HYPOSULPHITES. (Hydrosulphites.) Salts of the acid H.S.O. Hyposulphurous acid is known only in aqueous solution. Schützenberger (A. Ch. [4] 70, 351) obtained the Na salt, and an aqueous solution of the acid, in 1869; he gave the formula H.SO, to the acid; but Bernthsen (A. 208, 142; 209, 285; 211, 285) in 1881 showed that the acid has the composition $H_2S_2O_4$. This acid corresponds with the oxide S_2O_4 $(S_2O_4 + H_2O = H_2S_2O_4)$. Schützenberger prepared the Na salt by digesting conc. NaHSO_AQ with zinc shavings in a closed flask for c. half an hour, pouring the liquid from the crystals of the double Na-Zn sulphite which formed (this is a basic Zn-Na sulphite, according to Bernthsen, A. 211, 297) into a flask about three-fourths filled with conc. alcohol, allowing to stand for

some time in the closed flask, and then pouring off into flasks, each flask being quite filled with the liquid and then sealed; after a time crystals of the hyposulphite were formed, to which S. gave the composition NaHSO₂. Bernthsen (l.c.) after reducing NaHSO₂Aq by Zn, pouring off from the double sulphite formed, &c., removed sulphite and sulphate by adding $BaCl_{2}Aq$; he then filtered and titrated the solution by ammoniacal CuSO, Aq, which is decolourised by Na2S2O, Aq to a cuprous salt (and eventually to Cu₂H₂ and Cu₂S). For details of the method of analysis v. Bernthsen (B. 13, 2277; A. 211, 292 note). B. failed to obtain Na₂S₂O₄ free from Na₂SO₄, Na₂SO₃, Na₂S₂O, and Zn salts.

B. expresses the first part of the reduction of NaHSO, by the equation 4NaHSO, Aq + Zn = $Na_2S_2O_4Aq + ZnSO_3 + Na_2SO_3Aq + 2H_2O_4$, and the

complete change by the equation

 $10NaHSO_8Aq + 3Zn =$ $3Na_2S_2O_4Aq + Na_2SO_3Aq + Zn_3Na_2S_3O_{10} + 5H_2O$. Moist sodium hyposulphite is rapidly oxidised in air to Na₂SO₃; when dry it is unchanged in O; it is easily sol. water, less sol. dilute alcohol, and insol. conc. alcohol; but according to B. even approximately pure Na₂S₂O₄ has not been isolated. Na₂S₂O₄Aq is oxidised by iodine so-

lution to Na, SO, Aq.

By heating Na, S, O, Aq with an equivalent quantity of H,SO,Aq or H,C,O,Aq, an aqueous solution of hyposulphurous acid H,S,O,Aq is obtained; this solution is an energetic reducer, decolourising indigo, and reducing salts of Cu, Hg, Ag, &c.; the solution rapidly decomposes in air with separation of S. M. M. P. M.

SULPHO ... Use of this prefix applied to inorganic compounds: for sulpho- compounds and sulpho- salts v. the compounds or salts to the names of which sulpho- is prefixed; thus sulpho-molybdates are described under MOLYBDATES; but the prefix thio- is used in almost every case in preference to sulpho- to denote the replacement of O by S. Sulphocyanides and similar compounds are described under CYANIDES; sulphocyanic anhydride is described as Cyanogen SULPHIDE

SULPHO-ACETIC ACID C,H,SO,

SO₃H.CH₂.CO₂H. [75°].

Formation.—1. By adding SO₃ to cooled HOAc (Melsens, A. Ch. [3] 5, 392; 10, 370; A. 52, 276).—2. By the action of fuming H₂SO₄ on acetamide or acetonitrile (Buckton a. Hofmann, C. J. 9, 247).—3. By boiling chloro-acetic acid with aqueous K₂SO₃ (Strecker, Z. [2] 4, 214; Collmann, A. 148, 109).-4. By heating Ag₂SO₄ with AcCl at 120° (Kämmerer a. Carius, A. 131, 165).-5. By boiling chloro-acetic acid with ammonium sulphite solution (Hemilian, A. 168, 145).—6. By heating Ac.O with H₂SO₄ at 130° (Franchimont, C. R. 92, 1054; cf. Baumstark, A. 140, 83).—7. By heating CISO, H with HOAc at 140° (Baumstark).—8. By oxidation of isethionic acid by CrO, (Carl, B. 14, 63).

Properties. - Deliquescent crystals (containing 1 aq) giving off water of crystallisation at about 160° and decomposed at 230° into CO SO₂, acetic acid, and other products (Franchimont, R. T. C. 7, 25). V. sol. water. Its salts are sol. water, insol. alcohol. PCl, acting on the Nasalt forms CHCl(SO2Cl).COCl (B. Siemens,

B. 6, 659). Vol. IV.

Salts .- K.A" aq: hard crystals .- KHA": six-sided tables.—BaA" aq. S. 267 at 20° .-BaA"Et, SO, (Laube, A. 218, 258).—CaA"aq.—

PbA": radiating needles.—Ag, A" ag: flat prisms.

Mono-ethyl ether EtHA". Formed by
passing HCl into the Ag salt suspended in alcohol. Syrup, v. sol. water. - AgEtA": plates.

Ethyl ether Et, A". Oil. Formed, together with C₂H₁₄Ag₂S₃O₁₆ by the action of Ag₂A" on EtI (F.). May be distilled (Mauzelius, B. 21, 1550

Sulpho-diacetic acid v. DI-METHYL SULPHONE DICARBOXYLIC ACID.

Di-sulpho-acetic aldehyde C2H4S2O, CH(SO₃H)₂.CHO. By adding chloral hydrate to conc. KHSO,Aq at 80° there are formed crystals of C₂H₂K₂S₂O₇KHSO₃, and these when boiled with HClAq yields C₂H₂K₂S₂O₇aq, from which C.H.,BaS,O, 2aq may be prepared (Rathke, A. 161, 154). Boiling K₂CO₃ yields CH₂(SO₃K)₂, potassium formiate, and K₂SO₃.

SULPHO-AMIDO-BENZOIC ACID v. AMIDO-

SULPHO-BENZOIC ACID.

DI-SULPHO-ANILINE v. DI-AMIDO-DI-PHENYL DISULPHIDE. (B.) - SULPHO - ANTHRACENE - (A) - CARB-

OXYLIC ACID $C_{\sigma}H_{\bullet} < \stackrel{C}{C}(CO_{2}H) > C_{\sigma}H_{\bullet}(SO_{3}H)$. [above 360°]. Obtained by dissolving anthracene-(A.)-carboxylic acid in cold conc. H₂SO₄ and allowing to stand for 12 hours on ice (Behla, B.

20, 706). Minute yellowish prisms (from alcohol). V. sol. water and alcohol, insol. benzene and ether. The aqueous solutions of the acid and its salts have a strong blue fluorescence.-BaA": very soluble yellowish minute prisms.

Di-sulpho-anthracene-(A.)-carboxylic acid $C_1H_1(SO_3H)_2.CO_2H$. [above 360°]. Formed by dissolving anthracene-(A.)-carboxylic acid in cold fuming H_2SO_4 (Behla). Yellow prisms (from alcohol). V. sol. water and alcohol, insol. benzene. Its aqueous solution scarcely fluoresces, but the conc. H2SO, solution has a strong yellowish-green fluorescence. - A'", Ba: very sparingly soluble crystalline powder.

SULPHOBENZIDE v. DI-PHENYL SULPHONE. o-SULPHO-BENZOIC ACID C, H. SO. i.e.

C.H.(SO.H).CO.H. S. 50 in the cold. Formation.—1. Together with its imide, by oxidation of toluene o-sulphonic amide with KMnO₄ (Remsen a. Fahlberg, Am. 1, 433).—2. By heating the imide with HClAq at 100° (Fahlberg a. Remsen, B. 12, 472; Fahlberg a. Barge, B. 22, 754; Brackett a. Hayes, Am. 9, 399). By this means the acid ammonium salt is first obtained (F.; Remsen a. Dohme, Am. 11, 332).—3. By boiling o-diazo-benzoic acid with alcoholic SO_2 (Wiesinger, B. 12, 1349).—4. By oxidation of toluene o-sulphonic acid by KMnO.

Properties. - Trimetric crystals (containing 3aq or 4aq); a:b:c='551:1:'812, v. e. sol. water, sol. alcohol, insol. ether. According to Fahlberg (B. 22, 754), it crystallises with 3aq, and is decomposed at 105°. According to Remsen (Am. 11, 332), it crystallises with 4aq, melts at 69° in its water of crystallisation, melts at 130° when anhydrous, and gives a sublimate of long needles (containing 2aq). Resorcin (1.2 pts.) at 195° forms 'sulphofluorescein' C_{1s}H₁₂SO, 2aq, which fluoresces in aqueous and alkaline solutions (Bemsen a. Hayes, Am. 9, 872). By condensation of resorcin (4 mols.) with o-sulpho-benzoic acid there is likewise formed a similar body $C_{s1}H_{20}SO_{s}4aq$ (Fahlberg a. Barge, B. 22, 765). Potash-fusion gives salicylic acid.

Salts.—KHA": transparent plates, sol. water.— K_2A'' 2aq.— $Ba(HA'')_2$ 2½aq: needles, sol. water.— $Ba(HA'')_2$ 2aq.— $BaH_2A''_2$ 4½aq.—BaA'' 2aq.— $Ca(HA'')_2$ 6aq.—CaA'' 5aq.—CuA'' 3½aq.— Ag_2A'' .— $C_0H_4(SO_3NH_4).CO_2H$. Formed by heating the imide with HClAq at

Anhydride $C_0H_4 < \stackrel{CO}{>} 0$. [119°] (F.). [128°] (Remsen a. Dohme, Am. 11, 332). Formed by heating the K salt with PCl, at 180° or the acid with P₂O₅ at 130°. Monoclinic tables, insol. cold water. NH, passed into its solution in benzene forms C_eH₄(CONH₂)SO₂ONH₄ [256°], which crystallises from alcohol in needles.

Chloride C.H. (SO.Cl).COCl. [73°]. Prisms (from ether). When reduced by zinc-dust and H₂SO₄ it yields C₆H₄(SH).CO₂H (Delisle, B. 22,

2205).

Amic acid C6H4(CO.NH2)SO3H. Crystals (containing aq), v. sol. alcohol and water. Got by the action of H₂S on C₂H₄(CONH₂).SO₂Ag aq, which is obtained by adding silver nitrate to C6H4(CONH2).SO3NH4 prepared as above (Fahl-

berg a. Barge, B. 22, 760).

Amic acid C.H.(SO2NH2).CO2H. Sulphamine-benzoic acid. Formed by boiling toluene o-sulphonic amide with KaFeCy, and aqueous potash (Noyes, Am. 8, 178; cf. Fahlberg, Am. 1, 170). Slender needles or prisms, v. sol. water, alcohol, and ether. Melts at 155° when slowly heated, and at 167° when quickly heated. At 180° it is completely converted into imide. Gives rise to the salts MgA'₂6½aq, BaA'₂4½aq, BaA'₂9aq, AgA', and Ag₂C₇H₃NSO₄. By passing HCl into an alcoholic solution of the imide, the ether C_cH₄(SO₂NH₂).CO₂Et [83°] is formed. The methyl and propyl ethers have also been obtained and may be converted by aniline into C_oH₄(SO₂NH₂).CONHPh [189°], by o-toluidine into C_oH₄(SO₂NH₂).CONHC₂H₄ [193°], and by means of p-toluidine into the isomeric p-toluide $C_0H_1(SO_2NH_2)$. CONHC, H, [202°].

Imide C₈H₄ < SO₂ NH. Saccharin. [220°] (Remsen a. Fahlberg, Am. 1, 432). Formed by oxidising CH₃,C₆H₄,SO₄NH₂ with KMnO₄ in neutral solution (Remsen a. Fahlberg, B. 12, 469; Remsen a. Palmer, Am. 8, 223; Fahlberg a. List, B. 20, 1597; 21, 242; Remsen a. Linn, Am. 11, 73). Elongated triangular plates (by sublimation), sl. sol. cold water, v. sol. alcohol and ether. Has a very sweet taste (500 times that of cane sugar). The solubility of commercial saccharin is given by Gravill (Ph. [3] 18, 837) as S. (water) ·2 at 15°; ·6 at 100°; S. (alcohol) 5.4 at 15°; S. (ether) 1.3 at 15°. According to Stift (B. C. 18, 458) and Aducco and Mosso (C. C. 1887, 1148) it retards digestion, and is antiseptic. The imide is converted into C₈H₄(SO₂NH₂).CO₂H on evaporation with KOHAq. Conc. hydrochloric acid at 150° forms C_eH₄(SO₂NH₄).CO₂H. The imide gives rise to salts of the form C.H. CO NAg, of which

BaA'₂1½aq.—BaA'₂4aq (Noyes, Am. 8, 180).—AgA': needles (from hot water). To detec To detect 'saccharin,' a suspected liquid may be acidified, extracted with ether, the ether evaporated, the residue fused with NaOH, and the product tested for salicylic acid and for sulphate (C. Schmitt, Fr. 27, 396; Allen, An. 13, 105; Börnstein, Fr. 27, 165). 'Saccharin' evaporated to dryness on a water-bath leaves a residue which, when warmed with alcoholic potash, exhibits yellow, blue, and red colours (Lindo, C. N. 58, 51, 155).

Methylimide $C_eH_4 < SO_2 > NMe$.

Formed by heating $C_6H_4 < SO_2 > NAg$ with MeI at 100° (Brackett, Am. 9, 406). Long flat needles, sol. ether, alcohol, and hot water.

 $C_6H_4 < SO_2 > NEt.$ Ethylimide

Needles (from water). Converted by boiling alcoholic potash into C₆H₄(SO₂NHEt).ČO₂H [116°], which yields C₆H₄(SO₂NKEt).CO₂K crystallising in nacreous scales, Na₂C₆H₆NSO₄, CuA'₂ 2aq, and AgA' (Fahlberg a. List, B. 20, 1596).

Propylimide C_oH₄ CO NPr. [60°-70°].

m-Sulpho-benzoic acid [1:3]C₆H₄(SO₃H).CO₂H.

Formation.—1. By mixing benzoic acid (2 pts.) with SO₃ (1 pt.) (Mitscherlich, P. 31, 287; 32, 227; Barth, A. 148, 33).—2. By heating BzCl with Ag₂SO₄ at 150° (Kämmerer a. Carius, A. 131, 153; B. 4, 219).—3. By heating BzCl with H₂SO₄ (Oppenheim, B. 3, 735).—4. By the action of an alcoholic solution of SO₂ are midiscaled. action of an alcoholic solution of SO2 on m-diazobenzoic acid (Wiesinger a. Vollbrecht, B. 10, 1715).-5. By oxidation of m-sulpho-benzoic aldehyde with air at 95° (Kafka, B. 24, 796).

Properties.—Deliquescent crystalline mass. Yields m-oxy-benzoic acid when fused with

potash.

Salts.-NaHA" 2aq: triclinic crystals.-Na₂A"Me₂SO₄ (Stengel, A. 218, 260). Crystals.— Na A"Et₂SO₄: v. sol. water.—KHA" 2 aq.— KHA" 5aq (Otto, A. 122, 155).—BaA" 3aq: v. sol. water (Fehling, A. 27, 322).—BaH,A" 23aq. White monoclinic prisms. S. 5 at 20° (M.).—BaA"Me,SO..—BaA"Mo,SO. BaA"Me,SO,.—BaA"Me,SO, 3½aq. S. 34 at 21°. —BaA"Et,SO, 3½aq. S. 31 at 21°. Formed from the acid, NaHSO4, and alcohol (Stengel). Long needles.—BaA"Pr₂SO, 7aq. S. 11 at 19°.— PbA" 2aq: stellate groups of white needles .-PbA"Et,SO, $2\frac{1}{2}$ aq. — CuA"Me,SO, $2\frac{1}{2}$ aq. Crystals. — CuA"Et,SO, $2\frac{1}{2}$ aq. — AgHA": nodules (Limpricht a. Uslar, A. 106, 50).—Ag,A" aq: small yellowish prisms.

Mono-ethyl ether EtHA". Formed by the action of alcoholic NH₃ on Et₂A" or on the chloride (Limpricht a. Uslar, A. 102, 252). The free acid is unstable.-EtNH,A": four-sided triclinic tables (Keferstein, A. 106, 385).-NaEtA" xaq. — Ba(EtA"), xaq: efflorescent

tables.

Ethyl ether Et,A". Formed from the chloride and alcohol. Syrup, miscible with water. Saponified by heating with water.

Semi-chloride C.H.SO.Cl. Formed by

heating the acid (1 mol.) with PCl_s (1 mol.) (L. a. S.). Formed also by leaving the chloride the following have been described.—KA'aq: (L. a. Š.). Formed also by leaving the chloride prisms, v. sol. water.—NaA'2aq: tables.— C.H. (SO.Cl)(COCl) in contact with water for several weeks. Crystalline powder, sol. ether. Converted by hot water into the acid.

Chloride C.H. (SO,Cl).COCl. Formed by heating the acid (1 mol.) with PCl, (2 mols.) at 100°. Thick oil, slowly decomposed by water. Yields CaH, Cl. COCl on distillation and on heating with PCl, at 150° (Kämmerer a. Carius, A. 131, 159).

Benzoyl derivative C₁₁H₁₀SO₈. The first product of the action of BzCl (2 mols.) on Ag₂SO₄ (1 mol.) (K. a. C.). Amorphous, sol. ether. Resolved by water into benzoic and sul-

phobenzoic acids.

Amic acid C₆H₄(SO₂NH₂).CO₂H. m-Sulnhamine-benzoic acid. [235° uncor.] (R. a. P.); [247°] (N. a. W.). Formed by the action of conc. KOHAq at 100° on the amide (Limpricht a. Uslar, A. 106, 27). Prepared by oxidation of [1:3] C,H,Me.SO,NH, with alkaline KMnO,, with chromic acid mixture (Remsen a. Palmer, Am. 4, 143), or with alkaline K, FeCy, (Noyes a. Walker, Am. 8, 188). Scales (from water), sl. sol. cold water, v. sol. alcohol. Converted by PCl, at 150°-200° into the oily chloride C₆H₄(SO₂NH₂)COCl.—BaA'₂ 4aq.—AgA' aq.— Ag₂C,H₅NSO,: amorphous pp., sol. hot water.— EtA'. Monoclinic crystals, sl. sol. hot water.

Amide C₆H₄(SO₂NH₂)(CONH₂). [170°]. Formed from the chloride and conc. NH,Aq. Small anhydrous crystals (from alcohol) or needles (containing aq). V. sol. hot water and alcohol. PCl, at 100° forms C.H. (SO,NH.) CCl:NH, which is decomposed by distillation into mchloro-benzonitrile, SO2, and NH3; and by treatment with NH, Aq into C, H, (SO, NH,).CN [152°]

(Wallach a. Huth, B. 9, 428).

Anilide C. H. (SO2NHPh). CONHPh. Formed from the chloride and aniline. Small crystals (from alcohol).

Isomeride of the amic acid C.H.C(NH.):N.SO₃H. Benzamidine v-sulphonic acid. [247°]. Mol. w. 200 (obs.). Formed by passing SO₃ into cold benzonitrile (Engelhardt, J. pr. 75, 363; Eitner, B. 25, 471). Prisms and rhombohedra. Converted by HClAq into benzamidine sulphate. - CaA'2 - BaA'2 4aq.

Amorphous isomeride of the amic acid. A product of the action of PCl, on the amic acid (L. a. U.). Insol. water, alcohol, and ether. Forms amorphous salts. Water at 180° reconverts it into the ordinary amic acid.-

BaA'24aq: brittle mass

p-Sulpho-benzoic acid [1:4] C₆H₄(SO₃H).CO₂H. [c. 200°]. Formed, in small quantity, together with the m- isomeride, by the action of SO₃ on benzoic acid (Remsen, A. 178, 275; Z. [2] 7, 81, 199). Formed also by oxidation of toluene p-sulphonic acid with KMnO₄ (Hart, Am. 1, 342), and by the action of an alcoholic solution of SO2 on diazotised p-amidobenzoic acid (Wiesinger, B. 10, 1715). Needles, not deliquescent, decomposed by heat. fusion converts it into p-oxy-benzoic acid. The K salt heated with potassium formate yields terephthalic acid.

Salts.-NaHA" 2 aq. Stellate groups of prisms, m. sol. cold water.—BaA" 2aq: groups of needles, m. sol. cold water. - BaH2A", 3aq:

flat needles, v. sl. sol. cold water.

Amic acid C₆H₄(SO₂NH₂).CO₂H. [c. 280°]. Formed by exidation of C₆H₄Me.SO₂NH₂ and of

p-sulpho-cinnamic amide (Remsen, A. 178, 299; Fahlberg, Am. 1, 170; Palmer, Am. 4, 164; Noyes, Am. 7, 145; 8, 182). Flat prisms (from water), sl. sol. hot water, v. sol. alcohol. Decomposed on fusion.—NH,A': needles or long lamins, v. sol. water.—BaA'₂ 5aq.—AgA': v. sl. sol. hot water.—C₈H₄(SO₂NHAg).CO₂Ag: v. sol. hot water.—EtA'. [111°].

Di-sulpho-benzoic acid C,H,S,O, i.e.

 $C_6H_3(SO_8H)_2CO_2H$ [4:2:1]. above 285°1. Formed by oxidation of toluene disulphonic acid with chromic acid mixture (Blomstrand a. Hakansson, B. 5, 1088; Brunner, Sitz. W. [2] 78, 665) and by digesting $C_0H_3(SO_2NH_2) < SO_2 > NH$ with HClAq at 160° (Fahlberg, Am. 2, 190). Large crystals (from HClAq), sol. cold water, insol. alcohol and ether. Yields (4,2,1)-di-oxybenzoic acid on fusion with potash.

Salts.—K₂A" 2aq: prisms, v. e. sol. water.

—K₂HA"aq: large crystals, sl. sol. water.—
Ba₃A" 7aq.—Cu₃A" 7aq: minute crystals.

Amic acid C₂H₃(SO₂H)(SO₂NH₂).CO₂H

[1650] Ecomod by helling the mide

[2:4:1]. [165°]. Formed by boiling the imide of the amic acid with dilute HClAq for five hours. Large crystals, v. e. sol. water, sl. sol. HClAq, insol. ether.—HKA": crystals (from dilute HClAq), v. e. sol. water.

Di-amic acid C₀H₃(SO₂NH₂)₂CO₂H. [183°]. Formed by heating CoH, Me(SO2NH2)2 with aqueous potassium permanganate and by evaporating $C_6H_3(SO_2NH_2) < \frac{SO_2}{CO} > NH$ with KOHAq (Fahlberg, Am. 2, 186; B. 21, 246). Minute sating needles, v. e. sol. water and alcohol, sl. sol. ether. Tastes acid. Decomposes completely at 250°-260°. Its salts are crystalline. Those of alkalis and alkaline earths are v. sol. water.—BaA'25aq: monoclinic prisms.—CuA', 2aq: pale-blue silky needles.-AgA': white needles.-EtA'. [200°]. Silky needles (from water).

Imide of the amic acid

 $C_aH_3(SO_2NH_2) < SO_2 > NH.$ [285°]. Formed by warming C,H,(SO,NH,),CO,Et with KOHAq (Fahlberg a. List, B. 20, 1603; 21, 242). Tables, sl. sol. cold water, v. e. sol. alcohol and ether .-K2C,H4N2S2O,: deliquescent crystalline mass.- $(C_bH_1(SO_2NH_2):SCO_2:N)_2$ ls $3\frac{1}{2}$ aq: crystals. — $Cu(C_tH_3N_2S_tO_3)_2$ 4aq: minute blue needles. — $AgC_tH_3N_2S_tO_3$.

s-Di-sulpho-benzoic acid C₆H₃(SO₃H)₂.CO₂H. Formed by heating benzoic acid with H.SO., SO₃, and P₂O₅ (Barth a. Senhofer, A. 159, 217). Hygroscopic prisms (containing xaq). Turns brown at 140°. Yields s-di-oxy-benzoic acid on fusion with potash. The K salt forms isophthalic acid when fused with potassium formate.—K,A''' 1 aq: needles.—BaHA" 2aq: minute needles.—Ba,A", 7aq: white prisms.—Cu,A", 8½aq.—Ag,A" 2aq: crystalline pp.

References. - BROMO-, NITRO-, OXY-AMIDO, and OXY-, SULPHOBENZOIC ACIDS.

m-SULPHO-BENZOIC ALDEHYDE

C_eH₄(SO₂H).CHO. Sulphonic acid of benzoic alldehyde. Formed from benzoic aldehyde and fuming H₂SO₄ below 50° (Wallach a. Wüsten, B. 16, 150; cf. Engelhardt, J. 1864, 350). The Na salt reacts with (a)-naphthylamine forming C_dH₄(SO₂Na).CH:NC₁₀H₄, and forms with sodium naphthionate C.H. (SO, Na). CH: NC, H. SO, Na. Yields m-sulpho-benzoic acid on oxidation .-MgA'₂ (dried at 170°).-BaA'₂ (dried at 170°). Nodules.

The salt CoH, (SO, Na).CH:NOH Oxim.crystallises in plates, v. e. sol. water, m. sol. alcohol (Kafka, B. 24, 791).

Phenyl-hydraside.

The salt C₆H₄(SO₃Na).CH:N.NHPh crystallises in needles, sl. sol. cold water, v. sol. alcohol.

Di-phenyl-hydrazide

The salt C.H.(SO,Na).CH:N.NPh2 crystallises from water in plates.

SULPHOBÊNZOLIC ACID v. BENZENE SUL-

PHONIC ADID.

SULPHO - BENZYL - SULPHAMIC C.H. CH(SO.H).NHSO.H. The salt Na2A" 3aq is formed from CoH, CH: NOH and aqueous Na₂SO₃ (Pechmann, B. 20, 2541). It crystallises in needles and is decomposed by NaOHAq into

benzoic aldehyde, Na. SO., and H(NH.) SO..
(a) SULPHO-BUTYRIC ACID C4H. SO. CH2.CH(SO3H).CO2H. Formed from butyric acid and ClSO,H; and also by the action of (NH₄)₂SO₂ on α-bromo-butyric acid (Hemilian, B. 6, 196, 562; A. 176, 1; Franchimont, R. T. C. 7, 27). Formed also from crotonic acid and KHSO, at 130° (Beilstein a. Wiegand, B. 18, 483). Syrup. The Ba salt is converted by into CH3.CH2.CHCl.COCl. - CaA" 2aq: needles (from ether-alcohol), insol. alcohol.—BaA"2aq. S. 7·1 at 16°.—PbA"2aq.—CuA"4aq. -ZnA" 5aq. -Ag.A": prisms, insol. alcohol. a-Sulpho-isobutyric acid C4H8SO, i.e.

CMe₂(SO₃H).CO₂H. Formed by the action of Cl.SO, OH on isobutyric acid and also by oxidation of C₅H₈N₂SO, which is prepared by heating thio-urea with a-bromo-isobutyric acid (Andreasch, M. 8, 412).—Na2H" aq.—BaA" 4aq.

Ammonium sulphite appears to convert a-bromo-isobutyric acid into an isomeride which yields BaC, H, SO, 2aq, crystallising in needles, v.

sol. water.

8-Sulpho-butyric acid CH₂.CH(SO₂H).CH₂.CO₂H. Formed from β-chloro-butyric ether and (NH₄)₂SO₃ (H.). also by oxidising barium oxy-butane disulphonate with chlorine-water (Haubner, M. 12, 547). Amorphous.—BaA" aq (H.).—BaC, H.SO,. White amorphous mass, v. sol. water, almost insol. alcohol. Yields acetic acid by potashfusion.

Sulphodibutyric acid v. DI-PROPYL SULPHONE DICARBOXYLIC ACID.

SULPHOBUTYRIC ALDEHYDE C.H.SO. i.s. CH₂.CH(SO₂H).CH₂.CHO. Got by saturating a solution of crotonic aldehyde with SO, and distilling in vacuo, afterwards distilling with steam. On neutralisation of the residue with baryta the salt BaA'₂ is got (Haubner, M. 12, 546). The barium salt is converted by hydroxylamine into (CH(NOH).CH2.CHMe.SOs)2Ba, which is reduced by sodium - amalgam to (CH2(NH2).CH2.CHMe.SO3)2Ba, which on distillation with lime yields butylamine and a little CH₂.CH:CH.CH₂NH₂. Sodium-amalgam reduces the sulpho-butyric aldehyde to oxy-butane sulphonic acid CH, CH(SO, H).CH, CH, OH, whence NaA', which when distilled with lime gives a mixture of butyl and crotyl alcohols.

SULPHOCAMPHORIC ACID v. CAMPHORIC

SULPHOCAMPHYLIC ACID v. CAMPHORIC

ACID.

SULPHO-CHLORO BENZOIC CHLORO-SULPHO-BENZOIC ACID.

o-SULPHO-CINNAMIC ACID CoH,SO, i.e. C₆H₄(SO₅H).CH:CH.CO₂H. Formed, together with the p-isomeride, by the action of SO, or fuming H2SO4 on cinnamic acid (Marchand, J. pr. 16, 60; Herzog, J. pr. 29, 51; Rudneff, A. 173, 8). Prisms (containing 3aq), v. e. sol. water and alcohol. Yields m-oxy-benzoic acid on fusion with potash.—K,A": amorphous, v. sol. water.—KHA": groups of needles.—CaA" 12aq. -BaH₂A"₂. S. ·45 at 20°. -BaA" 1\frac{1}{2}aq. S. 4.2 at 16°.--Ag, A".

m-Sulpho-cinnamic acid

[1:3] C₆H₄(SO₃H).CH:CH.CO₂H. Formed from m-sulpho-benzoic aldehyde, NaOAc, and Ac₂O (Kafka, B. 24, 791).—BaA": nodules, v. sol. water, sl. sol. alcohol.

p-Sulpho-cinnamic acid

[1:4] C₆H₄(SO₃H).CH:CH.CO₂H. Formed Monoclinic prisms (containing 5aq). above. Yields p-oxy-benzoic acid by potash-fusion.— K₂A" laq. Nodules (from alcohol). Reduces KMnO₄ in the cold (Liebermann, B. 22, 782).— CaA" aq.—BaA" Baq. S. (of BaA") 12 at 20°.
—BaA" aq. S. 4 at 18°.—CuA" 6aq: greenishprisms, v. sol. water.

AmideC,H,(SO,NH,).CH:CH.CONH, [218°]. Needles, v. sol. hot water (Palmer, Am.

4, 163).

Amic acid C₆H₄(SO₂NH₂).CH:GH.CO₂H. S. 058 at 21°. Formed by heating the amide with NaOHAq. Needles (from water), sol. alcohol, sl. sol. ether. Decomposes at 250°. Yields CaA', aq and BaA', 2aq, both crystallising in needles.

SULPHO-CUMINIC ACID C10H12SO3 i.e. [160°]. Formed $C_{n}H_{s}Pr(SO_{s}H).CO_{s}H[4:3:1].$ by sulphonating cuminic acid (Widman, B. 22, 2276). Crystalline.—BaA" aq: sl. sol. water.— BaH₂A''₂ 4¹/₂aq. Prisms, m. sol. water. Chloride C₆H₃Pr(SO₂Cl).COCl. [56°].

Amide C,H,Pr(SO,NH,).CONH,

Tables, sol. hot water.

Amic acid C₀H₅Pr(SO₂NH₂).CO₂H. [246° cor.]. Formed by heating the amide with KOHAq at 100°. Formed also by oxidising (B)-cymene sulphonic amide [148°] with chromic acid mixture, an isomeric change taking place (Remsen a. Day, Am. 5, 158). Needles, sol. hot water. Yields BaA'2 3aq, v. e. sol. water.

Sulpho-n-cuminic acid C_zH_zPr(SO_zH).CO_zH [4:3:1]. Formed by sulphonation of n-cuminic acid (W.). V. sol. alcohol, ether, and water.—BaA" aq: needles, sl. sol. Aq. Chloride C.H.Pr(SO,Cl).COCl. [43°].

Amide C.H.Pr(SO2NH2).CONH2.

Tables (from water).

Amic acid C.H.Pr(SO,NH,).CO,H. [216° cor.]. Formed from the amide (W.), and also by oxidation of p-di-n-propyl-benzene sulphonic amide [103°] (Remsen, Am. 5, 158). Long needles (from water).—CaA', 6aq.—CuHA', 2aq. -AgA': flocculent pp.

Sulpho-cuminic acid. Amic acid C.HMe.(SO,NH2).CO,H[5:8:2:x:1]. Formed by oxidation of isodurene sulphonic amide by alkaline KMnO, (Jacobsen, B. 15, 1855). The K salt is amorphous.

Bulpho-cuminic acid. Amic acid C.HMe.(SO2NH2).CO2H [6:4:2:x:1]. Formed by oxidation of the corresponding isodurene sulphonic amide (J.). The K salt is crystalline. Reference.—OXY-BULPHO-CUMINIC ACID.

SULPHO-CYANIC ACID v. vol. ii. p. 303.

SULPHOCYANO-ACETIC ACID CaHanso, i.e. Cy.S.CH2.CO2H. Formed from potassium sulphocyanide and sodium chloro-acetate (Claesson, B. 10, 1347). Thick oil, changing to a solid polymeride when heated.—NaA' aq: prisms. Converted by AgNO, into HS.CH₂.CO₂Na. CuSO, forms, after a time, a black pp. of cuprous thioglycollate. — KA'aq. — BaA'₂ 4aq: tables. — BaA'₂aq: prisms.—CaA'₂ 2aq.—MnA'₂ 2aq.

Ethyl ether EtA'. (225°). S.G. 1·174.

Formed by boiling chloro-acetic ether with potassium sulphocyanide (Heintz, A. 136, 223). Liquid. Yields thioglycollic acid and thioglycollic ether on distillation with phosphoric acid. Polymerised by repeated distillation, many other products being formed at the same time. Boiling dilute HClAq forms NH2.CO.S.CH2.CO2H [143°], which is v. sol. alcohol and ether, yields MeA' [75°-80°] and CaA', 2aq, and is resolved by heat into cyanic and thioglycollic acids (Nencki, J. pr. [2] 16, 11). Etl at 120° forms ethyl sulphocyanide and iodo-acetic acid.

Isoamyl ether C₅H₁₁A'. (255°). Amide CH₂(S.CN).CONH₂. Needles.

Polymeride (C₃H₃SNO₂)₃. [200°]. Formed from potassium sulphocyanurate and potassium chloro-acetate (Claesson, J. pr. [2] 33, 121; B. 14, 732). Needles, sol. alcohol and ether. Decomposed by HClAq at 100° into thioglycollic acid and cyanuric acid.—Ba₂A'''₂ 6aq: small prisms, nearly msol. water.—BaHA''' 2aq: large prisms.—Et₂A'''. [81°]. Formed by distilling sulphocyano-acetic ether. Needles, nearly insol. hot water.

Persulphocyano-diacetic acid

 $C_2N_2S(S.CH_2.CO_2H)_2$. [177°]. Formed from sodium chloro-acetate, iso-persulphocyanic acid, and KOHAq (Klason, J. pr. [2] 38, 381). Plates. -BaA'' 3aq. — CaA'' 3 $\frac{1}{2}$ aq. — ZnA'' aq. — CdA''. — CuA" 6aq.—Et2A". Õil.

Amide. [125°]. Prisms.

SULPHOCYANO-ACETIC ALDEHYDE

 $CH_2(SCy).COH.$ [below -20°]. S.G. 12 1.47. Formed by heating iodo-acetic aldehyde with AgSCy and ether (Chautard, A. Ch. [6] 16, 193; C. R. 106, 1169). Oil, miscible with alcohol Decomposed by distillation, by and ether. treatment with Cl, and by HClAq. HNO, forms acetic acid, sulphocyano-acetic acid, and thio-Resinified by carbimido-acetic acid [128°].

SULPHOCYANO-ACETONE C.H.NSO i.e. CH₃.CO.CH₂SCy. S.G. ^Q 1.2; ^{2Q} 1.18. S. 7 at 15°. Formed from chloro-acetone and barium sulphocyanide (Tschnerniac, C. R. 96, 587; B. 16, 349; 25, 2607, 2623). Liquid, v. e. sol. ether, v. sl. sol. ligroin. Decomposed by heat. Dissolves in aqueous KHSOs. Boiling dilute HClAq converts it into oxy-methyl-thiazole. Ammonium sulphocyanide forms 'propimine sulphocyanide' CH, C(NH). CH, SCy [42] (231°), which yields B', H, PtCl, B'HNO, [183°], B'H, SO, 2aq, B'HSCy [115°], B'MeI [160°], and CH, C(NAc). CH, SCy [134°].

Oxim CH₂.C(NOH).CH₂.SCy. [135°]. Yellowish prisms (Arapides, A. 249, 18). Produces great irritation on the skin.

SULPHOCYANO-ACETOPHENONE v. PHEN-

ACYL SULPHOCYANIDE.

SULPHOCYANO-BARBITURIC ACID

 $C_bH_sN_sSO_s$ i.e. CO < NH.CO > CH.SCy. Formed from di-bromo-barbituric acid and alcoholic KSCy in the cold (Trzcinski, B. 16, 1058). The free acid splits up into thiodialuric acid and other products. Hot KOHAq also forms thiodialuric acid.—NH, A'.—KA': tables (from water). -AgA': crystalline pr

y-SULPHOCYANO-BUTYRICACID. Nitrile. CH₂(SCy).CH₂.CH₂.CN. (c. 195° at 40 mm.). Formed by boiling potassium sulphocyanide (1 pt.) with γ -chloro-butyronitrile (1 pt.) and alcohol (Gabriel, B. 23, 2490). Liquid. Partially decomposed by distillation under atmospheric pressure. Converted by conc. H.SO.

into S2(CH2.CH2.CH2.CO.NH2)

a-SULPHOCYANO - ISOBUTYRIC ALDE -HYDE CMe₂(SCy).CHO. S.G. 13 1.63. Formed from a-iodo-isobutyric aldehyde and AgSCy (Chautard, A. Ch. [6] 16, 198). Liquid with nauseous odour, decomposed by heat, by acids, and by alkalis.

SULPHOCYANO - ETHANE SULPHONIC ACID C₃H₅NS₂O₃ i.e. CH₂(SCy).CH₂.SO₃H. Formed from CH2(SCy).CH2Cl and aqueous Na₂SO₃ in sunlight (James, J. pr. [2] 26, 881). SULPHOCYANO-FORMIC ETHER

CyS.CO₂Et. The compound (CyS.CO₂Et)EtOH is got by mixing chloro-formic ether with ammonium sulphocyanide and alcohol (Delitsch, J. pr. [2] 10, 118; cf. Henry, J. pr. [2] 9, 464). It crystallises in prisms [44°], insol. water, v. e. sol. alcohol. Alcoholic potash forms C₆H₁₀KSNO₈ crystallising from alcohol in plates. KOHAq forms KSCy, alcohol, and K2CO,

SULPHOCYANO-HEPTOIC ALDEHYDE. Formed from iodo-heptoic aldehyde and AgSCy in ether (Chautard, A. Ch. [6] 16, 198). Yellowish liquid with feetid odour. Decomposed by

heat, by acids, and by alkalis.

β - SULPHOCYANO - PROPIONIC HYDE CH2(SCy).CH2.CHO. Formed by treating \$\beta\$-iodo-propionic aldehyde dissolved in ether with AgSCy (Chautard, A. Ch. [6] 16, 197). Liquid with feetid odour. Decomposed by heat. and resinified by alkalis and by mineral acids.

ω-SULPHOCYANO-TOLUIC ACID. Nitrile. CH₂(SCy).C₆H₄.CN. [86°]. Formed from o-cyanobenzyl chloride and KSCy in alcohol (Gabriel a. Day, B. 23, 2479). Needles. Converted by conc. HClAq at 180° into thiophthalide C.H.SO [57°]. Conc. E₂SO₄ at 30°-70° forms cyano-benzylmercaptan C.H.NS [62°].

SULPHOCYANO-ISOVALERIC ALDEHYDE. Formed from iodo-isovaleric aldehyde and AgSCy in ether (Chautard, A. Ch. [6] 16, 198). Liquid, with feetid odour, easily decomposed by heat, by HClAq, and by alkalis.

SULPHO-DURIDE v. DI-DURYL SULPHONE.

SULPHO-ETHYL-BENZOIC ACID. Amia acid C,H11NSO, i.e. C,HEt(SO,NH2).CO,H. [262° cor.]. Formed by oxidising the sulphonic amide of p-di-ethyl-benzene with CrO. (Remsen s. Noyes, Am. 4, 202). Needles.—BaA's Saq.

SULPHO-FUMARIC ACID C,H,SO, i.e. CO2H.CH:C(SO2H).CO2H. Formed by the action of Br or of HNO, on the salts of (δ)-bromo-(8)-sulpho-pyromucic acid (Hill a. Palmer, Am. 10, 409).—Ba₂A'''₂ xaq: v. sl. sol. water.— Ag₃A''' xaq.

SULPHO-HIPPURIC ACID C, H, NSO, i.s. C₈H₄(SO₃H).CO.NH.CH₂.CO₂H. Formed from hippuric acid and SO₃ (Schwanert, A. 112, 59). Amorphous, deliquescent mass. Converted by nitrous acid into sulpho-benzoic acid.—BaA"aq:

needles.-Pb2OA"

SULPHO-ISATIC ACID v. ISATIC ACID.

SULPHO-TRIMELLITIC ACID C.H.SO. i.e. $C_6H_2(SO_2H)(CO_2H)_3$ [5:4:2:1]. Formed, together with the amic acid $C_aH_1(SO_2NH_2)(CO_2H)_3$, by oxidation of $C_aH_1Me(SO_2NH_2)(CO_2H)_2$ [2:5:4:1] by KMnO₄ (Jacobsen a. H. Meyer, B. 16, 192).— KH, A1 Baq: prisms, m. sol. cold water. Potashfusion yields oxy-trimellitic acid.

SULPHO-TRIMESIC ACID. Amic acid $\mathbf{C_9H_7NSO_8}$ i.e. $\mathbf{C_9H_2(SO_2NH_2)(CO_2H)_3}$. [x:5:3:1]. Formed by oxidation of the amic acid of o- or p-sulpho-mesitylenic acid (Jacobsen, A. 206, 203).—KH₂A''' 2aq: crystalline mass, m. sol. cold water. Conc. HClAq at 210° forms NH₂, H.SO., and trimesic acid. Potash-fusion gives

oxytrimesic acid.

α-SULPHO-MESITYLENIC ACID C.H. SO. i.e. $C_aH_2Me_2(SO_3H).CO_2H$. Formed, together with a (8)-isomeride which yields CaA" 4aq, by the action of SO, on mesitylenic acid (Remsen a. Brown, Am. 3, 218). Yields o-oxy-mesitylenic acid on fusion with potash. By successive treatment with PCl, and NH, it is converted into an amide [288°].—CaA" 4aq. Less sol. water than the salt of the (β) -isomeride.

Sulpho-mesitylenic acid. AmicC₀H₁₁NSO₄ i.e. C₀H₂Me₂(SO₂NH₂).CO₂H [5:3:6:1]. [262° cor.]. Formed, together with the (5,3,4,1)-isomeride, by oxidising mesitylene sulphonic amide with chromic acid mixture or alkaline KMnO₄ (Hall a. Remsen, Am. 2, 131; B. 10, 1040; Jacobsen, B. 12, 604; A. 206, 167). Short prisms (from water), sl. sol. hot water, sol. alcohol and ether. Decomposed by heating with conc. HClAq at 200° into H₂SO₄ and mesitylenic acid. Soda-fusion forms mesitylenic acid and m-xylene sulphonic amide [137°].-CaA'₂6aq: long flat plates.—CaA'₂5aq.— BaA'₂8aq. S. 3·3 at 0°; 14 at 21°.—CuA'₂4aq: blue needles.—CuA', 3aq.—AgA': pp.
Sulpho-mesitylenic acid. Amic acid

 $C_0H_2Me_2(SO_2NH_2).CO_2H$ [5:3:4:1]. [276° cor.]. Formed as above (J.). Long needles, v. sol. alcohol and ether, more sol. hot water than its isomeride. Conc. HClAq at 200° forms mesityl-Soda-fusion gives oxy-mesitylenic 2aq. S. 2.05 at 0°.—CaA'₂ 2aq: enic acid. acid. -BaA', 2aq.

prisms.—CuA', aq: blue monoclinic prisms. SULPHO - DI - METHYL - BENZOIC ACID.

Amic acid C,H, NSO, i.e. C₆H₂Me₂(SO₂NH₂).CO₂H [4:2:5:1]. [268°]. Formed by oxidation of the sulphonic amide of ψ -cumene (Jacobsen a. Meyer, B. 16, 190). Long needles (from water). Yields di-methyl-benzoic acid on

heating with conc. HClAq at 210°.—KA'aq.
Sulpho-di-methyl-benzoic acid. [180°-190°].

Amic acid C_uH₂Me₂(SO₂NH₂).CO₂H [1:2:5:3]. (a) - Sulphamine - hemimellitic acid. [238°]. Formed, together with the more soluble (β)-iso-

meride, by oxidising the sulphonic amide of hemimellithene with alkaline KMnO, (Jacobsen, B. 19, 2519). Needles, sl. sol. cold water. HClAq at 150° forms sulpho-di-methyl-benzoic acid and, finally, hemimellitic acid [144° Potash-fusion yields an oxy-hemimellitic acid. -BaA', 5aq: small tables, m. sol. water.

Sulpho-di-methyl-benzoic acid. Amic acid $C_6H_2Me_2(SO_2NH_2).CO_2H[1:3:5:2].[174^\circ].$ Formed as above. Stellate groups of minute needles. Converted by heating with HClAq into a very soluble sulphonic acid, and finally into m-xylene. Potash-fusion gives an easily soluble oxyhemimellitic acid.—BaA', 4aq: needles.

Isomeride v. Sulpho-mesitylenic acid.

SULPHO-METHYL-TEREPHTHALIC ACID. A mic acid $C_aH_2Me(SO_2NH_2)(CO_2H)_2$ [2:5:4:1]. [295°-300°]. Formed by oxidation of ψ -cumene sulphonic amide by alkaline KMnO, (Jacobsen a. H. Meyer, B. 16, 190). Small needles (from water).—BaA" 2 aq: stellate groups of prisms.

SULPHONAMIDE v. SULPHAMIDE, p. 567; and IMIDO-SULPHONAMIDE v. SULPHIMIDO-AMIDE, p. 587.

SULPHONAMIDES. Amides of sulphonic acids. They contain the group SO2.NH2 attached by S to carbon.

SULPHONAMIDO- compounds v. Amic acids derived from Sulpho- compounds.

SULPHO-NAPHTHALENE-AZO- compounds v. Azo- compounds.

('α')-SULPHO-(α)-NAPHTHOIC ACID

 $C_{11}H_sSO_s$ i.e. $C_{10}H_s(SO_3H).CO_2\Pi$. [235°]. Formed, together with the (8)- and (γ)- isomerides, by warming (a)-naphthoic acid with fuming H₂SO₄ at 70° (Battershall, A. 168, 119; Stumpf, A. 188, 1). Prisms, v. sol. water. Not deliquescent. Yields (a)-oxy-naphthoic acid by potash-fusion. — K₂A''2aq. — CaA'' 3aq. — BaA" 4aq. Monoclinic crystals, sl. sol. water.
—BaH₂A"₂ 2aq: prisms, more sol. than BaA".

(\$\beta\$)-sulpho-(\$\alpha\$)-naphthoic acid. [218°-222°]. Crystalline mass, v. e. sol. water.—BaA" $3\frac{1}{2}$ aq:

needles, m. sol. water. $-\text{BaH}_2\text{A''}_2$ 4aq. (γ)-Sulpho-(α)-naphthoic acid. [182°-185°]. Needles, v. sol. water.—K,A".—BaA" 1½aq: m. sol. water.—BaH,A", aq: almost insol. cold Aq.

Sulpho-(a)-naphthoic acid. Nitrile C₁₀H_aCy.SO₃H. Formed by sulphonating (α)-naphthonitrile (Dutt, B. 16, 1251; Armstrong a. Williamson, C. J. Proc. 3, 43).— BaA'2 6aq: glistening plates.—KA'3aq: prisms.

Chloride of the nitrile C10H8Cy.SO2Cl. Prisms, sol. benzene.

 (α') -Sulpho- (β) -naphthoic acid. [230°]. Formed, together with the (β') - isomeride, by

sulphonating (\$\beta\$)-naphthoic acid (B.; S.). Crystalline.—BaA" aq.—BaA" 6\frac{1}{2}aq.—BaH_2A"_2\frac{6}{2}aq. ('β')-Sulpho-(β)-naphthoic acid.—BaA" 2aq. Less sol. water than the acid salt.

Reference. - OXY-SULPHO-NAPHTHOIC ACID.

SULPHO- NAPHTHYL ETHYL DI-THIO-CARBONATE. The salt SO, K.C, He.S.CS.OEt, formed from potassium xanthate and diazotised naphthionic acid, crystallises from water in colourless plates, converted by boiling dilute alcoholic potash into S₂(C₁₀H_s.SO₂K)₂ (Leuckart, J. pr. [2] 41, 218). The corresponding salt from (8)-naphthylamine (8)-sulphonic acid crystallises in small plates.

SULPHO · (β) - NAPHTHYL - PHOSPHORIC ACID SO₃H.C.₁₀H₈·O.PO(OH)₂. Formed, together with the anhydride $O(PO(OH).C_{10}H_8.SO_3H)_2$, by heating potassium (β) -naphthol sulphonate (1 mol.) with PCl₅ (2 mols.) at 100° (Claus a. Zimmermann, B. 14, 1482). Decomposed by boiling alkalis into phosphate and (β) -naphthol sulphonate. The Ba salt is a white powder, sl. sol. water.

SULPHONES. Compounds of the form R.SO₂.R' where R and R' are attached by means of C to S. They may be formed from the corresponding sulphides and sulphoxides by oxidation with KMnO, (Beckmann, J. pr. [2] 17, 475). Aromatic sulphones can be prepared by the action of AlCl₃ on a mixture of a sulphonic chloride and an aromatic hydrocarbon (Beckurts a. Otto, B. 11, 472, 2066), and by heating a sulphonic acid with a hydrocarbon and P2Os at 200° (Michael a. Adair, B. 10, 583). Sulphones are often formed by the action of sulphuric acid on aromatic hydrocarbons. The sulphones are volatile, and are not reduced by nascent hydro-They are not attacked by PCl₅ or KMnO₄. Chlorine in daylight converts di-phenyl-sulphone into chloro-benzene (1 mol.) and C₈H₅.SO₂Cl, while in sunlight it yields chloro-benzene (2 mols.) and SO₂Cl₂. Disulphones CHR(SO₂R')₂ are not attacked by alcoholic potash at 140°, while disulphones of the form R'.SO₂.CH₂.CH₂.SO₂.R' are converted by alcoholic potash into a sulphinic acid and an oxy-sulphone (Otto a. Rössing, B. 20, 185). The disulphones CMe₂(SO₂ Et)₂ CHEt(SO₂Et)₂, (trional), and CHMe(SO₂Me)₂, (sulphonal), $\text{CEt}_2(\text{SO}_2\text{Me})_2$, $\text{CMeEt}(\text{SO}_2\text{Et})_2$ (trional), and $\text{CEt}_2(\text{SO}_2\text{Et})_2$ (tetronal) are powerful hypnotics (Baumann a. Kast, H. 14, 52).

SULPHONIC ACIDS (organic). Acids containing the group SO2. OH united to C by S. Fatty sulphonic acids may be formed by the action of silver sulphite on alkyl iodides, the resulting ether being saponified. Fatty sulphonic acids may also be got by boiling alkyl iodides with K₂SO₃ or (NH₄)₂SO₃Aq for a long time. Both fatty and aromatic sulphonic acids may be got by oxidation of the corresponding mercaptans, sulphides, and sulphocyanides. The sulphonic acids of fatty acids and alcohols are formed by the action of SO3 or ClSO3H on fatty acids and of SO, on alcohols; SO, H taking the a-position (Hemilian, A. 176, 1). Aromatic compounds readily yield sulphonic acids on treatment with SO₃, with H₂SO₄, or with ClSO₅H. As a rule not more than two SO,H groups enter a benzene nucleus. Aromatic sulphonic acids may also be obtained by the action of SO2 on diazo- compounds. On adding NaCl to the product of sulphonation of aromatic compounds, the Na salt of the sulphonic acid frequently separates in crystalline form (Gattermann, B. 24, 2121). Nitric acid does not attack fatty sulphonic acids, while usually it nitrates aromatic sulphonic acids, although it sometimes displaces SO,H by NO2. PCl3 forms sulphonic chlorides which are reduced by zinc and dilute H2SO4 to mercaptans. Aromatic sulphonic acids are partially converted into the corresponding amido- compounds by fusion with NaNH₂. In this way benzene sulphonic acid yields 15 p.c. of aniline (Jackson a. Wing, Am. 9, 75). Aromatic sulphonic acids are decomposed by superheated steam at 2000210° into the hydrocarbon and H₂SO₄ (Kelbe, B. 19, 92). Hydrolysis may also be effected by passing superheated steam into a solution of the sulphonic acid containing H₂SO₄ or H₃PO₄ (Armstrong; Friedel a. Crafts, C. R. 109, 95). Aromatic sulphonic acids when fused with potash yield K₂SO₂ and phenols. Fusion with KCN or K₄FeCy₅ yields the corresponding nitriles. Fusion with sodium formate displaces SO₃H by CO₂H.

SULPHONIC ACIDS AND DERIVATIVES. Several inorganic acids, and derivatives of these acids, will be described here, most of which are regarded as derived from SO2.OH.OH by replacing one OH by a monovalent radicle-such as Cl, F, or NH2-or as derived from SO. OH.H by replacing H by a monovalent radicle. A few sulphonic acids must be formulated as derived from 2(SO₂.OH.OH), or 2(SO₂.OH.H), by replacing 2OH, or 2H, by a divalent radicle; and one as derived from 3(SO₂.OH.OH), or 3(SO₂.OH.H), by replacing 30H, or 3H, by a trivalent radicle (v. supra). The sulphonic acids, therefore, are all looked on as compounds of the monovalent radicle SO2.OH. The term sulphonic is also frequently applied to organic acids containing this radicle; but in this dictionary such acids are described as sulpho- acids, e.g. sulpho-benzoic acid C₆H₁(SO₂.OH)CO₂H. The compositions of the sulphonic acids are expressed by the formulæ $R^{t}SO_{x}OH$, $R^{tt}(SO_{x}OH)_{x}$, and $R^{ttt}(SO_{x}OH)_{x}$, where $R^{t} = NH_{x}$, Cl, F, NO_{x} , NH. OH, or N(NO. OH); $R^{II} = NH$, or N.OH; and $R^{III} = N$.

AMIDOSULPHONIC ACID AND SALTS $NH_2(SO_2.OH)$ and $NH_2(SO_2.OM)$. These compounds are described in the article Sulphamic acid and sulphamates $(q.v.\ p.\ 567)$.

CHLOROSULPHONIC ACID AND SALTS Cl(SO₂.OH) and Cl(SO₂.OM). The acid has also been called chlorhydrosulphurous acid, monochlorosulphuric acid, sulphuric chlorhydrate or hydrochloride, sulphuric chlorhydrin, and sulphuryl hydroxyl chloride.

Chlorosulphonic acid Cl(SO,OH).

Formation.—1. By the direct union of SO₃ and HCl (Williamson, Pr. 7, 11; Baumstark, A. 140, 75; Williams, C. J. [2] 7, 304; Dewar a. Cranston, C. N. 20, 174; Michaelis, J. Z. 6, 235, 292).—2. By the reaction of PCl., POCl., or PCl., with conc. H₂SO₄ (Müller, B. 6, 227; Geuther, B. 5, 925; Thorpe, C. J. 37, 358).—3. By distilling fuming H₂SO₄ with P₂O₅ in a current of HCl (Müller, L.c.).—4. By the reaction of HCl with crystallised fuming H₂SO₄ (Beckurts a. Otto, B. 11, 2058).—5. By the action of Cl, S₂Cl₂, or S₂O₃Cl₄, with conc. H₂SO₄; or of Cl with moist SO₂ in presence of Pt at a red heat. 6. By warming S₂O₃Cl₂ with water (Billitz a. Heumann, B. 16, 602).

Preparation.—1. Three parts of the most conc. H₂SO₄ (preferably the residue from preparing SO₃) are warmed in a long-necked flask, and two parts PCl₅ are added little by little; when HCl ceases to be given off the contents of the flask are fractionated, the part boiling between c. 152°-156° being collected apart.—2. Very conc. oil of vitriol, 200 parts, is mixed with POCl₃, 226 parts, the mixture is gently heated and then distilled [2(SO₂.OH.OH) + POCl₃ = 2(Cl.SO₂.OH) + HPO₃ + HCl] (Thorpe, Lc.).—

8. A mixture of P2Os and fuming H2SO4 is distilled in a current of HCl, and the product is fractionated; this method removes all traces of P chlorides (Müller, l.c.).—4. Commercial crystalline fuming H.SO, (approximately pure H₂S₂O₇) is placed in a retort connected with a good condenser, dry HCl gas is passed in so long as absorption occurs, the liquid is distilled, and the product fractionated; the yield is very satisfactory (Beckurts a. Otto, B. 11, 2058).

Properties. - A colourless liquid, fuming much in air, and having a powerful, penetrating odour. S.G. $\frac{0^{\circ}}{4^{\circ}} = 1.78474$ (Thorpe, *C. J.* 37, 358); Michaelis (J. Z. 6, 235, 292) gives 1.776 at 18°. Boils at 155.3° at 760 mm. pressure (T., l.c.; v. also M., l.c.; Clausnitzer, B. 11, 2008; Beckurts a. Otto, l.c.; Behrend, B. 8, 1004). Ogier (C. R. 96, 646) gives H.F. [SO3 solid, HCl] =14,400. Baumstark (A. 140, 75) found V.D. 59.3 (calc. = 58.12) (temp. is not given); Williams (C. J. [2] 7, 304) found V.D. at 216° to be 32.8; Heumann a. Köchlin (B. 16, 602) found V.D. 34.7 at 184° and 30.4 at 440°. The gas is therefore dissociated at a temperature not much above its b.-p. When ClSO, OH is repeatedly heated above 158° it is partly resolved into SO, Cl2; distilled, it gives SO₂Cl₂ (Williamson, C. J. 10, 97). By boiling the compound for a long time with an inverted condenser, Clausnitzer (B. 11, 2008) obtained H2SO4, SO2, and Cl; Beckurts a. Otto (l.c.) heated CISO2.OH for a long time in a sealed tube at c. 170°, and found that part remained unchanged, and that Cl and SO, were produced. Heumann a. Köchlin (l.c.) think that the decomposition at c. 440° is represented as follows: $2(CISO_2 \cdot OH) = SO_2 + SO_3 + Cl_2 + H_2O$; they regard the high value obtained by Baumstark for the V.D. as due to presence of S₂O₆Cl₂ (v. also Claesson, J. pr. [2] 19, 235).

Reactions.—1. Action of heat, v. supra, Properties.—2. Water produces HClAq and H,SO,Aq. 8. Hydrogen sulphide reacts at ordinary temperatures, forming HCl, S₂Cl₂, S, and H₂SO₄ (Prinz, A. 223, 871).—4. Carbon disulphide, at 100°, produces HCl, SO₂, S, and COS.—5. Phosphorus pentoxide or pentachloride produce S, O, Cl, (Billitz a. Heumann, B. 16, 482; Konswaloff, C. R. 96, 1146).—6. Heating with silver nitrate forms AgCl and NO. SO. OH (Thorpe, C. J. 41, 297) .- 7. Fusion with potassium sulphate forms K2S2O, and HCl; heating with sodium chloride forms HCl and Cl.SO, ONa (Müller, B. 6, 227). 8. The reactions of ClSO, OH with several elements were examined by Heumann a. Köchlin

(B. 15, 416). Chlorosulphonates. The salts of Cl.SO2.OH, which reacts as a monobasic acid, are formed by the reaction of the acid with metallic chlorides. It is very difficult to obtain these salts pure. When heated they form sulphates, giving off SO2 and Cl; with water they decompose to chlorides and acid sulphates; and with alcohol HCl and salts of Et.H.SO, are formed (Müller, B. 6, 227). No accurate description of any salts has yet been given.

SO₂.Cl CHLOBOSULPHONIC ANHYDRIDE >0; 80,.C1 Pyrosulphuryl chloride, under SULPHUR OXY-

ENCORIDES, p. 618.

FLUOSULPHONIC ACID AND SALTS F.SO₂.OH and F.SO₂.OM. Only the acid has been isolated.

Fluosulphonic acid F.SO₂.OH. Isolated and examined by Thorpe a. Kirman (C. J. 61, 921 [1892]). Prepared by placing pure SO, in a Pt vessel, surrounded by ice and CaCl₂ (for description of apparatus v. Thorpe a. Hambly, C. J. 55, 163), leading in excess of pure HF (made by heating KHF2), and removing excess of HF by passing in a stream of dry CO₂ for many hours, the liquid being kept at 25°-35°. A thin, colourless liquid, with a slightly pungent smell, and fuming in air; boils at 162.6, a little being decomposed with formation of H,SO,, and probably also SO₂F₂. Reacts rapidly with Pb, forming PbSO₄ and PbF₂; slowly attacks glass. Reacts violently, and sometimes explosively, with water, forming H2SO Aq and HFAq.

IMIDOSULPHONIC ACID AND SALTS NH(SO₂·OH)₂, NH(SO₂·OM)₂, and NM(SO₂·OM)₂. By passing dry NH, into SO, H. Rose (P. 32, 81; 47, 41; 49, 183 [1834-40]) obtained two compounds, which he named sulphatammon and parasulphatammon, and to both of which he assigned the composition 2NH₃.SO₃. These compounds were examined by Jacquelain (A. Ch. [3] 8, 293 [1843]), Woronin (J. R. 3, 273 [1859])and others; in 1875 Berglund (Lunds Universitets Arsskrift, 12 and 13; Bl. [2] 25, 455; 29, 422) showed that Rose's parasulphatammon was diammonium imidosulpĥonate NH(SO2.ONH4)2 and that sulphatammon was probably the tri-ammonium salt N.NH₄(SO₂.ONH₄)₂. Berglund's conclusions were confirmed, partly by Raschig (A. 241, 161 [1887]), and Mente (A. 248, 232 [1888], and partly by Divers a. Haga (C. J. 61, 943 [1892]). A full discussion of the constitution of Rose's compounds, with reference to the work of other chemists and an historical summary of researches on the imidosulphonates, will be found in the memoir by D. a. H.

Imidosulphonic acid NH(SO,OH),Aq. This acid is known only in solution, which is obtained by suspending lead imidosulphonate in water, decomposing by a stream of H,S, and rapidly filtering from PbS. The solution gives a pp. with excess of BaOAq, soluble in HNO₅Aq; and a pp. of NH(SO₂.OK)₂ with K.C₂H₂O₂Aq. The acid solution is very unstable, soon becoming changed to NH2(SO2-OH)Aq and H2SO4Aq Jacquelain, l.c.; Fremy, A. Ch. [3] 15, 408; D. a. H., l.c. p. 945).

Imidosulphonates. The normal salts are of the types NH(SO₂.OM)₂ and NM(SO₂.OM)₂; besides these many basic salts have been isolated. The di-alkali imidosulphonates are prepared by mixing solutions of alkali nitrites and sulphites, dissolving the pp. which forms in water, heating this solution for some time, allowing to crystallise, and then boiling the nitrilosulphonate, N(SO₂OM), thus obtained (v. p. 601, NITRILO-SULPHONATES) with acidified water (for instance

 $N(SO_2.OK)_2 + H_2O = NH(SO_2.OK)_2 + KHSO_4);$ also by heating the alkali amidosulphonates (e.g.

2(NH₂,SO₂OK) = NH(SO₂,OK)₂ + NH₃); the diammonium salt NH(SO₂,ONH₄)₃ is also obtained by the reaction of NH₃ with SO₃, ClSO₂OH, SO₂Cl₂, or S₂O₃Cl₂. Most of the other di- salts are obtained by double decomposition from the alkali salts. The tri-salts

are generally prepared by dissolving the di-salts in excess of an aqueous solution of the base and crystallising. A number of mixed salts, such as N(NH₁)(SO₂.O)₂Ba, have been prepared, generally by double decomposition. Basic salts, chiefly of such slightly positive metals as Pb or Hg, have also been isolated. For references v. Divers a. Haga (C. J. 61, 943). The di-alkali salts give off NH₂, N, and SO₂ when heated; water reacts to form amidosulphonates NH₂(SO₂.OM)₂ and sulphates.

The chief imidosulphonates are those of NH₄, Ba, Ca, Hg, Pb, K, Ag, and Na; basic salts of Pb and Hg are known; mixed salts containing NH₄ and Na, NH₄ and Ba, Ba and Hg, Ba and Na, Ca and Na, Hg and K, and Ag and Na have been isolated; some of the imidosulphonates also form double compounds with KNO₃ and NaNO₃. Full accounts of the preparation and reactions of the chief imidosulphonates are given in the memoir

ef D. a H.

NITRILOSULPHONIC ACID AND SALTS N(SO, OH), and N(SO, OM). The acid has not been isolated, and only a few salts are known.

Potassium nitrilosulphonate N(SO₂.OK)₃. Solutions of K₂SO₃ and KNO₂ are mixed in the ratio 4K₂SO₃.KNO₂; after crystallisation occurs the mixture is heated at 100° till the crystals dissolve, a little water is added, and heating at 100° is continued for a little; on cooling the salt crystallises in well-formed crystals with 2H₂O. The water of crystallisation is removed at 100°-110°; at a higher temperature decomposition occurs to K₂SO₄, (NH₄)₂SO₄, SO₃, and SO₂. The salt may be crystallised from dilute KOHAq; it is decomposed by water at 40° to NH(SO₂.OK)₂ and KHSO₄.

Claus prepared this salt in 1871 (B. 4, 186), giving it the formula NH₂(SO₃K), and the name potassium trusulphammonate; Raschig (B. 20, 584 [1887]) showed that the salt was probably N(SO₂OK), and this composition was confirmed both by the earlier work of Berglund (B. 9, 252, 1896 [1876]) and by the more recent work of

Divers a. Haga (C. J. 61, 943).

For description of a salt NO(SO₃K)₂ v. Raschig (A. 241, 225); this is the trisulphooxyazotate of Claus and Raschig. R. regards it

as $(SO_2.OK)_3N < O > N(SO_2.OK)_3$.

Sodium nitrilosulphonate N(SO₂.ONa)₃. Formed by passing SO₂ into a solution of NaNO₂ and Na₂CO₃ (in the ratio 2NaNO₂:3Na₂CO₃, 10aq) until the solution is feebly acid to litmus (v. D. a. H., Lc.).

NITROSULPHONIC ACID NO₂(SO₂.OH), AND DERIVATIVES. The acid and its potassium salt have been isolated. The anhydride SO₂(NO₂).O.(NO₂)SO₂, the chloride NO₂(SO₂.Cl), and an oxy-anhydride S₂O₅O(NO₂)₂have also been

obtained.

Nitrosulphonic acid NO₂(SO₂·OH). (Nitrosylsulphuric acid SO₂·OH.O(NO). Lead chamber crystals. Nitrosyl hydrogen sulphate NO.H.SO₄.)
This acid is produced by the reaction between H₂SO₄ and any oxide of nitrogen except N₂O (Henry, P. 7, 135; A. Rose, P. 50, 161; Reibling, J. 1861. 152; Kuhlmann, A. Ch. [8] 1, 116; Sestini, Bl. [2] 10, 226). It is also formed in the leaden chambers in making H₂SO₄ (first observed by Clement a. Desormes,

A. Ch. [2] 59, 329). The acid is produced by the reaction of SO₂ with NO or NO₂ in presence of water (Davy); also by burning 1 part S mixed with 2.5-3 parts KNO₂ in moist air (Girard a. Pabst, Bl. [2] 30, 531; Reinsch, N. J. P. 12, 3).

The acid is best prepared by leading NO₂ into well-cooled fuming H₂SO₄ till the whole solidifies, and then drying over H₂SO₄ (Weber, J. pr. 85, 423; 100, 37; Tilden, C. J. 28, 630); or by adding NO₂ in excess to conc. H₂SO₄, washing the crystals with liquid NO₂, and drying in a stream of dry air at 20°-30° or in vacuo (Müller, A. 122,

1; Gaultier de Claubry, P. 20, 467).

Colourless rhombic plates, melting at 73° (Tilden, l.c., gives m.p. 85°-87°), and easily remaining liquid below this temperature (Weltzien, l. 115, 213; De la Provostaye, A. Ch. [2] 73, 362). On melting in air the anhydride (S₂O₃(NO₂)₂) is formed, and the water given off causes decomposition of part of the acid to H₂SO₄ and oxides of N (Michaelis a. Schumann, B. 7, 1075; Fremy, C. R. 70, 61). With dry NaCl reacts to give NaHSO₄ and NOCl; on heating for a little HCl is given off (Tilden, l.c.). Dissolves unchanged in H₂SO₄ (Döbereiner, S. 8, 239; cf. Lunge, B. 12, 1058; 21, 67). SO₂ is without action on dry NO₂(SO₂.OH), but in presence of water, or H₂SO₄Aq with S.G. less than 1.55, decomposition occurs with formation of H₂SO₄, N₂O₃, and other oxides of N, according to Lunge (l.c.).

Potassium nitrosulphonate NO₂(SO₂.QK) seems to be obtained by the reaction of SO₃ on KNO₂, also by adding liquid SO₂ to dry KNO₃; the salt cannot be prepared by neutralising the acid by KOHAq. The salt is decomposed by

acid by KOHAq. The salt is do water (Schultz-Sellack, B. 4, 113).
Nitrosulphonic anhydride

 $S_2O_a(NO_2)_2$ $O < SO_2(NO_2)_1$. Formed by heating $NO_2(SO_2.OH)$ (Michaelis a. Schumann, B. 7, 1075); also by the reaction of SO_2 with NO in absence of O and moisture (H. Rose, P. 47, 605; Brüning, A. 98, 377); also by adding liquid NO_2 to liquid SO_2 without warming (De la Provostaye, A. Ch. [2] 73, 362); by passing electric sparks through a dry mixture of N, O, and SO_2 , or of S vapour and N_2O or NO (Morren, A. Ch. [4] 4, 293; Chevrier, C. R. 69, 136). Hard, regular plates; S.G. 2·14; melts at 217°; may be distilled unchanged at c. 360°. Decomposed by water to NO, H_2SO_4Aq , and HNO_2Aq . $NO_2(SO_2.OH)$ crystallises from a solution in H_2SO_4 .

OXYNITROSULPHONIC ANHYDRIDE

 $S_2O_3O(NO_2)_2$ $\left[O < \frac{SO_2(NO_2)}{SO_2 \cdot O.(NO_2)}\right]$. A white, fusible, crystalline solid; formed by leading vapour of NO_2 into SO_2 till saturated. Gives $S_2O_3(NO_2)_3$ when heated (Weber, P. 123, 833; cf. Thorpe, C_2 , J_3 , J_4 , 297).

C. J. 41, 297).

Nitrosulphonic chloride NO₂(SO₂.Cl). (Nitrosulphuryl chloride.) A white, crystalline solid; formed by the action of SO₂ on NOCl in absence of moisture, also by the reaction of AgNO₂ with SOCl₂ (Thorpe, C. J. 41, 297); dissolves unchanged in fuming H₂SO₄; dissolves in conc. H₂SO₄, giving off HCl, and forming Cl(SO₂.OH) on heating; decomposed by moist air or by water, giving HClAq, H₂SO₄Aq, HNO₂Aq, and NO (Weber, Lc.).

NITROSO-OXY-AMIDOSULPHONIC ACID AND SALTS N(NO.OH)(SO₂.OH). (Nitrosohydroxylamine sulphonic acid

N(NO)(SO, OH)OH. Dinitroso-sulphuric acid SO(NO), (OH)₂ or (SO, OH).(NO), H.) The acid has not been isolated, but the NH, Ba, Pb, K, and Na salts are known. These salts are described as Nitrososulphates under SULPHATES,

p. 581.

AND OXY-AMIDOSULPHONIC ACID (Hydroxylamine SALTS NH(OH)(SO₂.OH). Sulphydroxylamic sulphonic acid [Raschig]. acid [Claus]. Sulphazidic acid [Fremy].) The K salt of the acid is obtained by the action of water on N(OH)(SO₂.OK)₂ (v. OXY-IMIDOSUL-PHONIC ACID AND SALTS, infra). The acid itself is known only in aqueous solutions. Two Ba salts, a K salt, and a Na salt have been isolated. The salts have been investigated by Fremy (A. Ch. [3] 15, 408), Claus a. Koch (A. 152, 336; 158, 52, 194), Raschig (A. 241, 161), and Divers a. Haga (C. J. 55, 760).

Oxy-amidosulphonic acid

NH(OH)(SO,OH)Aq. An aqueous solution of this acid (the acid has not been isolated) is prepared by heating an aqueous solution of N(OH)(SO,OK), (v. infra) to boiling (whereby NH(OH)(SO,OK)Aq, H,SO,Aq, and KHSO,Aq are formed), neutralising by NH,Aq, adding BaCl,Aq, filtering off BaSO, adding BaOA, to ppt. Ba(N(OH)SO,O,Ba.H.O, washing this pp., adding enough H,SO,Aq to ppt. half the Ba in the salt as BaSO, filtering, and so getting a solution of the (soluble) salt (NH(OH)SO,O),Ba. On now heating this solution with an equivalent quantity of H,SO,Aq (the Ba in solution must be estimated) and filtering, a solution of the acid is obtained (Fremy, modified by D. a. H.). The solution of NH(OH)(SO,OH) is fairly stable, but slowly decomposes; in presence of hot acid the decomposition is more rapid, giving 2NH,OH.H,SO,Aq and H,SO,Aq (Raschig, confirmed by D. a. H.).

Oxy-amidosulphonates. The normal of the form NH(OH)(SO₂.OM¹) The normal salts $(NH(OH)SO_2O)_2M^{11}$, where $M^1=K$ or Na, and $M^1=Ba$; there is also a dibarium salt M¹¹ = Ba; there is also a dibarium sait Ba N(OH).SO.O Ba.H₂O. The salts are best N(OH).SO₂O Ba.H₂O. obtained from the solution of the soluble salt (NH(OH).SO,O),Ba.H₂O (v. supra) by adding the equivalent quantity of a sulphate. The dibarium salt is itself prepared as described under the acid (supra); the monobarium salt is obtained by decomposing the di-salt by enough H2SO, Aq to ppt. half the Ba, filtering, and evaporating over H2SO4 (D. a. H.). The oxyamidosulphonates are fairly stable; they are decomposed by heating with acid into hydroxylamine sulphate and H₂SO₄Aq; caustic alkalis produce only sulphite and hyponitrites (the latter rapidly undergo further change, giving off N₂O) (D. a. H., l.c.; v. also C. J. 61, 988 note). Basic oxides, such as CuO and Ag₂O, in presence of alkali produce sulphite, sulphate, and N2O, and at the same time the basic oxide is reduced (D. a. H., l.c., p. 770).

Raschig's sulphasinate (A. 241, 197) (SO_..OK)N(OH).O.(OK)N.(SO_..OK) may be derived from 2NH(OH)(SO_..OH) by replacing 2H

hy O.

OXY-IMIDOSULPHONIC ACID AND SALTS N(OH)(SO₂.OH)₂.. (Hydroxylamine disulphonic acid N(SO₂.OH)₂OH (Raschig). Disulphydroxy-asic acid (Claus).) Only the potassium salt has been isolated.

Potassium oxy-imidosulphonate

N(OH)(SO₂OK)₂·2aq (Claus, A. 158, 83; Raschig, B. 20, 584; cf. Divers a. Haga, C. J. 51, 659). Prepared by passing a rapid stream of SO₂ through well cooled KNO₂Aq made strongly alkaline by KOHAq; allowing N(SO₂OK)₃ (v. Nitritosulphonates, p. 601) to crystallise out, pouring off, and allowing to stand. Large, lustrous crystals; almost insol. cold water, more sol. water at 40°-60°; heated with water gives KHSO₄Aq and NH(OH)(SO₂·OK)Aq (v. supra, OXY-AMIDOSULPHONIC ACID). Three Na salts and several other oxy-imidosulphonates have been prepared by D. a. H. (C. J. Proc. 1893-4. 61).

The salts described by Fremy (l.c.) as sulphazotates, and further examined by Raschig (A. 241, 211), and formulated by him as N(OH)(SO₂.OK)₂.N(OM)(SO₂.OK)₂, where M=K or Na, and N(OK)(SO₂.OK)₂.N(OK)(SO₂.OK)₂, may be regarded as derived from oxy-imidosul-

phonic acid N(OH)(SO2.OH)2.

The oxysulphazotale of Claus (sulphazilinate of Fremy) examined by Raschig (l.c., p. 223) and formulated by him as

(SO₂.OK)₂N

O

N(SO₂.OK)₂, may be looked on as derived from oxy-imidosulphonic acid by the removal of 2H from 2N(OH)(SO₂.OH)₂.

M. M. P. M.

SULPHONO-DI-ACETIC ACID v. DI-METHYL SULPHONE DICARBOXYLIC ACID.

SULPHONO-DI-BUTYRIC ACID

SO₂(CHEt.CO₂H)₂. [152]. Formed by oxidation of sulphido-dibutyric acid (5 g) in neutral solution by KMnO₄ (5 g.) in water (500 g.). Its ether is got from SO₂(CH₂·CO₂Et)₂, Na, and EtI (Lovén, J. pr. [2] 33, 104). Octahedra.

Sulphono-di-isobutyric acid

SO₂(CMe₂·CO₂H)₂· [182°-186°]. Formed by oxidising S(CMe₂·CO₂H)₂ (Lovén). Plates.—BaA" 2½aq. Groups of needles (from hot water). SULPHONO-DIPROPIONIC ACID v. Dr-

ETHYL-SULPHONE DICARBOXYLIC ACID.
SULPHONO-DI-ISOVALERIC ACID

SO₂(C₄H_a,CO₂H)₂. Formed by oxidation of S(C₄H_a,CO₂H)₂ and by the action of PrI (2 mols.) on sulphono-di-acetic ether (1 mol.) and NaOEt (2 mols.) at 120°; the product being saponified by baryta (Lovén, J. pr. [2] 33, 114).—BaA" 7aq.

SULPHO-OXY-BENZOIC ACID v. Ox SULPHO-BENZOIC ACID.

SULPHO-PHENYL-ACETIC ACID C₈H₈SO₃ i.e. CHPh(SO₃H).CO₂H. Formed by saponifying the product of the action of K₂SO₃Aq on a-bromophenyl-acetic ether (Papilsky, J. 1880, 856). Very deliquescent mass.—Salts: K₂A".—CaA".—BaA": plates, m. sol. hot water.—ZnA".—PbA".—CuA": blue plates.—Me(NH₄)A".—CHPh(SO₃NH₄).CO₂Et. Tables, v. sol. water.—CHPh(SO₃K).CO₂Et: thin plates, v. e. sol. Ag.

SULPHO-PHENYL-AMIDO-ACETIC ACID C₃H₂NSO₅ aq i.e. SO₃H.C₂H₄.NH.CH₂.CO₂H. [185°]. Formed by heating phenol (1 pt.) with hippuric acid (1 pt.) and H₂SO₅ (3 pts.) at 140° (Zehenter, M. 5, 882; 6, 523). Monoclinic prisms (containing aq); a:b:c=98:1:1·28. Sol.

water and alcohol. Coloured violet by FeCl₃. Decomposed by HClAq at 140° into phenol, glycocoll, and H₂SO₄. Agua regia gives [1:2:8:5].—BaH₂A"₂ ½aq.- $\mathbf{C}_{\mathbf{a}}\mathbf{H}_{2}(\mathbf{OH})\mathbf{Cl}(\mathbf{NO}_{2})_{2}$ AgHA" 3aq: concentric groups of needles

p-SULPHO-PHENYL-CARBAMIC ACID. The acid ether C₆H₄(SO₃H).NH.CO₂Me [188°] is formed by dissolving methyl phenyl-carbamate in fuming H.SO₄ (Hentschel, B. 18, 979) and also by adding NaOH to a cooled mixture of ClCO₂Me and aqueous p-amido-benzene sulphonic acid (Noelting, Bl. [2] 50, 622).

SULPHO-PHENYL-GLYCOCOLL v. SULPHO-

PHENYL-AMIDO-ACETIC ACID

m-SULPHO-PHENYL-PROPIONIC ACID CoH10SO i.e. $C_6H_4(SO_3H).CH_2.CH_2.CO_2H$. Formed by treating bromo-sulpho-phenyl-propionic acid with sodium-amalgam (Göring, C. C. 1877, 793, 808). Yields m-oxy-benzoic acid by potash-fusion.

exo-Sulpho-phenyl-propionic acid

 $C_6H_5.C_2H_3(SO_3H).CO_2H.$ Formed by boiling cinnamic acid with aqueous K2SO3 for 12 hours (Valet, A. 154,62). Cinnamic aldehyde is converted by K₂SO₃ into PhC₂H₃(SO₃K).CH(OH).SO₃K, which crystallises in needles (containing 2aq) and is converted by boiling dilute H2SO, into exo-sulpho-phenyl-propionic acid (Heusler, B. 24, 1805). Crystals, v. sol. water and alcohol. Converted by boiling conc. KOHAqinto cinnamic acid. Not affected by boiling dilute H₂SO,— KHA". S. 4 at 15°.—K₂A" xaq. Efflorescent crystals.— CaA" xaq. Plates.— BaA" aq.— -Ag.A" aq : white crystalline pp. SULPHO-PHENYL-THIO-CARBAMIC ACID.

Anhydride C,H,NS,O, i.e. C,H,<NH,OS [183°]. Formed from phenyl-thio-carbimide and SO₃ (Magatti, B. 11, 2267). Crystals (from benzene), insol. water, alcohol, and ether. Insol. acids and alkalis. Decomposed by water at 100° into H2S, CO2, and amido-benzene p-sulphonic acid.

SULPHO-PHLORETIC CaHioSO ACID Formed from phloretic (oxy-phenyl-propionic) acid and SO₃ (Nachbaur, J. pr. 75, 45). Sour syrup. — Na₂A" xa₄. — BaA" 3a₄. — MgA" 5a₄. — CaA" 4a₄. Crystalline.

c-SULPHO-PHTHALIC ACID

 $C_6H_3(SO_2H)(CO_2H)_2[3:2:1]$. (a)-Sulpho-phthalic acid. Formed by oxidising naphthalene (a)-sulphonic amide by KMnO₄ (Remsen, Am. 5, 107), and got also, in small quantity, together with the (β) - acid, by the action of fuming H_2SO_4 on phthalic acid (Rée, C. J. 49, 514). Minute crystals, v. sol. water, m. sol. alcohol. Sodafusion gives c-oxy-phthalic acid.—Ba,A'''₂ 8aq. Needles, sl. sol. hot water.—PbHA'''1½aq.—Ag₂KA'''2aq. Ppd. by adding AgNO, to a solution of the K salt (Stokes, Am. 6, 280).

Amic acid CaH, NSO, i.e. [155°-160°]. $C_6H_3(SO_2NH_2)(CO_2H)_2$ Formed by oxidation of naphthalene (a)-sulphonic arride by alkaline KMnO. Thick needles (containing aq). At 155° it splits up into H₂O and the anhydride. Conc. HClAq at 150° forms c-sulphophthalic acid.—KHA": slender needles, v. sl. sol. cold water.-K2A". [300°]. Amorphous, v. e. sol. water. Yields, when heated, the compound C.H.(COCI) SO₂ N.POCl₂ [120°-126°] whence

MeOH produces $C_eH_s(CO_2Me) < \frac{SO_2}{C(OMe)_a} > NH$ [144°] (Stokes, Am. 6, 274).—PbA".—AgHA": needles.—Ag₂A". Insol. hot water.

Anhydride of the Amic Acid

 $C_6H_3(CO_2H) < SO_2 > NH.$ Formed as above. Begins to sublime at 200°, but is not melted at 240°. Its silver salt C,H,AgNSO, is converted by MeI into $C_6H_3(CO_2H) < SO_2 > NMe [191° cor.]$.

The compound $C_6H_3(CO_2Me) < SO_2 > NMe$ [180°

cor.] has also been prepared. i-Sulpho-phthalic acid

 $C_6H_3(SO_3H)(CO_2H)_2[4:2:1]$. (β)-Sulpho-phthalic acid. (γ)-Sulpho-phthalic acid. [140°] (when hydrated). The chief product of the sulphonation of phthalic acid or anhydride at 100°-200° (Loew, A. 143, 257; Rée, B. 18, 1629). Formed also by oxidation of naphthalene (3)-sulphonic amide (Remsen, Am. 5, 110) and by the action of hot HNO₃ (S.G. 1-3) on potassium di-nitro-(a)-naphthol sulphonate (naphthol yellow S) (Graebe, B. 18, 1126; Rée, C. J. 49, 516). Crystalline (containing aq), very hygroscopic, v. sol. water and alcohol, insol. ether. At 180° it yields the anhydride C₈H₄SO₆. Soda-fusion forms i-oxyphthalic acid. The K salt fused with sodium formate yields trimellitic acid. Heated with resorcin it yields fluorescein sulphonic acid. PCl_5 forms $C_8H_3(SO_2Cl)(CO_2H)_2$ [170°] C_eH_s(SO₂Cl) < CO >O and C_eH_sCl < CO >O. The mono-chloride is converted by NH_s into $C_6H_3(SO_2NH_2)(CO_2H)_2$ [192°-202°] which crystallises in plates, sol. water, alcohol, and ether.

Salts.— KH_2A''' 2aq. Needles, v. sol. water. $-K_2HA'''$ 2aq.— $(NH_4)_2HA'''$ $\mathbf{1}_2^1$ aq. At 200° it yields $C_0H_3(SO_3NH_4) < \stackrel{CO}{CO} > NH$ [c. 300°]. Crystallising in monoclinic prisms.—Ba₃ Λ''' , 2aq.—BaH₄ Λ''' ₂5aq. S. 5 at 15°; 50 at 100°. A 250° it yields Ba($C_8H_3SO_9$)₂.—BaH Λ''' 2aq.

s-Sulpho-isophthalic acid $[258^{\circ}].$ $C_aH_3(SO_3H)(CO_2H)_2[5:3:1].$ Obtained by sulphonation of isophthalic acid (Aronstein a. Kramps, B. 13, 489; Lönnies, B. 13, 704). Long deliquescent needles (containing 2aq). Potash-fusion yields s-isophthalic acid. KH₂A''' 3aq: long needles.—K₃A''' xaq: prisms. -Ba,A''',8aq: needles, v. sol. water.

i-Sulpho-isophthalic acid [244°]. Formed by $C_6H_3(SO_3H)(CO_2H)_2[4:3:1].$ oxidation of m-xylene (a)-sulphonic acid (Jacobsen a. Lönnies, B. 13, 1556), and by oxidation of C_sH₃Me(SO₂NH₂).CO₂H (Remsen a. Iles, Am. 1, 114; Remsen a. Coale, Am. 3, 206). Hygroscopic needles (containing 2aq), v. e. sol. water. Potashfusion yields (a)-oxy-isophthalic acid.—Salts: KH,A" 2aq: needles, sl. sol. cold water. S. 1.59 at 26°.-K, A'".-CaHA" 4}aq. Crystals.-BaHA''' 3aq: small needles. S. 073 at 23.5°.

—BaHA''' 4aq.—Ba₃A'''₂ 3aq.

Amic a cid C₆H₂NSO₆ i.e.

C₆H₃(SO₂NH₂)(CO₂H)₂. Formed by oxidation of C₆H₃Me(SO₂NH₂).CO₂H by KMnO₄ (Remsen, B. 11, 464; 12, 1436; Am. 1, 122; 3, 209). When set free from its salts it changes at once to the anhydride C₆H₂(CO₂H)<SO₂NH [284°], S. 45

at 10°.—KHA" aq. S. 2.3 at 26°.—K2A" 4aq: v. e. sol. water. — CaH_A"_2 2aq. — CaA" 5aq. — BaH_A"_2 2aq: monoclinic tables.—BaA" 3aq.—Ag.C₂H₄NSO₆: crystalline pp. (Jacobsen, B. 12, 2320).

c-Sulpho-isophthalic acid. Amic acid C₆H₃(SO₂NH₂)(CO₂H)₂[2:3:1]. Formed by oxidation of the corresponding m-xylene sulphonic amide (Jacobsen, B. 11, 902). Its acid potassium salt is sl. sol. water.

a-SULPHO-PROPIONIC ACID C3H6SO5 i.e. CH₃.CH(SO₃H).CO₂H. Formed by boiling achloro-propionyl chloride with aqueous ammonium sulphite and also by warming propionic acid with CISO, H (Kurbatoff, B. 6, 563; A. 173, 5). Syrup, v. sol. water and alcohol.—K2A" aq: needles (Rosenthal, A. 233, 27).—(NH_J)₂A" aq : prisms.—BaA" 2aq. S. (of BaA") 7.45 at 18°.
—CaA" 2aq.—CdA" 2aq.—Ag₂A" : small needles. Got also by mixing propionic anhydride with SO₃ (Franchimont, R. T. C. 7, 27).

B-Sulpho-propionic acid CH₂(SO₃H).CH₂.CO₂H. [68°]. Formed by boiling B-iodo-propionic acid with an aqueous solution of (NH_a)₂SO₃. Got also by the action of ammonia-cal AgNO₃ on the compound of acrolein with NaHSO, (Rosenthal, A. 233, 15) and obtained likewise by the oxidation of thiohydantoic acid NH:C(NH₂).S.CH₂.CH₂.CO₂H (Andreasch, M. 6, 838; 7, 169). Hygroscopic crystals, sol. water and alcohol. Decomposes at 150°. Successive treatment with PCl, and with tin and HClAq converts it into sulphydro-propionic acid.

Salts.— K₂A" aq.— KHA" aq.— Na₂A" aq.— (NH₁)₂A" 4aq. Hygroscopic. — HNH₁A". — BaA" 5aq.—BaH₂A"₂.—SrA" 5aq. — CaA" 2½ aq. — CaA" aq.—MgA" 4aq. — ZnA" 4aq. — CuA". — MnA" 4aq.—PbA". — CdA" aq. — Ag₂A" ½ aq. — HAgA" \(\frac{1}{2} \text{aq}. \)
Ethyl ether \(\text{Et}_2 \text{A"}. \)
Liquid.

Sulpho-dipropionic acid is DI-ETHYL-SUL-PHONE DICARBOXYLIC ACID.

SULPHO-PROPYL-BENZOIC ACID v. SULPHO-CUMINIC ACID.

SULPHO-ISOPROPYL-SUCCINIC ACID [c. 167°]. $CMe_2(SO_2H).CH(CO_2H).CH_2.CO_2H.$ Got by action of HNO₃ on sulphocamphylic acid (Königs a. Hoerlin, B. 26, 2044). When heated in vacuo at 170° it splits up into water, SO2, and terebic acid. Tables (containing Saq), v. e. sol. water.

SULPHO-PYROMUCIC ACID C.H.SO. i.e. CH:C(CO₂H) >0. Formed by dissolving pyromucic acid in cold fuming H2SO4, and got also by the action of zinc-dust and ammonia on dibromo-sulpho-pyromucic acid (Schwanert, A. 116, 268; Hill a. Palmer, B. 18, 2095; Am. 10, 373, 409). Deliquescent prisms.—K,A''4aq.—KHA''.—Na,A'' 5aq.—NaHA''aq.—BaA''4aq. Small prisms. Yields fumaric acid when heated with excess of bromine.—Salts: BaH2A", 4aq. BaH₂A", 6aq.—CaA" 8aq.—PbA" 2aq.—Ag₂A".

Amide C, H2SO4(NH2), [213°]. Crystalline.

(β)-Sulpho-pyromucic acid C,H2O(SO2H)(CO2H). Formed by sulphonation of bromo-pyromucic acid and debromination of the product by zinc-dust and NH, (H. s. P.).—
Salts: K₂A"2½aq.—CaA"2aq.—BaH₂A", 8aq.
—BaA"8aq.—BaA"aq: small concentric prisms.

SULPHO - PYROTARTABIC ACID C,H,SO, i.e. C₅H₅(SO,H)(CO.H)₂. Formed by boiling ita-, citra-, and mesa- conic acids with aqueous K₂SO₂ (Wieland, A. 157, 34). Crystalline, v. e. sol. water.—Ca₂A'''₂7aq. Sl. sol. cold water.
SULPHO-QUINOLINE CARBOXYLIC ACID

(a)-Sulpho-cinchoninic $C_9H_5N(SO_3H).CO_2H.$ acid. Formed by heating cinchonic acid with SO₃ or with H₂SO₄ and P₂O₅ at 180° (Weidel a. Cobenzi, M. 1,844). Triclinic crystals (containing aq), m. sol. hot water, insol. alcohol and ether. Tastes bitter. Potash-fusion gives (a)oxy-cinchoninic acid.—(NH,)2A" 2aq. Monoclinic crystals; a:b:c = 1.19:1:3.53; $\beta = 95.14.$ CaA" 2 aq. Monoclinic crystals.—BaA" 3aq.—

cinchoninic acid. Formed at the same time as the preceding acid (Weidel, M. 2, 565) and prepared by heating cinchoninic acid (1 pt.) with H₂SO₄ (7 pts.) at 300° (Von Georgievitch, M. 8, 639). The same acid (crystallising with aq) appears to be formed by oxidation of benzylidenelepidine sulphonic acid by alkaline KMnO, (Busch a. Koenigs, B. 23, 2683). Colourless needles (containing 2aq), v. sol. hot water. Very bitter. Potash-fusion yields (3)-oxy-cinchoninic acid.—(NH₄)HA'' 2aq.—BaA''aq.—PbA'' 4aq.

SULPHO-SALICYLIC ACID v. OXY-SULPHO-BENZOIC ACID.

SULPHO-SUCCINIC ACID C.H.SO, i.e.

 $CO_2H.CH_2.CH(SO_3H).CO_2H.$

Formation.—1. By exposing cooled succinic acid to SO_s vapour (Fehling, A. 88, 285; 49, 203).—2. By boiling fumaric and maleïc acids with aqueous K₂EO₃ (Credner, Z. [2] 6, 77; Strecker a. Messel, A. 157, 15; Z. [2] 6, 459, 671).—8. From succinyl chloride and Ag₂SO₄ (Carius a. Kämmerer, A. 131, 167).—4. By oxidation of thio-malic acid with HNO, (Carius, A. 129, 9).

Properties.-Deliquescent mass, v. e. sol. water, alcohol, and ether. Yields fumaric acid

when fused with potash.
Salts.—K,A" aq: efflorescent crystals.— Salts.—K₃A''' aq: efflorescent crystals.—
K₃A''' 2½aq.—K₃A''' 1½aq. Crystals.—K₄A'''.

k₂HA''' 2aq.—K₄A''', 1½aq.—(NH₄)₂A''' aq.—
(NH₄)H₂A''' aq: crystals.—Ba₃A'''₂ (dried at 100°).

Pp.—Ca₃A'''₂ 6aq.—Pb₄A'''₂ 4aq.—Pb₂A'''₂ 3aq.—
Pb₄OA'''₂.—Pb₅O₂A'''₂.—Ag₅A''': m. sol. water.

SULPHO-TEREPHTHALIC ACID C₂H₃SO₇

i.s. C₀H₃(SO₃H)(CO₂H)₂[2:4:1].

Formation.—1. By heating terephthalic acid with fuming H₂SO, at 250° (Ascher, A. 161, 2; Schoop, B. 14, 223).—2. By oxidation of sulphop-toluic acid (Remsen a. Burney, Am. 2, 410; Weber, B. 25, 1740), of p-xylene sulphonic acid (Remsen a. Kuhara, Am. 2, 414), and of $C_8H_2Me(SO_2NH_2).CO_8H$ (Hall a. Remsen, B. 12, 1432; Am. 2, 56).

Properties.—Hygroscopic needles or tables.
Salts.—KH_A''' aq. Needles (Remsen a. Keiser, Am. 5, 170).—KH_A''' aq (W.).—K_A''' aq.
—BaHA''' aq.—CaHA''' 1½aq.—BaHA''' 1½aq.—BaHA''' 2aq.—Ag,HA'''.

Amic acid CaH_(SO_NH_2)(CO_2H)_r. Formed by oxidising payleng sulphonic amide with alka-

by oxidising p-xylene sulphonic amide with alkaline K.FeCy. (Noyes a. Walker, Am. 9, 94).

 $Imids C_{s}H_{3}(CO_{2}H) < SO_{2} > NH. [284°] (W.);$ [299° cor.] (N. a. W.). Formed by oxidation of $C_8H_3Me(SO_2NH_2).CO_2H$ by $KMnO_4$. Short thin prisms (from ether), m. sol. cold water. AgNO₃ gives a pp. insol. HNO₃.—KC₃H₄NSO₅ aq.—BaC₃H₃NSO₅ 3aq. Scales, v. sl. sol. water.—Ag₂C₅H₃NSO₅.

SULPHO-TOLUENE DICARBOXYLIC ACID. A mic acid $C_0H_2Me(SO_2NH_2)(CO_2H)_2[1:4:3:5]$. [c. 800°]. Formed by oxidation of ψ -cumene sulphonic amide (Jacobsen a. Meyer, B. 16, 190). Needles, sol. water, alcohol, and ether. -BaA" $2\frac{1}{2}$ aq: small prisms, sl. sol. water.

SULPHO-0-TOLUIC ACID C,H,SO, C₆H₃Me(SO₃H).CO₂H [2:3:1]. Formed by heating o-toluic acid with H₂SO₄ (5 pts.) for 3 hours at 160° (Jacobsen a. Wierss, B. 16, 1960). Crystalline, v. e. sol. water. Potash-fusion yields o-oxytoluic acid.

Sulpho-o-toluic acid. Amic acid C₆H₃Me(SO₂NH₂).CO₂H [2:5:1]. [243°]. Formed, together with about an equal quantity of the (2,4,1)- isomeride, by oxidation of m-xylene o-sulphonic amide by alkaline KMnO₄ (Jacobsen, B. 14, 38). Long needles, sol. alcohol, ether, and water. Potash-fusion yields the corresponding oxy-toluic acid. Further oxidised by KMnO, to C,H3(SO2NH2)(CO2H)2.

Sulpho-o-toluic acid. Amic acid C₆H₂Me(SO₂NH₂).CO₂H [2:4:1]. [217°]. Formed as above. Long needles, sl. sol. cold water, v. sol. alcohol and ether.—KA': crystals.

s-Sulpho-m-toluic acid C_eH₃Me(SO₃H).CO₂H [3:5:1]. Formed, together with the (3,4,1)- isomeride, by sulphonating m-toluic acid with fuming H2SO, at 180° (Jacobsen, B. 14, 2355).

Sulpho-m-toluic acid. Amic acid $C_6H_3Me(SO_2NH_2).CO_2H$ [3:4:1]. [248°]. 15°. Formed by exidation of m-xylene sulphonic amide (Remsen, Am. 1, 37; 3, 205; Jacobsen, B. 11, 895). Needles (from water), m. sol. alcohol, sl. sol. ether. Potash-fusion gives (4,3,1)-oxy-m-toluic acid. KMnO₄ yields sulpho-isophthalic acid.—CaA'₂1½aq: small concentric needles.—BaA'₂4aq.—BaA'₂5aq.—AgA': needles (from hot water).

Sulpho-m-toluic acid. Amic acid C₂H₃Me(SO₂NH₂).CO₂H [3:2:1]. [203°]. Formed by oxidation of the corresponding m-xylene sulphonic amide [96°] (J.). Converted by potash-fusion into the corresponding oxy-toluic acid.

Sulpho-p-toluic acid CaH, Me(SO,H).CO2H [4:3:1].

Formation. — 1. By oxidation of thiocarvacrol (Flesch, B. 6, 480; Bechler, J. pr. [2] 8, 170).—2. By oxidation of cymene sulphonic acid (Remsen, Am. 2, 411; R. Meyer a. Baur, A. 220, 18).—3. From p-toluic acid and SO₃ (Fischli, B. 12, 616).—4. By oxidation of p-xylene sulphonic acid (Remsen, Am. 8, 264). Needles (containing 2aq). Not hygroscopic. V. sol. water, insol. ether. Decomposes at 185°–190° without melting. Potash-fusion gives oxylenged 1900° Containing. Conc. HClAq at 190° forms p-toluic acid [204°]. n-toluic acid.

Salts. - KHA" Saq. - KHA" 2sq.-

Concentric needles, sol. hot water. Not melted at 310°.—KHA" ½aq.—BaA" aq: nodules.

Imids C.H.(CO.H) $< \frac{SO_2}{NH}$, [284°] (W.):

Amide C₆H₃Me(SO₂NH₂).CONH₂ aq. [218°]. Amic acid C, H, Me(SO, NH,).CO, H. [2670]. Formed by oxidation of cymene sulphonic amide and of p-xylene sulphonic amide by chromic acid mixture (Remsen a. Hall, Am. 2, 50; B. 11, 229; Noyes a. Walker, Am. 9, 98). Needles, sol. cold alcohol, insol. ether, sl. sol. cold water.-BaA'₂2aq. — CaA'₂4aq. — MnA'₂5aq: needles. Probably the same amic acid [242°] is formed by oxidising p-butyl-toluene sulphonic amide by KMnO, (Kelbe a. Baur, B. 16, 2565). Sl. sol. water, nearly insol. alcohol.

Sulpho-p-toluic acid C.H.Me(SO.H).CO.H [4:2:1]. [182°] (W.); [190°] (F.). Got by oxidising thio-thymol (Fittica, A. 172, 329) and by evaporating its imide with HClAq (Weber, B. 25, 1741). Crystals (containing Baq).—NH,HA".
—MgA".—Ag₂A".

Anhydride $C_6H_8Me < {}^{CO}_{SO_2} > 0$. [97°]. Got by treating the acid with AcCl.

Amic acid CaHaMe(SO3H).CONH2. [186°]. Prisms (containing aq).—NH,A'. Got from the anhydride in benzene by the action of dry NH2.-AgA' aq: scales (from water).

Amic acid C₆H₈Me(SO₂NH₂).CO₂H. [185°]. Got by boiling the imide with water (Weber, B. 25, 1739). Yellowish crystals. Probably identical with the preceding amic acid.—BaA', 2aq.—AgA'.—MeA'. [145].—EtA'. [95°].

Imide C₆H₃Me<CO $^2>$ NH. Methyl - sac- $\begin{array}{lll} \textit{charin.} & \textbf{(246°]}. & \textbf{Got from } p\text{-toluidine sulphonic} \\ \textbf{aoid} & \textbf{vid} & \textbf{C}_0\textbf{H}_3\textbf{MeCy.SO}_2\textbf{H}, & \textbf{C}_0\textbf{H}_3\textbf{MeCy.SO}_2\textbf{Cl}, \\ \textbf{C}_0\textbf{H}_3\textbf{MeCy.SO}_2\textbf{NH}_2, & \textbf{C}_0\textbf{H}_3\textbf{Me}(\textbf{CO}_2\textbf{H})(\textbf{SO}_2\textbf{NH}_2), \\ \end{array}$ the last body being heated (Kreis, G. P. 48,583; B. 22, Ref. 719; Weber, B. 25, 1737). Sl. sol. cold water, v. sol. alcohol, ether, and alkalis. Yields C.H. Me CO2 NAg, whence MeI forms $_{\rm GH_2Me}$ $<_{\rm CO}^{\rm SO_2}$ NMe [153°], while EtI gives

 $C_eH_aMe < SO_2 > NEt$ [106°] (Weber). The imide is converted by warming with aqueous KOCl into p-toluidine sulphonic acid.

Di-sulpho-o-toluic acid C₈H₂Me(SO₃H)₂CO₂H [2:3:5:1]. Formed by sulphonation of o-toluic acid with H₂SO₄ containing SO₅ (Jacobsen a. Wierss, B. 16, 1960). Minute needles, v. e. sol water.— *Ba₃A'''₂: amorphous, v. sol. water.

Di-sulpho-p-toluic acid $C_0H_2Me(SO_2H)_2CO_2H$. Formed by heating p-toluic acid with fuming H,SO, and P,O, (Weinreich, B. 20, 982).—BaHA''' 5aq. Crystals.

Reference.-NITRO-SULPHO-TOLUIC ACID.

C,H,80, SULPHO-UVITIC ACID Formed by $C_8H_2Me(SQ_8H)(CO_2H)_2$ [5:6:3:1]. evaporating its amic acid with conc. HClAq (Jacobsen, A. 206, 185). Needles (from dilute H.SO.). Potash-fusion yields oxy-uvitic acid.—
KH.A" 2aq.—Ba.A". S. 3·23 at 12·5°. Needles.

A mic a cid C.H.Me(SO.NH.)(CO.H).

product of oxidation of mesitylene sulphonic amide with KMnO, (Hall a. Remsen, Am. 2, 136; Jacobsen). When set free from its salts it changes at once into the anhydride C.H.NSO. [270°], S. 5 at 100°.-KHA".-BaA".-BaA"8aq.

SULPHO-ISOVALERIC ACID CaH10SO, i.s. C.H. (SO.H).CO.H. Formed by heating isovaleric acid (1 pt.) with CISO H (1 pt.) at 150° (De Varda, G. 18, 91). Deliquescent crystalline Its aqueous solution partially decom-

poses when heated.—BaA" aq.—PbA" 2aq. SULPHOXIDES. Organic compounds R.SO.R' formed by the action of conc. HNO3 on sulphides. Decomposed by heat. Reduced by Zn and H2SO4 and by HI to sulphides. Attacked by PCl, sulphides being formed. They reduce KMnO4, being converted into sulphones. Sulphoxides containing monovalent alcohol radicles form unstable compounds with HNO3, such as Et2SO,HNO3 (Beckmann, J. pr. [2] 17, 475). SULPHO-XYLENE CARI

CARBOXYLIC ACID. Amic acid $C_0H_2Me_2(SO_2NH_2).CO_2H$ [1:3:4:5]. [268° cor.]. Formed by oxidising ψ -cumene sulphonic amide (Jacobsen a. Meyer, B. 16, 190). Long needles, sl. sol. hot water. - KA'aq.

BaA. 22 aq.

SÜLPHUR. S. At. w. 31.98. Mol. w. 63.96; probably also 191.88 and 255.84 (v. infra). The following data apply to ordinary rhombic S. Melts at c. 115°; different observers give from 111° to 115° (v. Helff, Z. P. C. 12, 219; Spring, A. Ch. [5] 22, 170; Kopp, A. 93, 129; Brodie, J. pr. 62, 336; Gernez, C. R. 83, 217; for m.p. at high pressures v. Hopkins, J. 1854.
48). Boils at 440° (Dumas, A. Ch. [2] 36, 83;
Troost a. Hautefeuille, C. R. 76, 76, 219). Callendar a. Griffiths (T. 182, 119) give 444.53° as · within ·1° of the true temperature of the vapour of sulphur boiling freely under a pressure of 760 mm.' (This determination was made with Pt resistance thermometers with great care, but unfortunately no special precautions were taken to insure the purity of the S used. For other determinations v. Carnelley's Melting- and Boiling-point Tables, 1, 11. For b.p. under different pressures from 1 to 760 mm. v. Monckman, Pr. 46, 136.) S.G. c. 203 (for numerous data v. Clarke's Table of Specific Gravities, 2nd ed., 9). S.G. molten S 1.801 to 1.815 (Playfair a. Joule, C. S. Mem. 3, 76); S.G. at b.p. 1.46 to 1.51 (Ramsay, C. J. 35, 471). V.D. varies from c. 122 to c. 62; v. infra, Molecular Weight of S. S.H. (17°-45°) ·163 (Kopp, T. 1865. 71); ·(0°-100°) ·1776 (Bunsen, P. 141, 1). S.H. (liquid) ·2346 (Person, A. Ch. [3] 21, 295). Latent heat of fusion = 9.368 (for 1 g. S) (Person, A. S.Y. (15°-20) (Per son, Lc.). S.V.S. 15-9. S.V. of S in combina-tion varies from 28-6 to 22-6; S.V. at b.p. 21-6 (Ramsay, C. J. 35, 471). C.E. (linear) 00006413 (at 40°) (Fizeau, C. R. 68, 1125; v. also Kopp, A. 93, 129; Pisati, G. 1874. 93; Spring, J. 1881. 1085; Moitessier, J. 1866. 27, who gives C.E. for each c. 20° from 110° to 440°). S. in CS₂; 16.54 at -11°, 18.75 at -6°, 23.99 at 0°, 87.15 at +15°, 41.65 at 18.5°, 46.05 at 22°, 94.57 at 38°, 146.21 at 48.6°, 181.24 at 55°, 40°, 181.24 at 55°, 41°, 181. 181.34 at 55° (Cossa, B. 1, 138; Payen, C. R. 84, 456, 508). For S.G. of solutions in CS₂ v. 84, 456, 508). For S.G. of solutions in CS₂ v. Macagno (C. N. 43, 192). For S. in C₆H₆ and other solvents v. Pelouze (C. R. 68, 1179; 79,

Refraction-equivalent $\left(\frac{\mu-1}{d}$. at.w.) for

D line 16.0 (solid), 16.47 (liquid), 16.1 (gaseous, for C line), 16.0 (in solution), 16.0 (calculated from data for SCl₂), 16.1 (from data for S₂Cl₂)

(Gladstone, P.M.[5]35,204). H.C. $[S,O^7] = 71,080$ (Th. 2, 247). The E.C. of S is very small; it varies much with temperature (v. Monckman, Pr. 46, 136). The fundamental form of rhombic S is a simple pyramid, or elongated octahedron; a:b:c = 8106;1;1·898. For emission-spectrum v. Salet, C. R. 68, 404; 73, 559; Bl. [2] 11, 302; Mulder, J. pr. 91, 112; Barrett, J. 1865. 138; Seguin, C. R. 53, 1272; Chautard, C. R. 79, 1123; Plücker a. Hittorf, J. 1863. 109. For absorption-spectrum of S vapour v. Salet, C. R. 74, 865; Gernez, C. R. 74, 803; Ciamician, W. A. B. 77 [2] 839. For vapour-pressures of S vapour v. Regnault (J. 1863. 65). Ignition point c. 250° (Hill, C. N. 61, 125; Blount, C. N. 61, 153).

The following data apply to monoclinic S. Melts at 117.4° (Gernez, C. R. 83, 217); at 120° (Brodie, J. pr. 62, 336). S.G. 1.982 (Marchand a. Scheerer, J. pr. 24, 129); 1.958 (Deville, J. 1, 365). S.V.S. 16.4. H.C. $[S,O^2] = 71,720$ (Th. 2, 247; Petersen, Z. P. C. 8, 601). For volumes occupied at different temperatures v. Toepler (W. 47, 169). Crystallises in secondary forms of a monoclinic prism; a:b:c=1.004:1:1.004, angle $b:c=84^{\circ}$ 14'. Sol. CS_2 , from which solution

rhombic S crystallises out.

Insoluble sulphur is amorphous; insol. CS2; S.G. c. 2.04 (Troost a. Hautefeuille, C. R. 69, 248); S.G. after compression at 8,000 atmos. 1.9556 at 0°, 1.9643 at 100° (Spring, Bl. Acad. Belg. [3] 2, 83).

Cf. Allotropy of Sulphur (p. 609). Occurrence.—Native, in beds, in Sicily, Mexico, New Zealand, &c.; in the lava fissures of volcanic districts; in small quantities in the mud from the sea-bottom (Buchanan, Pr. E. 1891. 1). Many metallic sulphides also occur native-e.g. sulphides of Sb, As, Cu, Fe, Pb, Hg, and Zn. Sulphates occur in large quantitiese.g. gypsum, celestine, heavy spar; these and other sulphates are found in the earth's crust, in the sea, and in many river and spring waters. (For more details v. DICTIONARY OF APPLIED CHEMISTRY, iii. 682.) S is a constituent of albumen, hair, feathers, horn, and some other parts of animals; it is also found in many plants. Small quantities of S compounds are found in the atmosphere near volcanoes. According to Young sulphur occurs in the solar atmosphere (Am. S. [3] 4, 356).

Sulphur has been known from very early times. The quantitative work of Lavoisier, in 1772, on burning S showed this substance to be an element, but it was not finally placed on the list of elements until after the experiments of

Gay-Lussac and Thénard in 1809.

Formation.—1. By the interaction of SO_2 and H_2S ; $2SO_2+4H_2S=3S_2+4H_2O.—2$. By the partial oxidation of H_2S , either by incomplete combustion or by exposure of H2SAq to a limited quantity of air; $2H_2S + O_2 = 2H_2O + S_2$. Also by the oxidation of FeS by exposure to moist air; $2FeS + 3O = Fe_2O_3 + S_2$. By distilling certain metallic sulphides out of contact with air, e.g. $3FeS_2 = Fe_3S_4 + S_2 - 4$. By decomposing solution of an alkali or alkaline polysulphide by acid; $CaS_{a}Aq + 2HClAq = CaCl_{a}Aq + H_{a}S = 2S_{a}$ Also by adding acid to solution of a thiosulphate; e.g. $Na_{s}S_{2}O_{s}Aq + 2HClAq =$ $2NaClAq + H_{s}O + SO_{s}Aq + S.$ By decomSULPHUR. 607

posing S_2Cl_2 by water $(2S_2Cl_2+2H_2O+Aq=4HClAq+SO_2+3S)$.—6. When sulphates, e.g. $CaSO_4$, are in prolonged contact with decaying organic matter (such as plant-leaves) they are reduced to sulphides, and if these come into contact with acids S is among the products of the reactions that occur. For a fuller account of some of these processes, whereby native S is probably formed, v. DICTIONARY OF APPLIES (HEMISTRY, iii. 683. For methods of forming the various varieties of S v. infra, Preparation.

Preparation.—1. Ordinary S is dissolved in pure, dry CS₂; a part of the CS₂ is distilled off, and the rest is allowed to deposit crystals of S. This process is repeated once or twice; the S crystals are powdered, kept in vacuo for some days to remove all CS2, and then repeatedly distilled in vacuo, the middle portion of the distillate being collected each time.—2. Ppd. S is distilled several times in S2Cl2 to remove H compounds; the distillate is then repeatedly distilled in vacuo, the middle portion only being collected; the distilled S is finely powdered, repeatedly washed with water, dried, distilled over a few small pieces of pure Zn in vacuo (to remove traces of S₂Cl₂), and then repeatedly distilled in vacuo.—3. S is ppd. by adding HClAq to pure Na₂S₂O₃Aq, the pp. is repeatedly washed till free from chlorides, dried, distilled repeatedly, and finally distilled several times in vacuo (v. Monckman, Pr. 46, 149).

Ordinary rhombic sulphur is prepared by melting S and allowing to cool very slowly at 90° (Schützenberger, C. R. 66, 746), or by melting and throwing in a crystal of rhombic S when the liquid has cooled nearly to the crystallisation point (Gernez, C. R. 83, 217); also by crystallising from CS₂; and, in very well formed crystals, by saturating pyridine or picoline with H₂S and allowing to stand (Ahrens, B. 23, 2708).

Monoclinic sulphur is prepared by melting a considerable quantity of roll sulphur in a Hessian crucible, allowing to cool till a crust forms on the surface, piercing this crust, and pouring out the S that is still liquid; the walls of the crucible are covered with monoclinic crystals. This form of S can also be prepared by evaporating alcoholic solutions of (NH₄)₂S (Mallard, J. 1885. 383; Ruys, J. 1884. 336; Gernez, C. R. 100, 1539; 101, 312); also, along with rhombic crystals, from solution of S in boiling alcohol, benzene, &c. (Maquenne, Bl. [2] 41 238). By melting S, and throwing in a crystal of the monoclinic form when the liquid is near the crystallisation point, the S solidifies in monoclinic crystals (Gernez, C. R. 83, 217).

Soft soluble sulphur is prepared by decomposing S₂Cl₂ by water, or Na,S₂O₂Aq or a soluble polysulphide by a limited quantity of acid, or many metallic sulphides by fuming HNO₂, or SO₂Aq by H₂SAq (Weber, A. 141, 432; Rose, P. 47, 166; Deville, Ph. C. 1848. 200; Fordos a. Gélis, Ph. C. 1854. 294). This preparation is not homogeneous; it contains both soluble and insoluble S.

Insoluble sulphur (insol. in CS₂) is prepared by heating ordinary S nearly to boiling and then cooling rapidly (most easily by slowly pouring into a large quantity of cold water), rubbing the plastic mass so obtained with a glass rod, under water, till it becomes hard, and

removing soluble S by treatment with warm CS₂ (cf. Deville, Ph. C. 1848. 200). Insoluble S is generally present in 'flowers of sulphur' (which is formed by rapidly cooling vapour of S); by treating this with CS₂ the soluble S is removed, and the insoluble form remains. Soluble S becomes covered with a film of the insoluble variety by exposure to sunlight or electric light when melted (Berthelot, J. pr. 31, 396; Lallemand, C. R. 70, 182). Insoluble S is also said to be obtained by decomposing Na₂S₂O₃Aq by HClAq, dissolving the ppd. S in CHCl₃, evaporating, and keeping the crystals that separate for some time (for details v. Engel, C. R. 112, 866; Friedel, C. R. 112, 834). Insoluble S is also formed, mixed with the soluble variety, by the incomplete combustion of H2S or CS₂; by decomposing H₂S by fuming HNO₃, aqua regia, Fe₂Cl₂Aq, S₂Cl₂, or CrO₃Aq; and by the reaction of HNO, Aq, SO,, or halogens with melted S (v. Wöhler, A. 86, 373; Vogel, J. Ph. [3] 29, 433; Schiff, A. 115, 68; Nöllner, A. 108, 19; Dietzenbacher, C. R. 56, 39). Insoluble S is also formed by decomposing thiosulphates by acid, or S2Cl2 or S2Br2 by water, &c., and washing the soft magma so obtained with CS2 to remove soluble S (cf. Preparation of soft soluble sulphur, supra; and v. Weber, A. 141, 432; Rose, P. 47, 166; Deville, Ph. C. 1848. 200).

Colloidal sulphur, soluble in water, is prepared by passing H₂S into SO₂Aq at a little above 0° till all the SO₂ is decomposed, filtering, and concentrating over KOH in vacuo. The yellow solid so obtained is sol. water, but changes to ordinary S on keeping (Debus, C. J. 53, 282). According to Engel (C. R. 112, 866) this form of S exists in the solution obtained by adding 1 vol. Na₂S₂O₃Aq, saturated at the ordinary temperature, to 2 vols. HClAq saturated at 25°-30° and let cool to c. 10°, and filtering from NaCl that separates. This form of S has not been isolated in a state of purity.

Regarding the formation of the varieties of S one from the other, cf. Allotropy of sulphur, p. 609.

Properties.—S exists in several modifications. The chief are (1) soluble in CS_2 , (2) insoluble in CS_2 , (3) soluble in water.

1. Soluble sulphur exists in two, perhaps in three, varieties, differing in S.G., crystalline form, &c.

A. Ordinary rhombic (octahedral) sulphur is a pale-yellow, tasteless, very brittle solid. By crystallisation from CS2 it forms clear, yellow, transparent, lustrous crystals; the colour becomes paler at low temperatures, until at - 50° the crystals are almost colourless (Schönbein, J. pr. 55, 161); by immersion in boiling water it becomes easily powdered (Daguin, C. R. 20, 1667). The crystals are derived from the fundamental form of a rhombic pyramid; about thirty varieties are known. When held in the hand S emits a distinct odour, probably because of slight volatilisation; according to Berthelot (C. R. 100, 1326) S is wholly volatilisable at a temperature not much above the ordinary. Sis said to phosphoresce in air or O at 200° (Heumann, B. 16, 139). When S is rubbed it becomes strongly (negatively) electrified. S is a bad conductor of heat, and a very bad conductor of electricity; the conductivity varies slightly, 608 SULPHUR.

according as the light is or is not allowed to fall on the S (Monckman, Pr. 46, 136). Insoluble in water; easily sol. CS₂ (data at beginning of article); also sol. C₆H₆, CHCl₈, &c. Sol. warm conc. acetic acid (Liebermann, B. 10, 866); sol. liquid SO₂ (Sestini, Z. 1868. 718). As octahedral S changes to prismatic S near the m.p. of the former, it is evident that hot solutions of the former, in solvents which boil near to, or above, the m.p. of S must contain some prismatic S. Rhombic S slowly changes to monoclinic S when kept near its m.p. (v. p. 609, Allotropy of sulphur); the change is accompanied by absorption of heat (c. 650 g.-units for 32 grams S) and expansion of volume (v. Petersen, Z. P. C. 8, 601). According to Gernez (C. R. 83, 217), if a fair quantity of rhombic S is heated till melting begins on the surface, the interior portions change to microscopic monoclinic crystals, and the surface parts solidify again; this occurs only if the m.p. of monoclinic S (117.4°) is not exceeded. S melts to a clear yellow liquid; at c. 150° the liquid begins to darken and thicken; between 170° and 200° it is black, and so viscid that it does not pour out when the vessel containing it is inverted; at 330°-340° the liquid becomes thin, but it remains dark coloured until it boils at c. 444°. These changes are accompanied by very considerable changes in the electrical conductivity of S: the conductivity is almost constant from 270°-290°; it increases considerably up to 340°-350°, and then very rapidly up to the b.p. (v. Monckman, Pr. 46, 136). The expansion-coefficient, for each interval of 10°, decreases from 110° to 180°, and then increases gradually, but not regularly, to the b.p. (v. Moitessier, J. 1866. 27). Melted S may be cooled below 95° without solidifying (cf. Gernez, C. R. 97, 1298, 1366, 1433). For S.G., S.H., and other physical properties of S, v. beginning of this article.

B. Monoclinic sulphur, obtained by cooling molten S, forms transparent, yellowish brown needles; as obtained by crystallisation from solutions the crystals are nearly colourless. The crystals gradually become pale yellow and opaque, due to change into the rhombic form. The crystals are secondary forms of a monoclinic prism. This form of S is sol. CS2; rhombic S crystallises from this solution; it also dissolves in alcohol, C,H, CHCl, &c. On evaporation, crystals, both of rhombic and monoclinic S, separate (Maquenne, Bl. [2] 41, 238); monoclinic S may be recrystallised from alcoholic solution of (NH₄), S (for references v. p. 607, Preparation of monoclinic S). The change of monoclinic to rhombic S is hastened by scratching with a glass rod, or by covering with CS2; heated to 95.1° at 760 mm. pressure, monoclinic changes to rhombic S; the change is accompanied by appearance of heat and contraction of volume (v. p. 609, Allotropy of sulphur). For S.G. &c. of monoclinic S v. beginning of this article.

C. Soft soluble sulphur (v. Preparation of soft soluble sulphur, p. 607) is only partly sol. CS2; it contains both soluble and insoluble amorphous S; it is a very pale yellow, or nearly white, soft, amorphous, magma that gradually hardens; when heated it gives off H.S (Weber, A. 141, 432; Rose, P. 47, 166; Deville, Ph. C. 1848. 200; Fordos a. Gélis, Ph. C. 1854. 294).

Insoluble sulphur. It is doubtful whether more than one variety of this form of S exists, or whether substances described as varieties are not merely mixtures of ordinary in-

soluble S with impurities.

D. Plastic sulphur is a citron-yellow. soft, caoutchouc-like, amorphous solid, formed by suddenly cooling molten S (v. p. 607, Preparation of insoluble sulphur); as prepared from ordinary S it is dark brown, but Mitscherlich (J. pr. 67, 369) found that the dark colour is caused by the presence of small quantities of fatty substances (cf. Dietzenbacher, C. R. 56, 39). The S.G. of brown, plastic S is c. 1.91 to 1.96; but the substance is not homogeneous; it contains both soluble and insoluble S. Plastic S soon hardens and becomes yellow, more quickly when broken up and rubbed with a glass rod, or when immersed in turpentine, or when heated to c. 100°; heat is given out in this change (v. p. 609, Allotropy of sulphur). For determinations of the volumes occupied at different temperatures by plastic S v. Toepler (W. 47, 169).

E. Amorphous yellow sulphur is prepared from plastic S, or flowers of S, or the S obtained by decomposing thionates by acid, or H₂S by oxidisers &c. (v. p. 607, Preparation of insoluble sulphur), by washing with warm CS, till all soluble S is removed. This variety is an amorphous, buff-yellow powder; kept under CS2 it seems white; when dried and rubbed it forms a loose, flocculent powder. Insol. CS2; somewhat sol. CHCl₃, Et₂O, and alcohol, according to Deville (J. pr. 56, 359). Unchanged at the ordinary temperature, but slowly converted to crystalline soluble S at 100° (v. Berthelot, J. pr. 70, 941; 71, 364; Favre, J. Ph. [3) 24, 344; F. a. Silbermann, A. Ch. [3] 34, 447). S.G. 2.046 (v. data at beginning of this article). H.C. $[S,O^2] = 71,990$ (Petersen, Z. P. C. 8, 601). The accounts of this variety of S vary con-

siderably.

F. Black sulphur. When S mixed with a very little oil is thrown into a hot Pt dish, a black substance is obtained which has been looked on as a modification of S (Magnus, P. 92, 367; 99, 145; Dietzenbacher, P. 124, 644; Gross, B. B. 1879. 788; Jones, C. N. 41, 244; Keller, Bl. [2] 4, 346). Knapp (J. pr. [2] 88, 48; 43, 305) has shown that the black substance contains c. 55 p.c. S and c. 83 p.c. carbonaceous matter; K. thinks it is probably a modification of Sadhering to carbonised products of the oil. This substance is said to be non-volatile above the b.p. of S; it is a lustrous, amorphous, solid, insol. CS2, alcohol, Et2O, oils, H2SO4, &c.

8. Colloidal sulphur soluble in ter. This form of S is said to exist in water. Wackenroder's solution (the milky liquid formed by passing H₂S into SO₂Aq), and also in the soluby passing H₂S into 50₂A_{1/1}, and tion obtained by adding HClAq to Na₂S₂O₃Aq (v. p. 607, Preparation of colloidal sulphur). form of S has not yet been obtained free from impurities; it is a plastic, gummy, pale-yellow solid; it dissolves in water, forming a turbid liquid. By ppg. by NaCl, filtering, drying on bibulous paper, shaking with water, and repeating this treatment, Debus (C. J. 53, 284) obtained colloidal S that dissolved in water so as to form an almost clear, opalescent liquid, which became quite clear on warming, and turbid SULPHUR. 609

en cooling. This solution yielded ordinary S on addition of several salts; evaporation left a viscous, transparent residue; the S in solution did not diffuse through a porous membrane.

Other modifications of sulphur. Various experimenters have described forms of S different from those usually recognised, but there is much doubt as to whether any of these are really distinct varieties, or merely mixtures of known varieties (v. Maquenne, Bl. [2] 41, 238; Gernez, A. Ch. [6] B, 266; C. R. 100, 1326; Engel, C. R. 112, 866; Berthelot, C. R. 100, 1328; Brame, C. R. 101, 533, 639).

Allotropy of sulphur. The following table presents the best established allotropic forms of S:—

Insoluble in water.

Soluble in CS₂
Rhombic; S_a
Monoclinic; S_a
Amorphous, soft

Insoluble in CS₂
Plastic; S_a
Amorphous, yellow

Soluble in water. Colloidal; S.

The amorphous forms of S (soft, soluble; yellow, insoluble; and plastic, insoluble) and colloidal S have not been obtained pure; the soluble amorphous always contains insoluble, and the insoluble forms always contain soluble S; the colloidal contains S insoluble in water (cf. Magnus, J. pr. 70, 215; 72, 48; Weber, J. pr. 70, 354). It would probably be more accurate to make only three divisions of amorphous S: soluble in water, insol. water but sol. CS, insol. both water and CS. Berthelot (J. pr. 71, 364; 78, 244) holds that there are two main varieties of S: soluble in CS2, and insoluble in CS2. separated from compounds wherein it acts as the positive radicle, or part of the positive radicle (e.g. S₂Cl₂, SO₂Aq), is insoluble, according to B.; whereas when separated from compounds wherein it forms the negative radicle, or part of the negative radicle (e.g. H₂S, K₂S), S is soluble in CS₂. B. distinguishes the two varieties of S as electro-positive and electro-negative. Cloëz (J. pr. 74, 266; 78, 241) asserts that insoluble S is obtained from S.Cl. when this compound is rapidly shaken with water, but that soluble (crystalline) S separates from the same compound by the gradual action of moist air; and that insoluble S is obtained by electrolysing H.SAq provided the electrolysis is rapid (cf. Weber, P. 141, 432).

Any form of S changes gradually to the rhombic crystals (S_a); this change is hastened by raising the temperature within certain limits. Berthelot (J. pr. 71, 360) examined the amount of soluble (crystalline) S changed to insoluble at different temperatures; he found that rhombic S heated to 130°-140° and rapidly cooled was still wholly soluble in CS_a, that much insoluble S was formed by heating to c. 170°, and not much more at c. 280°. These results can be taken only as very roughly approximate, on account of difficulties in cooling, &c. (cf. Deville, J. pr. 56, 865; Frankenheim, J. pr. 54, 436). According to experiments of Gernez (C. B. 97, 1298, 1366, 1483; 100, 1343, 1382), Ruys (R. T. (C. 3, 1), and Reicher (J. 1885, 247), monoclinic S (S_B) changes to rhombic (S_a) in c. 12 days at - 36° Vol. IV.

to -15°, and in c. 30 minutes at 40°. Reicher (R. T. C. 2, 246; Z. K. 8, 593) says that the temperature of change of S, to S, is 95.6° at the ordinary pressure, and rises 05° for each increase of 1 atmosphere pressure; at 96.1° S. changes to S., and at 95.1° S. changes to S.. The change of any form of S to rhombic crystals is accompanied by the appearance of heat and contraction of volume; the data for the heats of combination of S., S., and amorphous insoluble S give c. 650 gram-units as the quantity of heat produced when 82 grams Se change to S., and c. 900 gram-units for the change of 32 grams insoluble amorphous S to S. (cf. Berthelot, C. R. 70, 941; and Mitscherlich, P. 88, 328). By keeping S molten for some time at 100°, and then inducing crystallisation by dropping in a crystal of S, either rhombic or monoclinic crystals can be obtained, according as the crystal dropped in is rhombic or monoclinic. rhombic S is melted and allowed to cool under ordinary conditions the solid contains both Sa and S_g; if the cooling is rapid, some amorphous S (both soluble and insoluble) is also produced; hence ordinary 'flowers of sulphur,' formed by rapidly cooling S vapour, contains both soluble and insoluble S, and generally also, when freshly prepared, both rhombic and monoclinic crystals. The crystals of S that separate from solutions in CS2 are rhombic; both forms of crystals separate from solutions in alcohol, CaH, CHCl, &c.; and monoclinic crystals separate from solutions in alcoholic (NH₄)₂S. Light brings about the change of soluble into insoluble S; if bright sunlight, or light from the electric arc, falls on molten S at c. 130°, a film of insoluble S is produced. Similarly, light concentrated by a lens and directed on to a cone. solution of S in CS, quickly causes the formation of a speck of insoluble S, which soon increases in size till the liquid becomes turbid (Berthelot, J. pr. 81, 396; Lallemand, C. R. 70, 182). No differences have been observed between the chemical behaviour of the soluble and insoluble varieties of S (v. Schmitz-Dumont, B. 25, 2659).

Atomic weight of sulphur. The at. w. of S has been determined (1) by converting AgCl into Ag,S (Berzelius, P. 65, 319 [1845]; Svanberg a. Struve, J. pr. 44, 820 [1848]); (2) by reducing Ag,SO, to Ag by H (Struve, A. 80, 203 [1851]; Stas, Stas R. 125 [1860]); (3) by direct synthesis of Ag,S (Dumas, A. Ch. [8] 55, 147 [1859]; Stas, Stas R. 53 [1860]); (4) by determining S.H. of S (Kopp, T. 1865, 71); (5) by determining V.D. of, and analysing, SH₃, SO₂, S₂Cl₂, &c. Molecular weight of sulphur. The V.D.

Molecular weight of sulphur. The V.D. determinations of Dumas (A. Ch. [2] 50, 170) gave c. 95 at 450°-500°, pointing to the molecular formula S_e; Bineau (C. R. 49, 799) found V.D. 39 from 714° to 743°, and 34 between 840° and 1160°, indicating the molecular weight S_e. The determinations by Deville a. Troost (C. R. 56, 891) at 860°-1040° confirmed the number 32 and the molecular weight S_e. Troost (C. R. 95, 30) got the number 42·5 at 665°, indicating a mol. w. of c. S_e (V.D. corresponding to S_e-48). In 1888 Biltz determined V.D. of S at intervals from 468° to 606°, and got results ranging from 118·2 at the lower temperature to 68·4 at the higher (S_e-112; S_e-64). The V.D. constantly decreased as temperature rose; the V.D. was constant only between 502° and 524°, but the

values were between those calculated for Se and S. Biltz concluded that the only molecules which exist as gas through any considerable range of temperature have the composition S. Ramsay (Z. P. C. 3, 67 [1889]) thought that Biltz's results did not negative the existence of gaseous molecules more complex than S. Schall (B. 23, 1701 [1890]) got numbers for V.D. varying from 115 to 122, at 573° and pressures from 10-20 mm., in an atmosphere of N or CO₂; S. concluded that molecules S₀ probably exist in the vapour of S. Riecke (Z. P. C. 6, 430 [1890]) regarded the decrease of V.D. with increasing temperature as a dissociation of S_s to S_s and S_z; by making certain assumptions, his calculated results agreed well with the experimental numbers of Biltz. In 1888 Paterno a. Nasini (B. 21, 2153) determined the depression in the freezingpoint of benzene caused by dissolving S therein; their results indicated S_s as the molecular formula of S in solution in benzene. By determining the increase in the boiling-point of CS, produced by dissolving S in that solvent, Beckmann (Z. P. C. 5, 76 [1889]) got values for the mol. w. of S varying from 245 to 280 ($S_s = 256$). Sakurai also used this method in 1892 (C. J. 61, 989); his values for mol. w. of S in CS, varied from 252.3 to 254.9. From observations of the effect of dissolving S in naphthalene on the f.p. of the solvent, Hertz (Z. P. C. 6, 358 [1890]) concluded that the mol. w. of S in solution in naphthalene is $256 = S_s$. Helff (Z. P. C. 12, 196) in 1893 also got S, for mol. w. of S in CS, and in molten P. These results leave little doubt that molecules S₈ exist in solutions of S in CS₂ and in naphthalene, and S₆ in solutions in benzene; that molecules of greater complexity than Saprobably as complex as S_s—exist in S vapour, but these are stable only through a small range of temperature, and gradually dissociate as temperature rises until all the molecules existing above c. 700° have the composition S₂. Biltz a. Meyer (B. 22, 725 [1889]) have shown that the V.D. of S corresponds with the mol. w. S, up to 1600°-1700°.

Reactions .- 1. Reacts with water at 100° to form H.S. Cross a. Higgin (C. J. 35, 249; B. 16, 1195) found that the solution contained thionic acids; they supposed that these were produced by reactions between the H2S and SO. formed by the action of the S on the water (cf. Payen, J. Ph. 8, 871; Mulder, J. 1858. 84; Meyer, C. R. 74, 195; Gélis, C. R. 56, 1004). Senderens (Bl. [8] 6, 800) regarded the production of H₂S as due to interactions between the S and the glass vessels employed; C. a. H. (C. J. 85, 252) found H_2S was produced when dilute H.SO.Aq was used in place of water, and they concluded that the alkaline constituents of the glass were without influence. Becquerel (C. R. 56, 287) says that when water containing S in suspension is electrolysed, H2SO4 forms at the positive, and H2S at the negative, electrode (cf. Colson, Bl. [2] 84, 66). When S is moistened with water and let stand in the air, H,SO,Aq is said to be produced (Polacci, C. C. 1884. 484; Böhm, M. 8, 224).—2. S is oxidised to H₂SO, Aq by heating with nitric acid, aqua regia, or potassium chlorate and hydrochloric acid.—8. Conc. hot sulphuric acid is reduced, giving off SO_s.—4. Conc. hydriodic acid gives H_zS and I (I decomposes dilute H.SAq to HIAq + S).-5. Chlorosulphonic acid when heated with S produces S.Cl., SO, and HCl (Heumann a. Köchlin, B. 15, 416).-6. S dissolves in boiling alkali solutions, also in molten alkalis and alkaline carbonates, to form mixtures of polysulphides and thiosulphates. Boiling ammonia solution forms polysulphides; heated in a sealed tube above 100° some (NH₄)₂S₂O₄Aq is produced (v. Brunner, D. P. J. 150, 371).—7. Sulphur trioxide dissolves S, forming a blue solution of S₂O₂ (v. Sulphur oxides).—8. Carbon dioxide reacts with boiling S to form COS (Cossa, B. 1, 117; Berthelot, Bl. [2] 40, 364). The same compound is produced by volatilising S, by an electric current, in carbon monoxide (Chevrier, C. R. 69, 56).—9. PSCl, is formed by the interaction of S and phosphorus trichloride at 130° (Henry, Bl. 13, 495). Phosphorus pentachloride forms PCl₂ and S₂Cl₂ (Goldschmidt, C. C. 1881. 489).— 10. S decomposes many sulphates and carbonates at high temperatures, forming sulphides and SO_2 , or CO_2 (v. Sestini, Bl. [2] 34, 490; Berthelot, Bl. [2] 40, 864).—11. Solutions of salts are very often reduced by boiling with S, sulphides being generally ppd. (v. Vortmann a. Padberg, B. 22, 2642; and, more fully, Senderens, Bl. [3] 6, 800; 7, 511).

Combinations.—1. Sulphur combines directly with most of the elements. The binary compounds of S—with the exception of those with Br, Cl, F, I, or O, and also compounds of S with more than one other element (except these be some of the five just mentioned)—are described under the least negative of the component elements; e.g. compounds of S with P and Cl are described (as sulphochlorides) under Phosphorus. For the conditions of formation of the binary compounds of S reference must be made to the various elements, except in the cases of bromidé, chloride, fluoride, iodides, and oxides of S, which are described in this article in their proper (alphabetical) places.—2. Chlorine monoxide Cl₂O is said to combine directly with S (suspended in S₂Cl₂), at —12°, to form SOCl₂ (Wurtz,

C. R. 62, 460).

Sulphur, acids of. Reference should be made to Hydrogen sulphide, vol. ii. p. 725; Sulphur oxyacids, this vol. p. 619; Sulphur oxyacids, nitrogen derivatives of, this vol. p. 619; Sulphonid acids and derivatives, this vol. p. 599; Thio-carbonic acid, vol. i. p. 703; Sulphonid acids, containing S and P, and S, P and O, v. Phosphorus sulphides, this vol. p. 145 (beginning of article); also Phosphorous sulphides, Reactions 7 and 8 (p. 146), Phosphoric sulphide, Reaction 8, and also end of that article (p. 147).

Reaction 8, and also end of that article (p. 147).

Sulphur, bromides of. It is very doubtful whether any definite compound of S and Br exists. Powdered S dissolves in Br with evolution of a little heat; the deep ruby-red liquid begins to distil over at c. 60° and the thermometer rises steadily to c. 190°, when it ascends slowly to c. 200°, after which it steadily rises till S remains; the fraction coming over at 190°-200° agrees fairly in composition with the formula S₂Br₂ (Pattison Muir, C. J. 28, 845). M. found that by passing CO₂ through a solution of S in excess of Br for some hours, at 15°, 50°, and 90°, the residual liquid had nearly the com-

position S₂Br₂; Hannay, however, by passing air through such a liquid for 400-500 hours found that the whole of the Br was removed and that pure S remained (C. J. 33, 284; 35, 16). It is, however, possible that at the temperature of this experiment, c. 15°, dissociation of S.Br. was induced by the stream of air. H. also found that the vapour given off by the supposed S₂Br₂, even at 0°, showed the absorption-spectrum of Br. Spring a. Lecrenier (Bl. [2] 45, 867) examined the reaction of the supposed S₂Br₂ with K₂SO₃. S. had before shown that S2Cl, and K2SO, produce KCl and K2S4O6; but Cl and S acting as free elements produce K₂SO₄, KCl, K₂S₂O₃ and SO₂. S₂Br₂ ought then to produce KBr and K₂S₄O₄, whereas S and Br would form K₂SO₄, KBr, K₂S₂O₃, and SO₄; by determining the quantities of the products S. a. L. calculated that the supposed S.Br. contains c. 27 p.c. of the elements uncombined. Various reactions of the supposed S2Br2 are described in the memoirs already referred to (v. also Rose, p. 44, 327; Michaelis, J. Z. 6, 297; Ogier, C. R. 92, 922).

Sulphur, chlorides of. Three compounds of S and Cl are known: S_2Cl_2 , SCl_2 , and SCl_4 . These compounds are liquids; S_2Cl_2 boils unchanged at c. 138°, SCl_2 begins to decompose to S_2Cl_2 and Cl at c. 10°, and the decomposition of SCl_4 to SCl_2 and Cl begins at c. -20° .

SULPHUR MONOCHLORIDE S₂Cl₂. (Sulphur protochloride, Sulphur sulphochloride, Sulphur sulphochloride, Sulphothionyl chloride, Thio-thionyl chloride.) Mol. w. 134.7. Boils at 138·12°; S.G. ⁶⁰/₆₂ 1·70941, S.G. at b.p. 1·49201 (Thorpe, C. J. 37, 356). S.V. 90·28 (T., l.c., p. 372). V.D. 68·4 (Dumas, A. Ch. [2] 49, 204; Marchand, J. pr. 22, 507). μ_{Ha} = 1·64449 (Haagen, P. 131, 117). H.F. [S²,Cl²] (from S_a) = 14,257 (Th. 2, 810); 17,600 (Ogier, C. R. 92, 922).

Formation.—1. By the interaction of S and Cl.—2. By distilling S with 9 parts SnCl₂ or 8.5 parts HgCl₂.—3. By reacting on PCl₃ with S (Goldschmidt, C. C. 1881. 489).—4. By decomposing PSCl₂ by Cl or by heating to redness.—5. By the interaction of P₂S₃ and SOCl₂, the other product being P₂O₅ (Carius, A. 106, 331).—6. By the action of CS₂ on ICl₂ (Weber, P. 128, 459).

Preparation.—A stream of dry Cl is passed into a flask, or retort, containing S (flowers or sticks) until most, but not quite all, S is dissolved to a yellowish-red liquid, the vessel being gently warmed. The liquid is repeatedly distilled until it boils constantly at 138°-139°. If the passage of Cl is continued until the S is completely dissolved, the liquid must be distilled from a little S (to decompose SCl₂ quickly).

Properties.—A yellowish-red, oily liquid, with a very characteristic, rather disagreeable, odour and an acid taste; the vapour affects the mucous membranes rapidly, causing tears, and affecting the breathing. Does not become visoid at -75° (Haase, B. 26, 1052). Fumes in air. Dissolves in CS₂ and C₂H₂, also in alcohol and ether with decomposition. S₂Cl₂ dissolves large quantities of S; solution of S₂Cl₂, containing free S, in CS₂ is used for vulcanising (v. DICTIONARY OF APPLIED CHEMISTEY, vol. iii. p. 706).

Reactions and Combinations.—1. S.Cl. sinks in water, and then decomposes, giving HClAq,

SO₂Aq, and S, with a little H₂SAq, H₂S₂O₂Aq, and some of the thionic acids .- 2. Hydrogen sulphide produces HCl and S; hydrogen iodide reacts at the ordinary temperature to form HCl, I, S.I., and then H.S (Hautefeuille, Bl. [2] 7, 198).—3. By passing S₂Cl₂ vapour, mixed with air or oxygen, through a red-hot tube, SO₂, SO₂, and Cl are formed (Donny a. Mareska, C. R. 20, 817; Spring a. Lecrenier, Bl. [2] 45, 867). 4. Heating with phosphorus produces PCl_s, PSCl_s, and S if a little P is used; or PCl_s, P sulphides, and red P if much P is used (Wöhler a. Hiller, A. 93, 274; Chevrier, C. R. 63, 1003).—5. Heated with many metals, S.Cl. gives metallic chlorides and S; the more volatile the metallic chloride the more readily does the the metallic chloride the more readily does the reaction occur (v. Chevrier, C. R. 64, 802).—6. S₂Cl₂ combines with chlorine, forming SCl₂ and SCl₄ (q.v., infra). Br and I are absorbed by S₂Cl₂; according to Evans a. Ramsay (C. J. 45,65) no compound is formed with Br.—7. With the second compound of compound second sec oxides of arsenic and antimony S₂Cl₂ forms AsCl₃ and SbCl₃, giving off SO₂ and separating S; selenion dioxide produces Se, Cl, and SO,; oxides of phosphorus, boron, and silicon do not react (Prinz, A. 223, 355).—8. S2Cl, absorbs sulphur trioxide, below 0°, forming a brownish-yellow liquid, from which excess of SO, crystallises out; a little above 6° SO₂ is given off, rapidly at 10°; S₂O₅Cl₂ is formed on distillation (Rose, P. 44, 291).-9. By passing vapour of S.Cl. along with the vapours from sulphonic acid through a hot tube, SO₂, H₂S, S, HCl, and Cl are produced (Brault a. Poggiale, J. Ph. 21, 140).—10. Heating S2Cl2 with sulphates generally produces SO2Cl2, chlorides, S, and SO,; heated with sulphuryl chloride to 250°, S,OCl, is formed (v. Schneider, J. pr. 104, 83; [2] 32, 18; Demarcay, C. R. 92, 726).—11. S,Cl, combines with ammonia gas to form S₂Cl₂4NH₃, stable in air, sol. alcohol, decomposed by water giving NH₄ClAq(NH₄)₂S₂O₃Aq, and S (Brault a. Poggiale, J. Ph. 21, 140).—12. Reacts with organic compounds containing oxygen, generally forming Cl compounds and small quantities of S compounds, and also HCl, SO2, and S.-13. Combines with unsaturated hydrocarbons; e.g. with C.H. forms C2H4S2Cl2

Sulphur dichloride SCl_x Mol. w. (in solution in benzene or acetic acid) $102\cdot72$. S.G. $1\cdot6482$ at $15\cdot4^\circ$ (Costa, G.20, 367). $\mu_{Ha}=1\cdot5716$ (Costa, l.c.). No constant b.p. $(v.\ infra)$. This compound was prepared by H. Rose in $1831\ (P.21, 431; 27, 107)$; the experiments of Dumas $(A.\ Ch. [2] 49, 204)$ and Marchand $(J.\ pr. 22507)$ led to its recognition as a definite compound $(v.\ also$ Carius, $A.\ 106, 291)$; Hünner a. Guerout $(Z.\ 1870.\ 455)$ isolated the compound satisfactorily, and their results were confirmed by Thorpe a. Dalzell, in $1871\ (C.\ N.\ 24, 159)$. Costa, in 1890° (G. 20, 867), determined the mol. w. of the compound in solution.

Preparation.—Dry Cl is led into S₂Cl₂, surrounded by a freezing mixture, until the liquid becomes deep red; excess of Cl is removed by a stream of dry CO₂. The temperature must not rise above 6°; the vessel containing the S₂Cl₂ should be kept filled with Cl continually, and should not be exposed to direct sunlight (H. a. G., Lc.; C., Lc.; T. a. D., Lc.; also Soubeiran A. Ch. [2] 67, 64; Michaelis, A. 170, 1).

Properties. — A thin, dark, brownish-red, tiquid, with smell and taste like S₂Cl₂; does not solidify at -30°; gives off Cl when exposed to sunlight, and should, therefore, be kept in a sealed tube in the dark. Easily decomposed by heat to S₂Cl₂ and Cl; Michaelis (B. 6, 995) gives a table showing amount of dissociation from 20° to 180°; the temperature whereat 50 p.c. of the SCl₂ is dissociated is c. 86°. SCl₂ reddens dry litmus paper. The depression of the freezing-point of benzene or glacial scetic acid by solution therein of SCl₂, shows that the mol. w. agrees with the formula SCl₂ (Costa, G. 20, 367).

Reactions and Combinations.—1. Heat, or direct sunlight, causes dissociation to S2Cl2 and Cl (v. supra, Properties).—2. Water produces HClAq and H₂S₂O₂Aq.—3. Oxidised violently by nitricacid to HClAq and H2SO, Aq. -4. Ammonia solution produces NH₄ClAq, N, and S; ammonia gas combines to form SCl₂2NH₃ and SCl₂4NH₄, soluble, unchanged, in absolute alcohol and ether (Soubeiran, A. Ch. [2] 67, 74).-5. Thionyl chloride (SOCl.), when heated with SOl, forms S2OCl4 (Ogier, Bl. [2], 37, 293).—6. Alkali sulphites produce chlorides and trithionates (Spring a. Lecrenier, Bl. [2] 45, 867).—7. Some metals, e.g. Cu, Fe, K, react with SCl₂, with production of heat and light, and formation of chlorides and sulphides.—8. Combines with arsenious chloride, forming AsCl. SCl. according to H. Rose (P. 27. 107).—9. SCl₂ reacts with oxygenised carbon compounds similarly to, but more readily than, S2Cl2; it combines with several unsaturated hydrocarbons (v. Heintz, A. 100, 870; Guthrie, C. J. 12, 109).

SULPHUE TETRACHLORIDE SCI,. This compound exists only at temperatures under -20°. The molecular weight is probably 178.46 (SCI,).

Preparation.—S₂Cl₂ is cooled to below -20°

Preparation.— \tilde{S}_2Cl_2 is cooled to below $\sim 20^\circ$ (c. -22°) and a slow stream of dry Cl is passed in until absorption of Cl ceases. Michaelis a. Schifferdecker (B. 6, 993) found that 67.5 g. S₂Cl₂, kept at -20° to -22° , absorbed 106 g. Cl in 10 hours, that absorption of Cl then ceased, and that the liquid had the composition SCl_4 .

Properties.—A mobile, yellowish-brown liquid; when removed from the freezing mixture used in the preparation, the liquid gives off Cl, and boils with an absorption of much heat. M. a. S. give the following data showing the dissociation of SCl, to SCl, and Cl:—

Temp.	P.o. SCL	P.c. SCL
- 22°	100.0	0.0
- 15	41.95	58·0 5
- 10	27.62	72.38
- 7	21.97	78.03
- 2	11.93	88.07
+0.7	8.87	91.13
+6.2	2.43	97.57

Reactions.—1. Water causes rapid decomposition to HClAq and SO₂Aq, with more or less S according to the mass and temperature of the water.—2. With sulphur trioxide, in the ratio SOl₄:SO₂ the products are SOCl₂ and S₂O₃Cl₃ in the ratio SOl₄:2SO₃ the products are SOCl₂, SO₂ and Cl (M. a. S., Lo.; also B. 5, 924). Sulphur dioxide does not react.—8. Chlorosulphonic acid reacts at a low temperature to form S₂O₃Cl₄ and HCl (M. a. S., B. 6, 996).—4. With some hydroxylic organic compounds, e.g. C₄H₄.OH and

C.H. CO.OH, OH is replaced by Cl and SOCl, is formed.

Combinations.—1. Compounds of SCl4 with

several metallic chlorides are produced by warming the chlorides with S₂Cl₂ and passing in Cl; the compounds 2AlCl₂.SCl₄, AuCl₂.SCl₄, and 2TiCl₄.SCl₄, are formed thus (Weber, P. 104, 421; Rose, P. 42, 517; Casselmann, A. 83, 267). The compound with AlCl₂ may be distilled in a closed \$\simes\$ shaped tube; the other two compounds give off Cl when heated. The compounds SbCl₂.3SCl₄ and SnCl₄.2SCl₄ are produced by passing Cl over SnS₂ and Sb₂S₃ respectively (R., Lc.; C., Lc.).—2. With iodine trichloride forms ICl₂.SCl₄ (Weber, P. 128, 459; cf. Jaillard, A. Ch. [3] 59, 454, who gives the composition SCl₂.2ICl₂). Prepared by passing Cl into I dissolved in CS; till the liquid is wine-red, coelling, and drying the red-yellow, prismatic, very deliquescent, crystals, in a tube in a stream of dry Cl. Also formed by passing Cl into a mixture of 2 pts. S and 1 pt. I. Melts to a brown liquid when heated in a closed tube; heated in an open tube forms Cl, ICl, and S₂Cl₂; decomposed by

Sulphur, chloro-iodide of, SCl,I(= SCl,ICl₂); v. supra, Sulphur tetrachloride, Combinations, No. 2.

water or dilute HNO,Aq.

Sulphur, chloronitride of, S₂CIN; v. Nitrogen sulphochloride, vol. iii. p. 571.

Sulphur, cyanides of, v. Cyanogen, sulphides, vol. ii. p. 358.

Sulphur, fluoride of. According to Gore (Pr. 20, 70 [1871]) the heavy colourless gas produced by fusing AgF with S is a fluoride of S. This gas is not liquefied at 0°, at the ordinary pressure; it fumes in the air, and attacks glass; no analyses are given by G.

Sulphur, haloid compounds of. The most stable of these compounds belong to the forms S_2X_2 ; where X = Cl, Br, or I. The chloride S_2Cl_2 , may be distilled unchanged; the bromide, S_2Br_3 , is exceedingly easily decomposed if, indeed, it has been isolated; the iodide, S_2I_2 , has been very slightly examined. The compounds SCl_2 and SCl_4 also exists; the former at temperatures below c. 10°, and the latter below c. -20°. The compound Sl_4 perhaps exists; it gives up I at the ordinary temperature. The only haloid compound of S that has been gasified unchanged is S_1Cl_2 .

Sulphur, hydrides of, SH₂ and (?) S₂H₂; v. Hydrogen sulphides, vol. ii. p. 725.

Sulphur, nitride of, SN; v. Nitrogen sulphide, vol. iii. p. 570.

Sulphur, iodides of. S and I probably combine when heated together, but no definite compound has been isolated by this method (of. Gay-Lussac, G. A. 48, 872; H. Rose, P. 27, 115; Lamers, J. pr. 84, 849; Sestini, J. 1863. 153; Wheeler a. Lüdeking, Fr. 26, 602).

SULFRUE MONO-RODIDE S.I. Guthrie (C. J. 14, 57) obtained this compound by placing 2½ parts Etl and 1 part S₂Cl₂ in a scaled tube, opening after 12 hours, evaporating EtCl by warming with the hand and removing excess of Etl by gently warming; the S.I. crystallised out in lustrous crystals resembling I. (One analysis is given, but no full description of properties.)

Ogier (C. R. 92, 922) gives H.F. [S², I²] (from

gases) = 10,800.

SULPHUR HEXA-IODIDE SI. This compound is said to be formed by slowly evaporating, at a low temperature, a solution of I with slight excess of S in CS₂. Grey-black crystals; resembling, and isomorphous with, I; I evaporates on exposure to air and eventually only S remains; alsohol, conc. KOHAq, or KIAq, also withdraws all I (G. vom Rath, P. 110, 116; Lamers, J. pr. 84, 349). It is very doubtful whether the crystals examined by vom Rath were a compound or only I retaining a little S.

By passing H₂S into a dilute solution of KCl.ICl₂, Lamers (l.c.) obtained an orange-red pp. which dried in an exsiccator to a brown amorphous mass containing S and I in the

ratio S,I2

By subliming SnS₂ with I, Schneider (J. 1860. 186) obtained a compound to which he gave the

composition SnSI2.SI2.

Sulphur, iodochloride of, SICl, (= SCl, ICl₂) v. Sulphur tetrachloride, Combinations No. 2.

Sulphur, oxides of. Four compounds have been isolated; S₁O₂, SO₂, SO₃, and S₂O₇. S and O combine directly to form SO₂ and SO₂; S dissolves in SO₃ to form S₂O₃; S₂O₇ is formed by the action of an electric discharge on a mixture

of SO₂ and O, or SO₃ and O.

SULPHUE SESQUIOXIDE S.O. (Hyposulphurous anhydride). The production of a blue substance, on distilling fuming H₂SO₄ and S, was noticed by Buchholz in 1804 (Gehlen's J. 3, 7); Vogel, in 1812, obtained the blue body by bringing together S and SO₄ (S. 4, 121); the substance was examined by Wach (S. 50, 1), Berzelius (Lehrbuch [5th edit.] 1, 485), and Stein (J. pr. [2] 6, 172). Weber, in 1875, isolated the compound and found its composition to be S₂O₅ (P. 156, 531).

S.O. is prepared by adding well-dried flowers of sulphur, little by little, to SO, quite free from H.SO., more S being added when the previous quantity has all combined. The process is conveniently conducted in a test tube, covered with a small porcelain crucible to exclude moisture, and plunged into water cooled to c. 12°; the S disappears in the SO, and blue drops are formed; the tube should be tilted so that the blue drops come into contact with the sides of the tube where they form a thin crust (if the drops solidify as they float in the SO, the solid S,O, retains 80, from which it cannot be freed). When c. 1g. S has been added, excess of SO, (which should be colourless) is poured off, the last traces being removed by gently and carefully warming not above 35°; the solid S₂O₈ is scraped out of the tube by a glass rod.

S₂O₂ is a blue-green solid, consisting of a mass of microscopic crystals. Insoluble pure SO₂; if a little H₂SO₂ is present the S₂O₂ dissolves to a deep-blue liquid. Melts with decomposition; decomposes slowly in dry air, quickly on heating, to SO₂ and S; deliquesces in moist air to a brown liquid from which S soon separates; reacts violently with water, giving H₂SO₂Aq, SO₂Aq, H₂SO₂Aq, polythionic soids, and S; decomposed also by alcohol and ether, with separation of S, also by ordinary H₂SO₃.

S₂O₂ may be regarded as the anhydride of H₂S₂O₄; although the acid has not been ob-

tained from the oxide, nor the oxide from the

For the compounds SSeO, and STeO, v. Selenion thio-oxide (p. 441) and Tellurium thio-oxide (under Tellurium oxides).

SULPHUE DIOXIDE SO₃. (Sulphurous oxide. Sulphurous anhydrids. Thionyl oxide.) Mol. w. 63·9. Melts at c. -76° (Faraday, T. 1846. 155); c. -80° (Mitchell, A. 87, 356). Boils at c. -10° (Bunsen, P. 46, 97; Andréeff, J. 1859; Pictet, C. C. 1877. 81). S.G. (liquid) 1·4911 at -20·5°, 1·4384 at -2·08°, 1·4252 at +2·8°, 1·3769 at 20°, 1·3258 at 30°, 1·2872 at 52°, 1·1845 at 82·4°, 1·1041 at 102·4°, ·956 at 180·3°, ·7317 at 151·75°, ·52 at 156° (Cailletet a. Mathias, C. R. 104, 1563). S.G. (gas at 0° and 760 mm.) 2·2639 (Leduc, C. R. 117, 219). V.D. 32·28 (Gay-Lussac, Berzelius, Buff; Biltz, B. 21, 2769). S.H. (equal wt. water = 1) ·1544; (equal volume air = 1) ·3414 (Regnault, J. 1863. 84). Ratio of S.H. constant pressure to S.H. constant volume =1·2562 (Müller, W. 18, 94). C.E. ·00423 for 1° between 0° and 10°, ·004005 10° to 20°, ·003846 at 50°, ·003757 at 100°, ·003718 at 150°, ·003695 at 200°, ·003685 at 250° (Amagat, C. R. 73, 183; cf. C. R. 68, 1170). Leduc (C. R. 117, 219) gives C.E. between 0° and 20° as ·00396. For C.E. of liquid SO₂ v. Drion (A. Ch. [3] 56, 5) and Andréeff (A. 110, 1).

ratio $\frac{PV}{P_1V_1} > 1$; Amagat (l.c.) gives these values,

1.0185 at 15°, 1.0110 at 50°, 1.0054 at 100°, 1.0032 at 150°, 1.0021 at 200°, 1.0016 at 250°. H.F. $[8,0^2]=71,070$ from solid rhombic 8; 71,720 from solid monoclinic 8 (Th. 2, 247). Heat of liquefaction of gaseous $S_2=5644$ (Favre, A. Ch. [5] 1, 209). Critical temperature = 156° (Cailletet a. Mathias, C. R. 104, 1563); 155.4° (Sajotscheffski, P. B. 1879. 741); 157°–161° (Ladenburg, B. 11, 821). Heat of vaporisation of liquid SO_2 , at $0^\circ=91\cdot2$, at $60^\circ=69$ (C. a. M., l.c.; cf. Chappuis, C. R. 104, 897). Regnault (C. R. 50, 1063) gives vapour pressures as follows:

Temp.	Vap. pressure	Temp.	Vap. pressure
- 25°	873.79 mm. Hg	80°	3131.80 mm. Hg
-20	479.46 ,,	40	4670.23 ,,
-10	762.49 ,,	50	6220.01 ,,
0	1165.06 "	60	8123.80 "
+10	1719.55 ,,	65	9221.40 "
90	2462-05		

Vapour-pressure at critical temp. = 78·9 atmos. (Sajotschefiski, P.B. 1879. 741). S. in water between 0° and 20° (gaseous SO₂) = 79·789 $-2\cdot6077t+\cdot029849t^2$; 1 vol. of the saturated SO₂Aq contains $68\cdot861-1\cdot87025t+01225t^2$ vols. SO₂; S. (water) between 21° and $40^\circ-75\cdot182-2\cdot1716t+01903t^3$; 1 vol. of the saturated SO₂Aq contains $60\cdot952-1\cdot88896t+00726t^2$ vols. SO₂ (Bunsen a. Schönfeldt, A. 95, 2). S. in alcohol, at 760 mm. = $328\cdot62-16\cdot95t+3119t^3$; S.G. of solution = $1\cdot11937-014091t+000257t^2$ (Carius, A. 94, 148). Sims (C.J. 14, 1) gives the table on the next page, which presents both the weight of SO₂ in grams and the volume of gaseous SO₂ in c.c. dissolved by unit weight of water at the normal pressure and different temperatures:—

	Grams SO.	c.c. 80 _a
Temp.	absorbed by	
8°	·168	58· 7
12	·142	49.9
16	·121	42.2
20	·104	86.4
24	•092	82.3
28	•083	28.9
82	.078	25.7
36	.065	22.8
40	.058	20.4
44	.053	18.4
48	.047	16.4
50	•045	15.6

The quantity of SO₂ dissolved by a given weight of water varies with pressure, and only at c. 40° and upwards is the quantity proportional to the partial pressure of the gas (v. Sims, C. J. 14, 1). For S.G. and composition from 5 to 10 p.c. of SO₂Aq v. Anthon (C. C. 1860.744); also Giles a. Schearer (S. C. I. 4, 303). [SO²,Aq] = 7,700 (gaseous SO₂) (Th. 2, 249); 1500 (liquid SO₂) (Berthelot, C. R. 96, 142, 208; Chappuis, A. Ch. [2] 19, 21). H₂SO₄Aq absorbs the following quantities of gaseous SO₂ (Kolb, D. P. J. 209, 270; cf. Dunn, C. N. 43, 121; 45, 270):—

s.G. H.80.Aq	Kilos. SO, per kilo. acid	Litres SO, per litre acid
1.841	•009	5.8
1.839	.014	8.9
1.540	.021	11.2
1.407	.032	15·9
1.227	•068	29.7

SO₂ is absorbed by charcoal; 1 c.c. charcoal = 1.57 g. was found by Favre (A. Ch. [5] 1, 209) to absorb 165 c.c. SO₂. Camphor absorbs c. 308 simes its vol., glacial acetic acid c. 318 times, SO₂Cl₂ c. 187 times, its volume of SO₂ (Schulze, J. pr. [2] 24, 168; Bineau, A. Ch. [3] 34, 326). $\mu_{Na} = 1.35$ at 15° (Bleekrode, Pr. 87, 359). For surface tension of liquid SO₂ v. Clark (C. N. 38, 294; 40, 8). For absorption-spectrum v. Liveing a. Dewar (C. N. 47, 121).

Occurrence.—In volcanic gases, and in river and spring water near volcanoes (v. Ricciardi, G.

18, 38).

Formation.—1. By burning S in O. Baker (Pr. 45, 1; C. J. 47, 349) found that the combustion of S in O is retarded the drier the O is; using very carefully dried O there was no flame. According to Hempel (B. 23, 1456) c. 2 p.c. of the gas formed when S is burnt in O at the ordinary pressure is SO₂, 98 p.c. being SO₂; at a pressure of 40°-50° atmos. c. half the S is burnt to SO₂ and half to SO₂; at higher pressures c. 70 atmos. the proportion of SO₄ increases.—2. By heating S with oxide of Pb, Mn, Hg, Zn, &c.—3. By heating various sulphates with S, also SO₂ with S, or conc. H₂SO₄ with C, Cu, Hg, or S; by heating Na₂SO₄ with charcoal and SiO₂.—4. By the decomposition, by acids, of alkali thiosulphate or polythionate solutions.—5. By burning H₂S, CS₂, and various organic compounds containing S.—6. By decomposing conc. H₂SO₄ by heat.—7. By the action of electric sparks on a mixture of sulphur with CO₂, NO, or N₂O (Chevrier, C. R. 69, 136).

Preparation. -1. Pure H2SO, mixed with half to two-fifths its volume of water, is heated with pure Cu; the SO, that comes off is passed through water in a large flask, and then through a couple of bottles filled with pumice, broken into small pieces and moistened; the pumice should be twice moistened with pure H,SO, and heated to redness before being used, to remove chlorides and fluorides (Stas, Chem. Proport. 115). The SO, may be dried by passing it through pure cone. H₂SO₄, and then over CaCl₂—2. A mixture of 3 parts pure CuO and 1 part 8 in powder is heated in a tube of hard glass; the anterior part of the tube contains CuO only, to oxidise 8 that may sublime (Marchand, P. 42, 144).—3. Neumann (B. 20, 1584) obtains a regular stream of approximately pure SO₂ by placing a mixture of three parts CaSO₃ and 1 part CaSO₄, made into small cubes, in a Kipp's apparatus, and decomposing by ordinary oil of vitriol. The acid should be allowed to come into contact. with sufficient cubes, and no more, to give the stream of gas that is required. Using c. 500 g. of the cubes, N. obtained a constant stream of SO, for 80 hours. The cubes are made by mixing the CaSO, and CaSO, moistening with water so that a semi-solid mass is obtained that can be worked with the fingers with some difficulty, pressing this in an iron mortar, hammering it into an iron frame c. 10-12 mm. in height, covering with oilcloth and pressing strongly, cutting into cubes while still in the frame, removing and drying at c. 20° (v. Winkler, B. 20,

184).

Properties.—A colourless gas, at ordinary temperatures and pressures, with the strong penetrating, choking odour of burning S; causes coughing and blood-spitting if inhaled in any quantity. Incombustible in air, and extinguishes flame; when mixed with O and passed over heated spongy Pt it is oxidised to SO₃. Dry SO, is said not to redden litmus paper (Wilson, C. J. 1, 332). Bleaches many vegetable colours; the colours are generally restored by alkalis. Bad conductor of electricity (v. Bleekrode, P. M. [5] 5, 375, 439). Is not decomposed by passing through a red-hot tube. Condensed at $c.-10^\circ$ under the ordinary pressure -by passing the dry gas into a tube surrounded by a mixture of snow and salt; may also be condensed at the ordinary temperature by evolving the gas (by warming charcoal saturated with SO₂, or a mixture of 1 part S and 5 parts SO, in one limb of a closed tube, the other limb being placed in cold water. Convenient apparatus for liquefying SO, is described by Wöhler (A. 137, 871) and Hofmann

(B. 2, 262).

Liquid SO₂ is a colourless, mobile liquid. Exposed to air it evaporates very rapidly, with disappearance of much heat; an air thermometer placed in liquid SO₂ at 10°, in an open vessel, falls to -57°. Liquid SO₂ dissolves P, S, I, Br, and many gums. It is miscible in all proportions with liquid SO₂; partially miscible with CS₃, Et₂O, CHCl₃, and C₅H₄ on warming; but immiscible with conc. H₂SO₄ (Sestini, Bl. [2] 10, 226). Liquid SO₂ is a non-conductor of

By evaporating liquid SO, rapidly in an airpump, part of it solidifies to a white, woolly

electricity (Magnus, P. 104, 538).

mass; the liquid may also be solidified by placing it in a mixture of solid CO₂ and ether.

Reactions and Combinations.—1. 80, is not decomposed by passing through a red-hot tube. But by heating to c. 1200° it is changed to SO. and S $(3SO_2 = 2SO_2 + S)$; the action stops when the SO, attains a certain vapour pressure, but if the SO, is removed (by conc. H₂SO₄) the whole of the SO₂ is decomposed (Deville, C. R. 60, 317). Induction sparks effect the decomposition of SO₂ to SO₃ and S (v. Buff a. Hofmann, A. 113, 129); light acts similarly (Morren, C. R. 69, 397).—2. H₂O and S are produced by passing SO₂ mixed with hydrogen through a redhot tube.—3. Chlorine combines with SO₂, in sunlight, to form SO₂Cl₂ (v. SULPHURYL CHLORIDE). SO₂Cl₂ is also formed by passing SO₂ and Cl into glacial acetic acid, even in the dark; also by passing SO₂ over charcoal saturated with Cl (Melsens, C. R. 76, 92).—4. Ozone produces SO₂; the same compound is formed by passing SO, mixed with oxygen over heated spongy Pt, or by burning SO, in O under a pressure of several atmos. (v. supra; Formation, No. 1). According to Berthelot (A. Ch. [5] 12, 463), S₂O, is formed by the action of the silent discharge on SO, mixed with oxygen. The compound S2O2N2 is said to be formed by passing electric sparks through a mixture of SO2 and air. This compound is probably S₂O₅(NO₂)₂ (v. NITROSULPHONIC ANHYDRIDE, p. 601). -5. Passed over red-hot carbon the products are CO, CO2, and S; COS and CS, are also sometimes formed, probably by secondary reactions (Scheurer-Kestner, C. R. 114, 296; cf. Berthelot, Bl. [2] 40, 362; Eilsart, J. 1885. 457).—6. Potassium burns when heated in SO₂, giving K₂SO₄, K₂S₂O₃, and polysulphides .-- 7. Many other metals, especially when in fine powder, burn in SO2; Sb and Sn form oxides and sulphides; Fe, sulphide and sulphate; Pd and Pt, sulphides and SO,; Au, S and SO, Mg, sulphate, sulphite, and sulphide; Al, Co, Ni, and Zn also burn; Bi and Hg do not react (Schiff, A. 67, 94; 117, 92; Uhl, B. 23, 2151).—8. Water dissolves SO₂, torming H₂SO₂Aq (v. Sulphurous acid, p. 614).
9. Several metallic peroxides combine readily with SO₂, forming sulphates; e.g. PbO₂+SO₂ = PbSO₄; in this reaction much heat is produced .- 10. Nitrogen dioxide is said to react with liquid SO₂ to form nitrosulphonic anhydride S₂O₅(NO₂)₂ (v. p. 601). With nitric oxide or dioxide, in presence of water, nitrosulphonic acid NO₂(SO₂OH) (q. v., p. 601) is formed. Nitric oxide in presence of alkali produces salts of SO2.OH(NO)2.H (v. NITROSOSUL-PHATES, p. 581).—11. Selenion dioxide does not react with SO₂ (Schulze, J. pr. [2] 32, 390).—12. Dry hydrogen sulphide does not react; moist H₂S, at the ordinary temperature, produces H₂S₅O₆, S, and H₂O (Cluzel a. Schmid, Chem. Zeitung, 11,50); there is said to be no reaction in presence of water above 400° (Mulder, J. 1858. 84).—13. Phosphoretted hydrogen reacts at the ordinary temperature to form H₂O and P₂S₃ (C. a. S., l.c.).—14. Ammonia combines with SO₂ probably producing SO.NH. (=SO.OH.NH₂) and SO₂2NH₂ (=SO.ONH₂NH₂) (v. Thionamic acid).—15. Hydrochloric or hydriodic acid produces H₂O, S, and Cl or I.—16. Many oxysalts, e.g. nitrates and chlorates, react with SO₂, when heated, to form sulphates and N oxides; at higher temperatures SO₂ is sometimes produced (v. Hodgkinson a. Young, C. N. 66, 199).—17. Alkali nitrites yield salts of nitrilosulphonic acid N(SO₂.OH)₂ (q. v., p. 601), and salts of oxy-imidosulphonic acid N(OH)(SO₂.OH)₂ (q. v., p. 602).—18. Combines with aluminium chloride at 50°-60°, forming AlCl₂.SO₂, which gives off SO₂ at a higher temperature (Adrianoffski, B. 12, 688).—19. Phosphorus pentachloride produces POCl₂ and SOCl₂ (Kremers, A. 70, 297; Schiff, A. 102, 111).

SULPHUR TRIOXIDE SO. (Sulphuric oxide. Sulphuric anhydride.) Mol. w. 79-86. Melts at 14-8° (Weber, B. 19, 3189; Rebs, A. 246, 356). Some other observers have given m.p. as high as 29-5° (v. Buff, A. Suppl. 4, 129; Bussy, A. Ch. [2] 26, 411; Fischer, P. 16, 119; Marignac, A. 88, 230). Boils at 46-2° (Weber, l.c.; v. also Buff, l.c.; Fischer, l.c.; Schultz-Sellack, P. 139, 480). S.G. 1-94 at 16° (Weber, l.c.); 1-97 at 20° (Bussy, l.c.); 1-9086 at 25° (Buff, l.c.). V.D. 39-78 (Schultz-Sellack, P. 139, 480; Perman, Pr. 48, 45). H.F. [S,O*] = 103,240 (liquid SO, from solid S; Th. 2, 254); [SO², O] = 32,160 (liquid SO, from gaseous SO,; Th., l.c.); [SO*, Aq] = 39,170 (liquid SO,; Th., l.c.). For heat of vaporisation v. Berthelot (C. R. 90, 1510). Schultz-Sellack gives vapour-pressure at 20° as 152 mm. Hg (P. 189, 480).

20° as 152 mm. Hg (P. 189, 480).

Formation.—1. By the interaction of ozone and SO₂. Also by the action of an induction current on a mixture of SO₂ and O (Buff a. Hofmann, A. 113, 129; Deville, Bl. [2] 3, 366); Berthelot (C. R. 86, 20) says that the product is S₂O₂.—2. By burning S in O under a pressure of several atmos. (v. SULPHUB DIOXIDE, Formation No. 1). Also by heating pyrites (Scheurer-Kestner, C. R. 99, 917; Lunge, B. 10, 1824).—3. By passing a mixture of SO₂ and O over heated Pt (Phillips a. Magnus, P. 24, 610), spongy Pt (Hodgkinson a. Lowndes, C. N. 57, 193), or certain oxides, e.g. Fe₂O₃, CuO, Cr₂O₃ (Mahla a. Wöhler, A. 81, 255).—4. By heating certain sulphates or bisulphates, e.g. Fe₂(SO₄)₃, Bi₂(SO₄)₃, Ag₂SO₄, or NaHSO₄.—5. By distilling Nordhausen sulphuric acid (v. Osann, D. P. J. 151, 158); also by heating conc. H₂SO₄ with P₂O₄ (Barreswil, C. R. 25, 30; Evans, Ph. 8, 127).

Preparation.-1. Furning sulphuric acid is heated at as low a temperature as possible; the impure SO, is collected in a dry retort and redistilled once or twice, the most volatile portion being collected each time. The partially purified SO, is then distilled into a small tube, c. 8-10 mm. wide and about the length of a finger, containing some dry P2Os; this tube is narrowed at one end and fused on to a thick-walled tube c. 1-1.5 mm. wide, which is bent round and fused to a receiving tube c. 12-15 mm. wide and 800 mm. long; the tube containing the SO, and P.O. is heated in a water-bath to 90°-100° for 6 or 8 hours, the receiving tube projecting upwards, so that the distillate flows back again into the distilling-tube. The position of the apparatus is then changed, so that the receiving tube is lower than the distilling tube, and the SO, is distilled from the P2O3 on a water-bath. When sufficient SO, has been collected, the narrow connecting tube is fused off, and the SO, is kept in the sealed receiving tube (Weber, B. 19, 3189) .-

2. A mixture of SO, and O, in the ratio SO₂:O, is passed over platinised asbestos heated nearly to redness; the SU, is condensed in a small dry tube, in which it is sealed up. The mixture of SO, and O is obtained by strongly heating conc. H₂SO₄and removing H₂O (H₂SO₄ = H₂O + SO₂ + O) by cooling and passing through coke moistened with 60 p.c. H₂SO₄Aq (Winkler, D. P. J. 218, 128). The asbestos is platinised by soaking in PtCl,Aq made alkaline by soda, and mixed with sufficient HCO2Na to reduce the Pt, drying at 100°, carefully washing out the salts with water, and drying (Thomsen, B. 8, 496). The SO₂ thus obtained is not quite pure. It may be purified by distillation with P₂O₂ (v. supra).

Properties.—A colourless, mobile liquid, solidifying at c. 15° to long, prismatic, transparent crystals, which melt at 14.8°. According to some observers (v. Marignac, Ar. Ph. 22, 225; 58, 228; Schultz-Sellack, P. 139, 480) the crystals of SO, slowly change to a mass of tough, opaque, lustrous needles, which melt above 50°, and constitute a distinct modification of SOs; but the experiments of Weber (P. 139, 480; B. 19, 3187; cf. Rebs, A. 246, 356) have shown that pure SO, exists only in one form, and that the substance with higher m.p. is formed by the action of traces of H2SO4 in the SO2; a small trace of moisture suffices to cause the gradual change. SO, is very acrid and poisonous; it chars paper, wood, and organic matter generally. SO, is extremely hygroscopic; it dissolves in water with a hissing sound and production of much heat ([SO,Aq] = 39, 170), forming H_2SO_4Aq . Non-conductor of electricity (Magnus, P. 104, 553); not decomposed by electric sparks, but by passing through a red-hot tube gives SO₂ and O. SO₂ does not redden dry litmus. It is a strongly acid-forming oxide (v. SULPHURIC ACID).

Reactions and Combinations .- 1. Heated to redness (by passing through red-hot tube) gives 80, +0.-2. Not decomposed by electric sparks (Magnus, P. 104, 553); electrolysis of SO, in H.SO, causes separation of S and O, and the solution becomes blue (Geuther, 4. 109, 129).—

8. Water reacts violently with SO, forming H₂SO, Aq (v. Sulphuric acid).—4. Hydrogen sulphide produces H₂SO, and S.—5. A mixture of SO, and oxygen, in the ratio 2SO,:0, forms S2O, when submitted to the silent electric discharge (Berthelot, A. Ch. [5] 14, 345).-6. Sulphur dissolves in SO, forming S₂O₃; selenion and tellurium react similarly (v. SULPHUR THIO-OXIDE, SELENION THIO-OXIDE, and TELLUBIUM THIO-OXIDE) .- 7. Phosphorus ignites in vapour of SO, forming PO, and S.—8. Iodine dissolves in SO, apparently forming several compounds (v. Wach, S. 50, 37; Schultz-Sellack, B. 4, 109), Weber (J. pr. [2] 25, 224) describes the compounds I(SO₂)₂, I(SO₂), and I₂(SO₂).—9. Iron and zinc react at red heat to give sulphides and oxides; mercury produces sulphate and SO₂ (d'Heureuse, P. 75, 255).—10. Several metallic sulphides—e.g. of Sb, Pb, K—form sulphates and SO₂ when heated with SO₂ (Weber, P. 130, 329).—11. Dry sulphur dioxide is absorbed by SO, at 0°, forming a thin, fuming liquid, approximately SO, 2SO, (H. Rose, P. 89, 178). According to Schultz-Sellack, liquid SO, and SO, are miscible in all proportions (P. 139, 480). 12. Selenion dioxide combines to form SO. SeO.

(v. Selenion dioxide, Combinations, No. 4, p. 441).—18. Nibric oxide, in absence of moisture and O, forms S₂O₂N₂, which is generally called nitrosulphonic anhydride (q. v., p. 601) and written S₂O₂(NO₂)₂. Nitrogen dioxide (made by heating Pb(NO₂)₂) seems to form SO₂,NO₂, and on heating this and passing in more NO, O is given off and the compound S,O,N, remains (v. Weber, P. 123, 337; Brüning, A. 98, 877). The formula of the first of these compounds is generally doubled, and the compound regarded as S_2O_a .O. $(NO_2)_2 = oxynitrosulphonic anhydride (q. v., p. 601); the second is described in some books as <math>S_2O_7(NO)_2$ and is called nitrosyl pyrosulphate, but from the descriptions of Weber and Brüning it is evidently the same as the compound S₂O₂N₂ formed from NO and SO₃, i.e. it is nitrosulphonic anhydride.— 14. SO, combines with iodine pentoxide to form SSO₃LO₃ and SO₃.5LO₃ (Weber, B. 20, 86; Kämmerer, J. pr. 83, 72). With phosphorus pentoxide 3SO₄.P₂O₃, decomposing at 30°, is formed (Weber, B. 19, 3185; 20, 86). Arsenious 632). According to Baker (C. J. Proc. 1893. 130), dry SO, does not react with dry oxides of barium, calcium, or copper.—15. The compound SO. N. O. 5H. SO. is said to be formed by passing SO, into pure, well-cooled, nitric acid (Weber, P. 123, 233; 142, 602).—16. Boric acid forms BH,O,3SO, [=B(HSO₄),] (D'Arcy, C. J. 55, 155; cf. Merz, J. pr. 99, 81; Schultz-Sellack, B. 4, 15).-17. Hydrobromic and hydriodic acids separate Br, and I, and form H2SO4 and SO2.— 18. Hydrochloric and hydrofluoric acids form Cl(SO₂.OH) and F(SO₂.OH) (v. Chlorosulphonic ACID, p. 599; and FLUOSULPHONIC ACID, p. 600). 19. Conc. sulphuric acid forms H,S,O, (v. Pyro-SULPHURIC ACID, p. 625).—20. Ammonia produces NH₂(SO₂.OH) and salts of this acid (v. SULPHAMIC ACID AND SALTS, p. 567); and also salts of NH(SO₂.OH)₂ (v. IMIDOSULPHONIC ACID AND SALTS, p. 600).—21. Phosphoretted hydrogen produces SO₂ and red P (H. Rose, P. 24, 140; Aimé, J. pr. 6, 79).—22. Phosphorus trichloride reacts violently, forming POCl₂ and SO₂ (Armstrong, B. 8, 732; Michaelis, J. Z. 6, 239). Phosphorus pentachloride produces pyrosulphuryl chloride S₂O₅Cl₂ (q. v. under Sulphur oxyohlorides, p. 617).—23. Boron chloride re-acts at c. 120° to form S₂O₅Cl₂ (Prudhomme, Bl. [2] 14, 385).—24. Nitrosyl chloride (produced by heating aqua regia) produces SO₂NOCl = NO₂(SO₂Cl) (v. NITEOSULPHONIC CHLOBIDE, p. 601). — 25. Pyrosulphuryl bromide S₂O₂Br₂ (q. v., p. 617) is perhaps formed by the reaction of bromoform with SO₂.—26. Selenion tetrachloride produces SSeO.Cl. (v. SELENION THIO-OXYCHLORIDE, p. 441).

SULPHURIO PEROXIDE S.O.. (Sulphur heptoxide. Persulphuric anhydride.) Mol. w. not known with certainty but probably 175.68 (S2O,). This oxide was prepared by Berthelot in 1877 A. Ch. [5] 14, 845) by passing the 'silent' electric discharge, for several hours, through a mixture of SO, and O in the ratio 2SO,:O, or of SO, and O in the ratio 280, 30. (For apparatus used v. B., A. Ch. [5] 12, 463.) Oily drops formed on the sides of the tube, and on reducing the temperature nearly to 0° they solidified to long, white, crystalline, needles. S.O., resembles solid SO, in appearance; it remains unchanged in a sealed tube for some days at 0°: when heated it gives SO, and O; dissolves in conc. H₂SO₄ forming a fairly stable solution; fumes in the air; in water it rapidly decomposes forming H.SO.Aq and giving off a brisk stream of O; with BaOAq forms BaSO, and O; and also some Ba persulphate, according to Berthelot (l.c.; no analyses given). By electrolysing H₂SO,Aq, containing water and acid in the ratio 10H₂O:H₂SO₄, B. (C. R. 90, 269) obtained a quantity of persulphuric acid (H2S2O2; v. under SuL-PHURIC ACID, p. 625) equal to from 88 to 123 g. S2O, per litre; using an acid of the concentration H₂SO₄:2H₂O to H₂SO₄:3H₂O, B. (l.c.) says that a compound S2O, 2H2O2 is formed, which is decomposed by excess of conc. H2SO4 chiefly to H2S2O8, and by dilution with water to H2S2O8Aq and H₂O₂Aq which after a time decompose to H.SO, Aq, H.O, and O.

S2O, is the anhydride of persulphuric acid $H_2S_2O_3$ (q. v., p. 625). Mendeléeff (B. 15, 242) regards S2O, as analogous in constitution to

OSO2 H₂O₂, and writes the formula oso₂⁄

looks on the corresponding acid as derived from H₂O₂ by replacing 2H by 2SO₂OH, thus: O.(SO2.OH)

O.(SO₂.OH)

According to Traube (B. 22, 1518; 24, 1764; 25, 95), the liquid obtained by electrolysing 40 p.c. H.SO,Aq contains an oxide to which he gives the formula SO, and calls sulphuryl hyperoxide (or holoxide), SO2O2. Traube did not isolate this compound; his argument for its existence is based on the results of indirect analyses. In the light of the criticisms of Carnegie (C. N. 64, 158) it may be concluded that the existence of SO, is extremely improbable. In a later communication (B. 26, 1481) Traube thinks it is probable that SO, does not exist.

Sulphur, oxyacids of. For compositions of these acids, and references to articles where they are described, v. SULPHUR, OXYACIDS OF (p. 619).

Sulphur, oxyacids of, nitrogen derivatives of. For a general statement regarding these compounds, and references to the articles where they are described, v. SULPHUR OXYACIDS,

NITROGEN DERIVATIVES OF (p. 619). Sulphur, oxybromides of. There is very little definite knowledge regarding the compounds of S, O, and Br. According to Michaelis (J. Z. 6, 239, 296) PBr, does not react with SO2, and a mixture of PCl, and Br (in the ratio PCl.Br.) produces POCl. and S bromides. Rose (P. 44, 327) and Clausnizer (B. 11, 2012) could not obtain an oxybromide by the reaction of SO, and S bromide, nor by the action of SO, and HBr.

In his memoir on S₂O₄Ol₂ prepared by the reaction of SO, with CHCl, Armstrong (B. 2, 712) states that CHBr, reacts with SO, similarly to these products are obtained by passing vapour CHCl, but he does not describe the product. If of SOCI, through a red-hot tube (Heumann a. the reaction of CHBr, with SO, is strictly similar Köchlin, B. 16, 1625).

to that of CHOL the product would be pyrosulphuryl bromide S.O.Br.. Odling (C. J. 7, 2) supposed that sulphuryl

oxy-bromide SO₂Br, was formed by the action of sunlight on SO₂ and Br, but Sestini (Bl. [2] 10, 226) and Melsens (C. R. 76, 92) failed to obtain an oxybromide by this method.

THIONYL BROMIDE SOBr. This compound was obtained, but not quite free from S bromides, by Hartog a. Sims (C. J. Proc. No. 118, 1893. 10) by the interaction of SOCl, and NaBr; it is described as a very hygroscopic, crimson liquid, S.G. 2.68 at 18°, decomposing at 150° to Br, S bromides, &c.

Sulphur, oxychlorides of. Five compounds have been isolated, SOCl₂, SO₂Cl₂, S₂O₄Cl₂, S₂OCl₄, and S₂O₂Cl₄.

THIONYL CHLORIDE SOCI, (Sulphurous oxychlorids). Mol. w. 118.62. Boils at 78.8°; S.G. 40 1.67673, S.G. at b.p. 1.52143 (Thorpe, C. J. 37, 354; for other data v. Carius, A. 106, 303; 111, 93; Wurtz, C. R. 62, 460). V.D. at c. 150°=56, at c. 440°=39 (Heumann a. Köchlin, B. 16, 1625). H.F. [S,O,Cl²]=40,800; heat of vaporisation = 6,480 (Ogier, C. R. 94, 82). S.V. 78.01 (Thorpe, l.c.).

Formation.—1. By the reaction of PCl_s with SO₂ (Kremers, A. 70, 297; Schiff, A. 102, 111), SO₂Cl₂, C₈H₃.SO₂Cl, and several other organic compounds containing S (v. Kekulé a. Barbaglia, B. 5, 875; Carius, J. pr. [2] 2, 262), sulphites, or thiosulphates; also by the reaction of POCl, with sulphates.—2. By the action of Cl₂O on CS₂, or on S in S₂Cl₂ at -12° (Wurtz, C. R. 62, 460).—3. By the interaction of SO. and SCl, (Michaelis a. Schifferdecker, B. 5, 924 6, 993). - 4. Along with SO₂Cl₂ by the gradual decomposition of S.O.Cl. (q. v., p. 619).

Preparation .- 1. About 100 g. dry PCl, are placed in a retort connected with a reversed condenser, and a stream of dry SO, is led in till the PCl, has liquefied; another 100 g. PCl, is added and the stream of SO2 is continued; when sufficient PCl, has thus been decomposed (PCl₃ + SO₂ = SOCl₂ + POCl₂) the liquid is heated for some time (the reversed condenser being still attached) to get rid of SO₂, and then fractionated repeatedly (SOCl, boils at 78.8° and POCl, at 107-2") (Carius, A. 106, 303; 111, 98; Wurtz, A. 139, 375; Michaelis, A. 170, 1; Thorpe, C. J. 37, 354).—2. S₂Cl₂ is placed in a small flask, cooled to c. —20°, a stream of dry Cl is led in till the liquid is saturated, and then SO, is distilled into the flask from a solution in conc. H2SO4; the weights of S2Cl2 and SO2 used should be in the ratio $S_2Ol_2:4SO_2$ (= 1:2.37); convenient quantities to use are 22 g. S.Cl., and 50 g. SO₂ (SCl₄ + 2SO₂ = SOCl₂ + S₂O₃ Cl₂). The liquid is fractionated finally from a small quantity of SO₂ (Michaelis a. Schifferdecker, B. 5, 924; 6, 998).

Properties.—A colourless, very refractive liquid, fuming in the air, with a very penetrating odour; vapour attacks the eyes and respiratory organs. V.D. 56 at c. 150°, but 89 at c. 440°; the smaller value corresponds with the decomposition 4SOCl₂ = S₂Cl₂ + 2SO₂ + 8Cl₂;

(O., l.c.).

Reactions and Combinations.—1. Moist air produces HCl and SO₂.—2. Water also forms HClAq and SO₂, warm water forming in addition H₂SO₄Aq and S (Carius, I.c.).—8. Hydrogen sulphide produces SO₂, HCl, and S (Prinz, A. 223, 871).—4. Phosphorus pentasulphide reacts to form SO₂, PSCl₃, and S, and some S₂Cl₂(Prinz, l.c.).—5. With antimony trisulphide the products are SbCl, SO, and S.-6. Sulphur, heated to 180° with SÖCl₂, forms S₂Cl₂ and ŠO₂; selenion forms S₂Cl₂ and SSeCl₂ (Prinz, *l.c.*).—7. Powdered antimony reacts without heating, forming SbCl, Sb₂S₃, and SO₂.—8. POCl₃, PSCl₃ and PCl₃ are formed by heating SOCl, with phosphorus tri-chloride (Michaelis, J. Z. 6, 239).—9. Heating with sulphur dichloride produces S2OCl, (q. v., p. 619) (Ogier, Bl. [2] 37, 293).—10. Ammonia does not produce thionylamide SO(NH₂)₂, but forms NS, NH₄Cl, (NH₄)₂SO₃, and NH₄ polythionates (M., l.c.).—11. Reacts rapidly with silver nitrate, forming NO₂(SO₂.Cl) (Thorpe, C. J. 41, 297).—12. SOCl₂ absorbs sulphur disciple and shorpe form (M. 1). dioxide and chlorine freely (M., l.c.).-13. For reactions with various organic compounds v. Michaelis (l.c.); Heumann a. Köchlin (B. 16, 1625); Böttinger (B. 11, 1407).

Sulphuryl chloride SO₂Cl₂ (Sulphuric oxychloride). Mol. w. 134·58. Boils at 69·95° (Thorpe, C. J. 37, 359; for other determinations v. Regnault, J. pr. 19, 243; Clausnizer, B. 11, 2010; Ogier, C. R. 94, 82). S.G. 6 1·70814; at b.p. 1·56025 (Thorpe, l.c.). V.D. 65 at 184°; 34 at 440° (Heumann a. Köchlin, B. 16, 602; cf. Regnault, J. pr. 18, 97; Behrend, J. pr. [2] 15, 23). S.V. 86·29 (Thorpe, l.c., p. 372). H.F. [S.O²,Cl²] = 89,780; [SO²,Cl²] = 18,700 (Th. 2, 310; cf. Ogier, C. R. 94, 82). S.H. (15°-63°) ·233 (Ogier, l.c.). Heat of vaporisation 7,060

Formation.—1. By the direct combination of SO₂ and Cl in sunlight, or by reacting on a mixture of C₂H₄ and SO₂ by Cl in sunlight (Regnault, J. pr. 18, 93; 19, 243). Schulze (J. pr. [2] 28, 351; 24, 168) recommends to allow SO₂ and Cl to react in presence of camphor. Melsens (C. R. 76, 92) passes SO₂ and Cl into glacial acetic acid.—2. By passing SO₃ over charcoal saturated with Cl (M., l.c.).—3. By the interaction of PCl₃ with H₂SO₄ or SO₂ (Williamson, P. M. [4] 7, 365; Schiff, A. 102, 111); according to Michaelis (J. Z. 6, 233, 292), SO₂ and PCl₃ form S₂O₃Cl₂.—4. By heating SO₃ and BCl₃, in the ratio 2SO₃:BCl₃, to 120° in a closed tube (Gustavson, B. 6, 9).—5. By heating Cl(SO₂.OH) to c. 200° for some time (Behrend, B. 8, 1004; cf. Otto a. Beckurts, B. 11, 2058).

Preparation.—1. Cl and SO₂ are passed into

Preparation.—1. Cl and SO, are passed into camphor, and the colourless liquid is fractionated (Schulze, l.c.).—2. Cl(SO₂.OH) (v. Chicorosulphonic acid, p. 509) is heated to c. 200°, in a closed tube, for some hours, when it decomposes thus, 2[Cl(SO₂.OH)] = SO₂Cl₂ + H₂SO₄; the liquid is distilled, the portion coming over at 67°-71° being collected separately and then repeatedly fractionated (Behrend, l.c.; cf. Thorpe, l.c.).

Properties and Reactions.—A colourless

Properties and Reactions.—A colourless liquid, furning slightly in air. 1. Decomposed by heating to dull redness into SO_2 and Cl; decomposition is complete at 440° (Heumann a. Köchlin, B. 16, 602); but does not begin at 250°

(Beckurts a. Otto, B. 11, 2060).—2. Water produces HClAq and H₂SO,Aq; a small quantity of water is said to form Cl(SO₂OH)Aq.—3. Excess of SO₂Cl₂ added to a little alcohol forms Cl(SO₂OEt); with excess of alcohol SO₂(OEt)₂ is produced (Behrend, B. 9, 1384).—4. Phosphorus pentachloride reacts to form POCl₃, SOCl₂, and Cl (H. a. K., B. 15, 1736).—5. Phosphorus, arsenic, and antimony produce PCl₄, AsCl₃, and SbCl₂ respectively, and SO₂ (H. a. K., l.c.).—6. Ammonia forms NH₃Cl and SO₂NH₂ONH₄.—7. With many organic compounds SO₂Cl₂ reacts like PCl₃ or POCl₄, substituting Cl for H (v. Dubois, Z. [2] 2, 705; Michaelis, A. 170, 1).

PYROSULPHURYL CHLORIDE $S_2O_5Cl_2 = Cl.SO_2.O.SO_2.Cl.$ (Disulphuryl chloride. $S_2O_5Cl_2 = Cl.SO_2.O.SO_2.Cl.$ (Disulphuryl chloride. $S_2O_5Cl_2 = Cl.SO_2.O.SO_2.Cl.$ (Disulphuryl chloride anhydride.) Mol. w. 214·38. Boils at 139·59° (Thorpe, C. J. 37, 360; for other determinations v. Rose, P. 44, 291; Michaelis, J. Z. 6, 235, 240, 292; Rosenstiehl, C. R. 53, 658; Schützenberger, C. R. 69, 352). S.G. $\frac{60}{40}$ 1·85846, at b.p. 1·60610 (Thorpe, l.c.). V.D. at slightly above b.p. = 104; between 160° and 240° = 54·8 (Heumann a. Köchlin, B. 16, 479; Kanowaloff, B. 16, 1127; cf. Rosenstiehl, C. R. 53, 658; Rose, P. 44, 291; Ogier, C. R. 94, 82, 217; 96, 648). H.F. [S²,O²,Cl²] = 180,600 (Kanowaloff, l.c.); Ogier (l.c.) gives 159,400. Heat of vaporisation 7,570 (K.‡ l.c.); 13,160 (O., l.c.). S.H. ·258 (K., l.c.).

Formation.—1. By the interaction of PCl₃ and SO₃ or Cl(SO₂·OH).—2. By heating P₂O₃ and Cl(SO₂·OH) (Billitz a. Heumann, B. 16, 483).—3. By heating SO₃ with NaCl (R., l.c.), with S₂Cl₂ (Rose, P. 44, 291), with CCl₄ (Schützenberger, C. R. 69, 352; Kanowaloff, C. R. 95, 1284), with C₂Cl₃ (Prudhomme, C. R. 70, 1137), with CHCl₄ (Armstrong, C. J. [2] 1, 244), or with SiCl₄ (Gustavson, B. 5, 332).—4. By the interaction of POCl₂ and SOCl₂ (Michaelis, J. Z. 6, 235, 240, 292).

Preparation.—1. P₂O₃ and Cl(SO₂.OH) are heated in a flask with a reversed condenser, using rather more Cl(SO₂.OH) than P₂O₃; when HCl and SO₂ cease to be given off, the liquid is distilled, and the portion coming over between 135° and 145° is fractionated (B. a. H., l.c.).—2. S₂Cl₂ in a flask is surrounded by a freezing mixture, and saturated with SO₃; the product is fractionated; 5SO₃ + S₂Cl₂ = 5SO₂ + S₂O₃Cl₂ (Rose, P. 44, 291). Thorpe (C. J. 37, 360) found that the product of this reaction boiled almost wholly between 135° and 138°.

Properties and Reactions.—A colourless, mobile, very refractive liquid, fuming in the air with formation of Cl(SO₂.OH).—1. Decomposed by heat to SO₃, SO₂, and Cl (Rose, P. 44, 291); the V.D. determinations of Heumann a. Köchlin (B. 16, 479) show that at 440° the decomposition is S₂O₃Cl₂=SO₃+SO₂+Cl₂.—2. Sinks in water, and is slowly decomposed to HClAq and H₂SO₄Aq; if a very little water is used Cl(SO₂.OH) is formed, thus S₂O₄Cl₂+H₂O=2Cl(SO₂.OH) (v. B. a. H., l.c.)—3. Reacts with phosphorus, sulphur, antimony, copper, iron, sinc, and some other metals, to produce chlorides, SO₃ and SO₂ (Heumann a. Köchlin, B. 16, 479).—4. Phosphorus trichlorides produces

PCl₂, POCl₂, and SO₂; phosphorus pentachloride forms POCl₃, Cl, and SO₂ (Geuther, B. 5, 925). 5. Selenion tetrachloride gives SSeO, Cl, (Clausnizer, B. 11, 2010).-6. Chromates react to form CrO,Cl, (Rosenstiehl, C. R. 53, 658).—7. Ammonia forms a white solid, which dissolves in water, producing NH₄Cl and a sulphamate of NH (Rose, l.c.).

SULPHUR MONOXYTETRACHLORIDE S.OCI. By heating equal weights of S2Cl2 and SO2Cl2 to 250° in a sealed tube, Ogier (Bl. [2] 37, 293) obtained a deep-red liquid resembling S2Cl2, S.G. 1.656 at 0°, decomposing under 100° to S₂Cl₂, SO₂, and S, to which he gave the composition S₂OCl₄. Decomposed by water to H₂SO₄Aq, SO₂Aq, HClAq, thionic acids, and S.

SULPHUR TRI-OXYTETRACHLORIDE S₂O₃Cl₄. Mol. w. not determined. This compound was obtained by Millon (A. Ch. [3] 29, 327) by the action of moist Cl on S₂Cl₂ (cf. Marchand, J. pr. 22, 507; Carius, A. 106, 295). It is conveniently prepared by the reaction SCl₄ + Cl(SO₂.OH) = HCl+S2O2Cl4; a stream of dry Cl is led into a mixture, in the ratio of equal numbers of molecules, of S_2Cl_2 and $Cl(SO_2.OH)$, kept at c. -20° until a semi-solid mass is obtained (a very wide tube must be used to lead in Cl); the temperature is then kept at c. -13° and the Cl is continued till the mass gets quite solid; the freezing mixture is then removed, and the Cl is continued till the mass is white, when the excess of Cl is removed by a stream of dry CO, (Michaelis a. Schifferdecker, B. 6, 996).

S2O2Cl4 is a white, crystalline solid, with a most disagreeable odour; the vapour acts on the eyes. Melts at 57°, giving off SO₂ and Cl, and leaving a liquid that solidifies chiefly to SOCl₂ and S₂O₃Cl₂; a portion sublimes in fine white needles. Ice-cold water produces SO₂Aq, H₂SO₄Aq, and HClAq; water at the ordinary temperature also separates S. Deliquesces in a partially closed vessel, giving off HCl, Cl, and SO, and leaving a liquid that gives SOCl, and S2O4Cl2 on distillation. Conc. H2SO4 reacts violently, forming HCl, SO2, and Cl(SO2OH); CS2 produces very rapid decomposition to COCl2, CO, SO₂, S.Cl₂, and SOCl₂. On long standing in a closed tube S₂O₂Cl₄ separates into equal numbers of molecules of SOCl, and SO,Cl, (M. a. Mathias, B. 6, 1452).

Sulphur, phosphides of, v. Phosphorus sul-**РИГОЕВ**, р. 145.

Sulphur, selenides of, v. Selenion sul-

PHIDES, p. 442.
Sulphur, silicides of, v. Silicon sulphides,

p. 462. Sulphur, tellurides of, v. Tellurium sul-

M. M. P. M. PHIDES, p. 655. SULPHUE, OXYACIDS OF. Several acids have been isolated, and salts of some others are known. The following list gives the names and compositions of the oxyacids of sulphur and their salts, with references to the articles where

these compounds are described:

HYPOSULPHUROUS ACID AND SALTS H2S2O4 and M.S.O.; v. Hyposulphurous acid under Sulphur-OUS ACID (p. 627), and hyposulphites under Sur-PHITES (p. 592).

SULPHUROUS ACID AND SALTS H2SO, MHSO, and M.SO.; v. SULPHUBOUS ACID (p. 626), and Sulphités (p. 587).

SULPHURIC ACID AND SALTS H2SO4, MHSO4 and M2SO4; v. SULPHURIC ACID (p. 620) and SULPHATES (p. 567).

Thiosulphuric acid and salts $H_2S_2O_3$, MHS_2O_3 , and $M_2S_2O_3$; v. Thiosulphuric acid (p. 708) and Thiosulphates (p. 705).

Thionic acids and salts $H_2S_2O_8$, $H_2S_4O_8$, $H_2S_4O_8$, $H_2S_4O_8$ (and salts); v. Thionic acids (p. 698) and Thionates (p. 695).

Pyrosulphuric acid and salts $H_2S_2O_7$, MHS₂O₇, and $M_2S_2O_7$; v. Pyrosulphuric acid under Sulphubic ACID (p. 625), and Pyrosulphates under Sulphates (p. 583).

PERSULPHURIC ACID AND SALTS H.S.O.Ag. M₂S₂O₈, and MS₂O₈; v. Persulphuric acid under SULPHURIC ACID (p. 625), and Persulphates under SULPHATES (p. 582). M. M. P. M.

SULPHUR OXYACIDS, NITROGEN DERI-**VATIVES OF.** In 1845 Fremy (A. Ch. [3] 15, 408) described the K salts of a number of acids obtained by the interaction of KNO, Aq and K₂SO₃Aq, and by boiling the products of this interaction with slightly acidified water; he called the compounds salts of sulphazotised acids. The examination of these salts was prosecuted by Claus a. Koch (A. 152, 336 [1874], and by Claus (A. 158, 52, 194 [1876]); Berglund (Bl. [2] 25, 455; B. 9, 252, 1896 [1875]); and Raschig (A. 241, 161; B. 20, 584 [1887]) also investigated the sulphazotised acids of Fremy; and the researches of Divers and Haga (C. J. Trans. 1885. 1886. 1889. 1892) have thrown much light on many classes of these compounds. The more recent investigations noted above have shown many of the formulæ given by Fremy to be inaccurate, and they have also led to a considerable revision of Claus's results. As the compounds that have been fairly thoroughly examined are noticed under the heading Sulphonic ACIDS AND DERIVATIVES (p. 509), it does not seem necessary to give an account of the other salts of sulphazotised acids, especially as it is very probable that the formulæ assigned to these compounds will require modification, and that some of the compounds will be found not to exist.

The chief sulphazotised compounds that have been investigated sufficiently to establish their composition satisfactorily are the following:-

Amidosulphonic acid and salts NH₂(SO₂.OH); described as SULPHAMIC ACID (p. 567).

Imidosulphonic acid and salts NH(SO, OH), (p. 600).

Nitrilosulphonates N(SO₂.OM), (p. 601).

Nitrosulphonic acid and salts NO, (SO, OH) (p. 601).

Nitrosulphonic anhydride (NO₂)₂S₂O₅ (p. 601). Nitrosulphonic chloride NO₂(SO₂.Cl) (p. 601). Nitroso-oxy-amidosulphonates

N(NO.OH)(SO.OM); described 8.6 sulphates under Sulphates (p. 581).

Oxy-amidosulphonic acid and salts (and NH(OH)(SO₂OH) salts derived therefrom) (p. 602).

Oxy-imidosulphonates (and salts derived

(p. 601). Sulphamide $(NH_2)_2SO_2$ (p. 567).

Sulphimide NH.SO, (p. 587).

Sulphimido-amide NH(SO, NH₂)₂ (p. 587).

Thionamic acid and salts NH₂(SO.OH)
(p. 695).

M. M. P. M.

SULPHURETTED HYDROGEN v. HYDROGEN SULPHIDE, vol. ii. p. 725.

SULPHURIC ACID; also PERSULPHURIC ACID and PYROSULPHURIC ACID.

SULPHURIC ACID H₂SO₄ (Oil of vitriol). l. w. probably 97.76. Melts at 10.85° Mol. w. probably 97.76. (Pickering, C. J. 57, 831); 10.5° (Thile, Chem. Zeitung, 16, 1688). Boiling begins at c. 290° SO, and H2O with some H2SO4.xH2O passing off; b.p. rises till 338° is reached, when it becomes constant and complete dissociation to SO₂ and H₂O occurs; for b.p. of H₂SO₄Aq from 95 to 5 p.c. H₂SO₄ v. Lunge (B. 11, 870). S.G. 1.8884 (Perkin, C. J. 63, 59; cf. Pickering, C. J. 57, 73, note; Kohlrausch, W. 17, 69; Schertel, J. pr. [2] 26, 246; Mendeléeff, B. 17, 30, 2536; Lunge a. Isler, Zeit. für anorg. Chemie, 1890. 129). V.D. 25 at 440°, corresponding with dissociation to $SO_s + H_2O$ (Deville a. Troost, C.R.56, 891; Wanklyn a. Robinson, Pr. 12, 507).
S.H. (22° to 80°) ·355; (22° to 170°) ·37
(Pfaundler, B. 3, 798); (16° to 20°) ·3315
(Marignac, Ar. Sc. 39, 217, 273). C.E. ·0005585
at 20° (Marignac, l.c.). H.F. [H², S, O¹] = 192,920; $[H'O, SO^3] = 21,320$ (formation of liquid H_2SO_4) (Th. 2, 255). Heat of fusion = 860 g. units (for 98 g. H₂SO₄) (Berthelot, C. R. 78, 716). T. C. v. Beetz (W. 7, 435), Weber (J. 1885. 121). $\frac{\mu-1}{d}$. Mol. w. = 32.45 (Gladstone, P. M. [4] 86, 311). For further data v. infra.

Occurrence. — Small quantities of H₂SO₄, from ·1 to ·5 p.c., are found in some rivers and springs in volcanic districts; in a pool in Texas, Mallet (C. N. 26, 147) found 5·29 g. H₂SO₄ per litre of water. The acid is also found in certain animal secretions; the salivary glands of some molluses are said to contain c. 2 p.c. (Bödeker a. Troschel, B. B. 1854, 486; De Luca a. Panceri, Bl. [2] 9, 400; Maly, M. 1, 205). The salts of sulphuric acid occur in large quantities (v. Sulphures, p. 567).

Formation.—The formation of an oily liquid by heating alum is spoken of by Geber in the eighth century; a similar liquid was formed in the fifteenth century by burning sulphur with saltpetre. These liquids were shown to be identical towards the end of the sixteenth century. Oil of vitriol seems to have been manufactured in fair quantities by heating ferrous sulphate from about the early part of the eighteenth century; the first manufactory for making the acid from sulphur was started at Richmond by Ward about 1740. The composition of sulphuric acid was determined by Lavoisier.

1. By long exposure of flowers of gulphur in the air (John, S. 14, 417; Wagenmann, P. 24, 601).—2. By heating S with H₂O to c. 200°; by electrolysing H₂O, using Pt electrodes with pieces of S attached, the H₂SO₄ forms at the positive pole (Becquerel, C. R. 56, 237).—3. By the oxidation of S in presence of H₂O by Cl, HNO₃, HCl and KClO₃, aqua regia, &c.; also by certain metallic salts (v. Wicke, A. 82, 146; Parkmann, Am. S. [2] 38, 328).—4. By oxidising SO₂Aq or SO₃ in presence of water, by Cl, Br, H₂O₂Aq,

NO₂, HNO₃, &c.; by passing moist SO₃ and air over hot Pt black (Phillips, A. 4, 171), through a red-hot tube containing pieces of glass (Magnus, P. 24, 610), or over red-hot sand (Blondeau, C. R. 29, 405).—5. By the oxidation of H₂S and polythionic acids (v. Cloëz a. Guignet, C. R. 46, 1110).—6. By the interaction of water and SO₂, SO₂Ol₂, SO₂Ol.OH, &c.

Preparation.—Commercial oil of vitriol is diluted until the S.G. is 1.58 (=50° Beaumé, or 106° Twaddell, =62.5 p.c. H_2SO_4 by weight); the acid is heated to 70° -80°, and $Na_2S_2O_2$ is added in quantity sufficient to decompose the As₂O₂ present (which should be estimated beforehand), according the equation to 8Na₂S₂O₃ + As₂O₄ = 8Na₂SO₄ + As₂S₄; the acid is stirred until the As₂S₄ separates in flocks (Thorn, D. P. J. 217, 495); the acid is filtered from As, S, through asbestos or glass-wool, and is then concentrated by evaporation in a Pt dish till it begins to fume, and then in a retort of Pt or hard glass. The conc. acid is now heated with a little pure (NH₄)₂SO₄, whereby HNO₂ and HNO₃ are decomposed (NH₄·NO₂ and NH. NO. are formed, and then decomposed to N and H₂O, and N₂O and H₂O). The acid is then distilled from a glass retort, heated by a ring-burner placed around the retort at about two-thirds of the distance from the level of the acid to the bottom of the retort. The distillate is re-distilled in small quantities from a Pt retort into a Pt receiver; the condenser being formed of Pt tubes, soldered with gold and fitting closely one into the other, and the temperature being such that the soid does not boil during the distillation (cf. Stas, Chem. Proport. 112). Various other methods for removing As have been proposed; for the preparation of H2SO4 free from As for chemico-medical use v. Bloxam (C. J. 15, 52), also Buignet a. Bussy (J. Ph. [3] 44, 177; 46, 252).

Commercial oil of vitriol is prepared by passing SO₂, mixed with a little HNO₃, into large leaden chambers into which jets of steam are also sent. The SO, is prepared by burning S or pyrites in a stream of air; the hot gas is passed over HNO, or over pots containing KNO. and H₂SO₄, so that it sweeps into the leaden chambers a certain quantity of HNO. The interactions between the hot SO, and HNO, probably produce NO and N₂O₄($88O_2 + 2HNO_2 + 2H_2O_3 + 2HO_3 + 2HO_3 + H_2O_4 + 2HO_3 + H_2O_5 + H_2O_3 + H$ $=2H_2SO_4+N_2O_3$; the NO then combines with O in the air of the chambers to form $N_2O_3(4NO+O_2=2N_2O_2)$, which interacts with SO_2 and H_2O to produce H_2SO_4 and $NO(N_2O_2+SO_2+H_2O=H_2SO_4+2NO)$:—if N_2O_3 is produced in the primary reaction of SO, with HNO,, this N.O. may take up O to form N.O., which in turn may react with SO, and H.O to produce H₂SO₄ and NO(N₂O₄+2SO₂+2H₂O = $2H_2SO_4$ +2NO). These cycles of changes proceed as long as SO, is sent into the chambers, and there is present a sufficiency of O, H₂O, and oxides of N. If there is not sufficient H₂O present to complete the decomposition of N.O. and SO, (or N₂O₄ and SO₂) to H₂SO₄ and NO, nitrosulphonic acid—NO₂(SO₂,OH)—is formed as a solid on the sides of the chamber $(3N_2O_3 + 2SO_3 + H_2O = 2NO_3(SO_2OH) + 4NO;$ or $3N.O_4 + 4SO_2 + 2H_2O = 4NO_2(SO_2OH) + 2NO)$

when steam is now sent into the chamber the nitrosulphonic acid is decomposed to H_2SO_4 and N_2O_3 (2NO₄(SO₂.OH) + $H_2O = 2H_2SO_4 + N_2O_3$). It is possible that the formation of NO₂(SO₂.OH) and the interaction of this with water may form normal stages in the series of reactions that occur in the chambers. Lunge (C. J. 47, 465) regards N_2O_2 as the chief carrier of O from the air to the SO₃; he formulates the two main interactions as

(1)
$$2SO_2 + N_2O_3 + O_2 + H_2O = 2NO_2(SO_2 OH)$$
,
(2) $2NO_2(SO_2 OH) + H_2O = 2H_2SO_4 + N_2O_2$;

moreover, he insists that the interactions take place for the most part between gases in contact with the surface of the vesicles of liquid water formed by the condensation of the steam sent into the chamber. (A fuller discussion of the reactions that occur in the leaden chambers will be found in the DIOTIONARY OF APPLIED CHEMISTRY, vol. iii. p. 726.)

Some portion of the oxides of N is swept through the chambers and so lost (part of this is recovered in the Gay-Lussac towers); some is lost by leakage, solution in the acid formed in the chambers, and in other mechanical ways; but some portion of the available oxides of N is also destroyed by secondary reactions that occur in the chambers, and result in the formation of N₂O and sometimes of N or NH₂ (v. Weber, P. 180, 329; Pelouze, A. Ch. [2] 60, 162; Lunge, B. 14, 2196; cf. DICTIONARY OF APPLIED

CHEMISTRY, vol. iii. p. 729). Properties.—A clear, colourless, oily, hygroscopic liquid; solidifying at c. 10.5°. The liquid may be cooled much below its freezing-point without solidifying, but addition of a crystal of H₂SO₄ or a small quantity of SO₂ causes crystallisation; the presence of a trace of water lowers the freezing-point to c.0°. H_2SO_4 is exceedingly acrid and burning, and is a violent poison. H.SO, fumes very slightly in air at the ordinary temperature; at 30°-40° it fumes much, giving off SO₂; according to Colefax (C. N. 63, 179), H₂SO₄ is probably slightly volatilised at ordinary temperatures. The mol. w. of H₂SO₄ has not been determined directly, but from the mol. weights of such derivatives as SO2Cl2, SO₂(OMe)₂, &c., the formula H₂SO₄ is most probably molecular. H₂SO₄ is decomposed when heated, and does not show a definite b.p.; boiling begins at c. 290°, and the temperature gradually rises to 888°, at which temperature dissociation to H₂O and SO₂ is complete. The residue has the composition H₂SO₄, ½H₂O; acid with this composition is obtained by boiling down either H₂SO₄xH₂O or H₂SO₄xSO₂ (Marignac, A. Ch. [3] 39, 184; Dittmar, C. J. [2] 7, 446). The sp.gr. of the vapour given off at c. 440° corresponds with that calculated for the dissociation of H₂SO₄ into SO₄+H₂O (Deville a. Troost, C. R. 56, 891; Wanklyn a. Robinson, Pr. 12, 507). The b.p. of H₂SO₄Aq, containing from 95 p.c. H₂SO₄ (b.p. = 295⁵), to 5 p.c. H₂SO₄ (b.p. 101°) is given by Lunge (B. 11, 370). H₂SO₄ freezes at 10.852° (v. Pickering, C. J. 57, 831; where a long series of determinations is given of f.p. of H₂SO₄Aq containing from '005 to 100 p.c. H₂SO₄). The S.H. and mol. heat $(=S.H. \times 98)$ are given by Pfaundler (B. 8, 798)

as follows:

Temp. interval		
from 22° to	8.H.	Mol. neat.
800	·855	84.790
90	.856	34.888
. 100	·858	85.084
110	•859	85.182
120	.860	85.280
180	.362	85.476
140	•364	85.672
150	•365	85·770·
160	.367	85 ·9 66
170	.370	86-260

For S.H. of H₂SO₄.xH₂O from 16° to 20°, x varying from 0 to 400, v. Marignac (Ar. Sc. 39, 217, 273); Bode (Zeit. fur anorg. Chemis, 1889. 244) gives S.H. of H₂SO₄Aq from S.G. 1842 to 1037; Pickering (C. J. 57, 160) gives S.H. of H₂SO₄Aq from c. 2 to c. 12 p.c. H₂SO₄ dissolves in water in all proportions; much heat is produced, and contraction occurs; for thermal measurements v. Thomsen (Th. 3, 44), Favre a. Silbermann (C. R. 24, 1081), F. a. Quaillard (C. R. 50, 1150), Pfaundler (J. 1869, 122), and, especially, Pickering (C. J. 57, 164). According to Pickering (C. N. 64, 14), the maximum contraction, calculated for unit weight of solution, changes from 67 p.c. at 8°, to 70.1 p.c. at 38°; but the maximum is constant at 76 p.c., from 8° to 38°, if calculated for unit volume; for fuller data v. P. (C. J. 57, 148). The expansion of H₂SO₄.xH₂O, at 20°, when x varies from 0 to 400, is given by Marignac (Ar. Sc. 39, 217, The refractive indices and dispersion of H₂SO₄Aq containing from 93.87 to 088 p.c. H₂SO₄ are given by Pickering (C. J. 63, 99; cf. van der Willingen, Ar. N. 8, 122). Perkin (C. J. 63, 57) gives Mol. R. for H₂SO₄Aq from 99.92 to 9.18 p.c. H₂SO₄ (= from H₂SO₄ + 53.87 H₂O to H₂SO₄ + 004 H₂O).

The electrical conductivities of H₂SO₄Aq of different concentrations have been measured by Kohlrausch (P. 164, 215; 159; 233; W. 6, 1; 17, 69), also by Pickering (C. J. 57, 158), who gives determinations for H₂SO₄Aq containing from 1 to 100 p.c. H₂SO₄ (v. also Bouty, C. R. 108, 393; also Grotrian, P. 151, 378).

The S.G. and composition of \dot{H}_2 SO₄Aq of varying concentrations have been determined by many observers. Pickering (C. J. 57, 152) gives S.G. and composition of \dot{H}_2 SO₄Aq containing from 5 to 100 p.c. \dot{H}_2 SO₄ at 0°, 2°, 4°, and each 2° up to 40°. Lunge a. Isler (Zeit. für anorg. Chemie, 1890. 129) give the following tables (cf. Mendeléeff, Z. P. C. 1887. 273; Pickering, C. N. 64, 311; Lunge, C. N. 65, 13):—

8.G. 15°	Degrees Beaumé Degrees Twaddell		we	arts by sight stain	1 litre contains kilos.		
(n vacuo	AA	T De	P.o. 80,	P.c. H _s 80 ₄	80,	н.80.	
1.000 1.005 1.010 1.015 1.020 1.025 1.030	0 0·7 1·4 2·1 2·7 8·4 4·1	0 1 2 8 4 5	0.07 0.68 1.28 1.88 2.47 8.07 8.67	0·09 0·83 1·57 2·30 8·03 8·76 4·49	0.001 0.007 0.013 0.019 0.025 0.032 0.038	0.001 0.008 0.016 0.023 0.031 0.039 0.046	
1.035 1.040	4·7 5·4	7 8	4·27 4·87	5·23 5·96	0.044 0.051	0.062	

48.G. $\frac{15^{\circ}}{4^{\circ}}$	Degrees Beaumé	Degrees Twaddell	we	arts by ight itain		contains los.	8.G. 15°	Degrees Beaumé	Degrees Twaddell	we	arts by eight stain		ont ains los.
in vacuo	28	PP	P.c. 80,	P.o. H _s 80 ₄	80,	H,804	in vacuo	AA	Pe	P.c. 80,	P.c. H ₂ SO ₄	80.	H,SO.
1.045	6.0	9	5.45	6.67	0.057	0.071	1.365	88.6	78	37.89	46.41	0.517	0.633
1.050	6.7	10	6.02	7.37	0.068	0.077	1.370	89.0	74	38.32		0.525	0.643
1.055	7.4	11	6.59	8.07	0.070	0.085	1.375	89.4	75	38.75		0.533	0.653
1·060 1·065	8.0	12 13	7.16	8·77 9·47	0.076 0.082	0.093	1·380 1·385	39·8 40·1	76	39.18		0.541	0.662
1.070	9.4	14	8.32	10.19	0.082	0.102	1.390	40.5	78	40.05	1	0.549 0.557	0.682
1.075	10.0	15	8.90	10.90	0.096	0.117	1.395	40.8	79	40.48		0.564	0.692
1.080	10.6	16	9.47	11.60	0.103	0.125	1.400	41.2	80	40.91		0.573	0.702
1.085	11.2	17	10.04	12.30	0.109	0.188	1.405	41.6	81	41.33	50.63	0.581	0.711
1·090 1·095	11·9 12·4	18 19	10.60 11.16	12·99 13·67	0.116	0.142	1·410 1·415	42·0 42·3	82 83	41.76		0.589	0.721
1.100	13.0	20	11.71	14.85	0.129	0·150 0·158	1.420	42.7	84	42.17	51.66 52.15	0.597	0.730 0.740
1.105	18.6	21	12.27	15.03	0.136	0.166	1.425	43.1	85	42.96	52.63	0.612	0.750
1.110	14.2	22	12.82	15.71	0.143	0.175	1.430	43.4	86	43.36	53.11	0.620	0.759
1.115	14.9	23	18.86	16.36	0.149	0.183	1.435	43.8	87	43.75	58.59	0.628	0.769
1·120 1·125	15·4 16·0	24 25	13·89 14·42	17·01 17·66	0·156 0·162	0.191	1·440 1·445	44.1	88	44.14	54·07 54·55	0.636 0.643	0·779 0·789
1.130	16.5	26	14.95	18.31	0.169	0.207	1.450	44.8	90	44.92	55.03	0.651	0.798
1.185	17.1	27	15.48	18.96	0.176	0.215	1.455	45.1	91	45.31	55.50	0.659	0.808
1.140	17.7	28	16.01	19.61	0.183	0.228	1.460	45.4	92	45.69	55.97	0.667	0.811
1.145	18.3	29	16.54	20.26	0.189	0.231	1·465 1·470	45.8	93	46.07	56.43	0.675	0.827
1·150 1·155	18·8 19·8	30 31	17·07 17·59	20·91 21·55	0.196	0.239	1.475	46·1 46·4	94	46·45 46·83	56·90 57·37	0.683 0.691	0.837 0.846
1.160	19.8	82	18.11	22.19	0.210	0.257	1.480	46.8	96	47.21	57.83	0.699	0.856
1.165	20.3	88	18.64	22.83	0.217	0.266	1.485	47.1	97	47.57	58.28	0.707	0.865
1.170	20.9	84	19.16	23.47	0.224	0.275	1.490	47.4	98	47.95	58.74	0.715	0.876
1·175 1·180	21.4	85 86	19.69	24.12	0.231	0.283	1·495 1·500	47.8	99	48.34	59.22	0.723	0.885
1.185	22·0 22·5	37	20·21 20·73	24·76 25·40	0.238 0.246	0·292 0·301	1.505	48·1 48·4	100 101	48·73 49·12	59·70 60·18	0·731 0·739	0·896 0·906
1.190	23.0	38	21.26	26.04	0.253	0.310	1.510	48.7	102	49.51	60.65	0.748	0.916
1.195	23.5	89	21.78	26.68	0.260	0.319	1.515	49.0	103	49.89	61.12	0.756	0.926
1.200	24.0	40	22.30	27.32	0.268	0.328	1.520	49.4	104	50.28	61.59	0.764	0.936
1·205 1·210	24·5 25·0	41	22·82 23·33	27·95 28·58	0·275 0·282	0.337 0.346	1.525 1.530	49·7 50·0	105 106	50.66 51.04	62·06 62·53	0·778 0·781	0·946 0·957
1.215	25.5	43	23.84	29.21	0.290	0.355	1.535	50.3	107	51.43	63.00	0.789	0.967
1.220	26.0	44	24.36	29.84	0.297	0.364	1.540	50.6	108	51.78	63.43	0.797	0.977
1.225	26.4	45	24.88	80.48	0.805	0.873	1.545	50.9	109	52.12	68.85	0.805	0.987
1·230 1·235	26·9 27·4	46 47	25·39 25·88	81·11 81·70	0·312 0·320	0·382 0·391	1.550 1.555	51·2 51·5	110 111	52·46 52·79	64·26 64·67	0.813 0.821	0.996 1.006
1.240	27.9	48	26.35	82.28	0.320	0.400	1.560	51.8	112	53.12	65.08	0.829	1.015
1.245	28.4	49	26.83	82.86	0.334	0.409	1.565	52.1	118	53.46	65.49	0.837	1.025
1.250	28.8	50	27.29	83.43	0.341	0.418	1.570	52.4	114	53.80	65.90	0.845	1.035
1.255	29.3	51	27.76	84.00	0.348	0.426	1.575	52.7	115	54.13	66.30	0.853	1.044
1·260 1·265	29·7 80·2	52 53	28·22 28·69	84·57 85·14	0.356 0.363	0·435 0·444	1.580 1.585	53·3	116 117	54·46 54·80	66·71 67·13	0·861 0·869	1·054 1·064
1.270	80.6	54	29.15	35.71	0.870	0.454	1.590	53.6	118	55.18	67.59	0.877	1.075
1.275	81.1	55	29.62	36.29	0.377	0.462	1.595	53.9	119	55.55	68.05	0.886	1.085
1.280	81.5	56	80.10	86.87	0.385	0.472	1.600	54.1	120	55.93	68.51	0.895	1.096
1.285	82·0 82·4	57	80.57	87.45	0.898 0.400	0.481	1.605	54.4	121	56.80	68.97	0.904	1.1107
1 290 1 295	32.8	58 59	81·04 81·52	88·03 88·61	0.408	0·490 0·500	1.610 1.615	54·7 55·0	122 128	56·68 57·05		0.918 0.921	1·118 1·128
1.300	88.8	60	81.99	89.19	0.416	0.510	1.620	55.2	124	57.40	70.32	0.980	1.139
1.305	33.7	61	82.46	89.77	0.424	0.519	1.625	55.5	125	57.75		0.988	1.150
1.810	34.2	62	32.94	40.85	0.432	0.529	1.630	55.8	126	58.09	71.16	0.947	1.160
1·815 1·320	84·6 85·0	68 64	88·41 88·88	40.93	0·439 0·447	0·538 0·548	1.635 1.640	56·8	127 128	58·48 58·77	71.57	0.955 0.964	1·170 1·181
1.825	85.4	65	84.85	42.08	0.455	0.557	1.645	56.6	129	59.10		0.972	1.192
1.330	85.8	66	34.80	42.66	0.462	0.567	1.650	56.9	180	59.45	72.82	0.981	1.202
1.335	36.2	67	85.27	48-20	0.471	0.577	1.655	57.1	181	59.78	78.23	0.989	1.212
1·840 1·845	86.6	68	85.71	48.74	0.479	0.586	1.660	57.4	182	60.11	78.64	0.998	1·222 1·238
1.850	87.0 87.4	69 70	86·14 86·58	44.82	0.486	0-596 0-605	1.665 1.670	57·6 57·9	188 184	60·46 60·82		1.007	1.244
1.855	87.8	71	87.02		0.502	0.614	1.675	58.2			74.97	1.025	1.256
1.860	88.3		87.45	45.88	0.209		1.680	58.4	186	61.57	75.42	1.084	1.267

s.g. $\frac{15^{\circ}}{4^{\circ}}$	Degrees Besumé	Degrees Twaddell	we	rts by ight tain	1 litre contains kilos.		
in vacuo	ÄÄ	P P	P.c. 80,	P.c. H,804	80.	H_80_	
1.685	58.7	137	61.93	75.86	1.043	1.278	
1.690	58.9	138	62.29	76.30	1.053	1.289	
1.695	59.2	139	62.64	76.73	1.062	1.301	
1.700	59.5	140	63.00	77.17	1.071	1.812	
1·705 1·710	59.7	141 142	63·35 63·70	77.60 78.04	1.080 1.089	1·823 1·334	
1.715	60·0 60·2	143	64.07	78.48	1.099	1.346	
1.720	60.4	144	64.43	78.92	1.108	1.357	
1.725	60.6	145	64.78	79.36	1.118	1.369	
1.730	60.9	146	65.14	79.80	1.127	1.381	
1.735	61.1	147	65.50	80.24	1.136	1.392	
1.740	61.4	148	65.86	80.68	1.146	1.404	
1·745 1·750	61·6 61·8	149	66.22	81·12 81·56	1·156 1·165	1·416 1·427	
1.755	62.1	150 151	66.94	82.00	1.175	1.439	
1.760	62.8	152	67.30	82.44	1.185	1.451	
1.765	62.5	153	67.65	82.88	1.194	1.463	
1.770	62.8	154	68.02		1.204	1.475	
1.775	63.0	155	68.49	83.90	1.216	1.489	
1.780	63.2	156	68.98		1.228	1.504	
1.785	63.5	157	69.47		1·240 1·252	1·519 1·534	
1·790 1·795	63·7	158 159	70.45	86.30	1.265	1.549	
1.800	64.2	160	70.94		1.277	1.564	
1.805	64.4	161	71.50		1.291	1.581	
1.810	64.6	162	72.08		1.305	1.598	
1.815	64.8	163	72.69		1.319	1.621	
1.820	65.0	164	73.51		1.338	1.639	
1.821	-	-	73.63		1.341	1.643	
1·822 1·823	65.1	-	73·80 73·98		1·345 1·348	1.647 1.651	
1.824	65.2		74.12		1.352	1.656	
1.825	00 2	165	74.29		1.356	1.661	
1.826	65.3	_	74.49		1.360	1.666	
1.827	_		74.69	91.50	1.364	1.671	
1.828	65.4	-	74.86		1.368	1.676	
1.829	_	-	75.03		1.372	1.681	
1·830 1·831	05.5	166	75.19		1.376	1.685	
1.832	65.5		75.58		1.384	1.695	
1.833	65.6		75.72		1.388	1.700	
1.834	_	-	75.96		1.393	1.706	
1.835	65.7	167	76.27	98.43	1.400	1.713	
1.836	-	-	76.57		1.406	1.722	
1.837	-	-	76.90			1.780	
1·838 1·839	65.8	_	77.28		1·419 1·426	1·789 1·748	
1.840	65.9	168	78.04			1.759	
1.8405		-	78.35		1.441	1.765	
1.8415		_	79.18	97.00		1.786	
1.8410		-	79.76	97.70	1.469	1.799	
1.8415		-	80.16	98.20	1.476	1.808	
1.8400	1	-	80.57		1.488	1.816	
1·8400 1·8395		-	80.98		1·490 1·494	1.825	
1.8390			81.38		1.494	1.834	
1.8385	1	_	81.59		•	1.838	
			1 0.	1 30 30	1	1 - 55	

For the properties of fuming H₂SO₄ v. Pyrosuppure Acrd, p. 625.

For a test for small quantities of free H₂SO₄ v. Carey Lea (Am. S. [3] 45, 478; abstract in C. J. 64 (ii.), 566).

Reactions. -1. Conc. H.SO, is decomposed

by heat; at c. 440° the products are SO, and H₂O, and by passing through a red-hot tube SO₂, O, and H₂O are formed (Deville a. Troost, C. R. 56, 891; Wanklyn a. Robinson, Pr. 12, 507; Redwood, Ph. [2] 5, 601; Deville a. Debray, A. Ch. [8] 61, 125).—2. Decomposed by the electric current; with cone. H₄SO₄ and a raised temperature O separates at the positive pole, and H and S at the negative; at 00 only H and O are given off (Faraday, T. 1834; Warburg, P. 135, 114; Luckow, Fr. 1880. 1; Berthelot, C. R. 86, 71; 90, 269). Gladstone a. Tribe (C. J. 85, 176; 43, 845) found that a little SO, was formed when the electrolysis of 98.2 p.c. H₂SO₄ was very slow. According to the concentration of the H2SO4Aq electrolysed, the strength of the current, and the current density, ozone, H_2O_2 , and S_2O_7 are formed, besides H and O. For an examination of the conditions favourable to the production of ozone v. McLeod (C. J. 49, 591, where references are given to earlier memoirs). cording to Bouty (C. R. 108, 393), acid of the concentration H₂SO₄+H₂O always gives H₂O₂ on electrolysis; Richarz (A. Ch. [2] 24, 183) says that the maximum production of H2O2 is obtained with acid of the concentration H₂SO₄+1.347H₂O; with acid more dilute than 60 p.c. H₂SO₄ very little H₂O₂ is obtained, but considerable quantities of S2O, and ozone (Bouty, l.c.; cf. Kuriloff, J. R. 23, 235). In connection with the electrolysis of H2SO4Aq, and the conditions of production of H and O only, or of H.O., ozone, and S.O., also, v. Armstrong (C. J. Proc. 1893-4. 188 [No. 127]).—3. By passing a mixture of the vapour of H2SO4 mixed with hydrogen through a red-hot tube, H2O is obtained along with SO₂, S, or H₂S, according to the quantity of H (v. Warner, C. N. 28, 13). Hydrogen occluded by palladium or platinum partially reduces 98.2 p.c. H₂SO₄, giving SO₂ (G. a. T., l.c. p. 177).—4. SO₂, CO₂, and H₂O are formed by heating conc. H₂SO₄ with carbon; and SO2 and H2O by heating with sulphur .-5. Phosphorus takes fire in boiling H.SO, with separation of S; red phosphorus reacts with the conc. acid when hot, forming SO2 and H2PO4.-6. Chlorine reacts with conc. H2SO, to form Cl(SO₂.OH) (v. CHLOROSULPHONIC ACID, p. 509). 7. Most metals interact; K and Na give off H, Zn and Fe and several other metals also give off H with more or less dilute H2SO,Aq at ordinary temperatures; on raising the temperature SO₂ and H₂S are generally evolved and S is often separated. Some metals, such as Sb, Bi, Cu, Hg, react only with warm fairly conc. H2SO4, giving off SO₂, and often also H₂S. Au, Ir, Pt, Rh, and W do not interact with H, SO. Regarding the conditions of the interaction of H.SO.Aq with metals—concentration, temperature, and products—v. Ditte (A. Ch. [6] 19, 68). Pattison Muir a. Adie (C. J. 53, 47) found that the only products of the interaction of approximately pure zinc and dilute H2SO,Aq (c. H2SO,12H2O to c. H₂SO₄.15H₂O) were ZnSO₄Aq and H; that with conc. acid (c. H₂SO₄.H₂O to c. H₂SO₄) SO₂ was the chief gaseous product, and that H2S was also produced at higher temperatures; and that with acid of intermediate concentration (c. H₂SO₄.2H₂O) both SO₂ and H₂S were formed in considerable quantities at c. 160°. For experiments on the rate of action of H.SO.Aq on different specimens of zinc v. Divers a. Shimidzu (C. J. 47, 598), who have also examined the production of NH₂OH by the interaction of zinc with H₂SO₄Aq and HNO₂Aq together (cf. HYDROXYLAMINE, vol. ii. p. 784). For qualitative experiments on the interaction of H2SO4 and tin, v. Pattison Muir a. Robbs (C. N. 45, 69, where references to other memoirs are given). Pullinger (C. J. 59, 815) found that pure sinc was not acted on by H.SO, Aq (1 part conc. acid + 8 parts water by weight) provided the acid had been boiled for a long time before use and the surface of the zinc was quite smooth; if the surface was rough some of the zinc was dissolved. P. also found that oxidisers increased the rate of solution of zinc, and reducers, if not containing S, almost stopped the solution. P. concluded that when the surface of the zinc was rough S2O, was probably formed, and then the action proceeded. Weeren (B. 24, 1785) noticed that there was scarcely any reaction between pure zinc and H2SO4Aq (c. 20 parts water); he supposed that a layer of H was condensed on the surface of the metal. W. thought that with impure zinc the H is given off from the surface of the more negative metal present, and so the surface of the zinc is left exposed to the action of the acid. W. found that the evolution of H proceeded more rapidly in vacuo, and that it was hastened by brushing the surface of the metal. The weight of pure zinc dissolved in 80 mins. was 2·1 mgm. at 0°, 9·3 mgm. at 98°, and 122·1 mgm. at the b.p. of the acid used. W. obtained similar results with A1, Cd, Co, and Fe. Sodium amalgam or zinc amalgam, with conc. H2SO4, gives off H, then quantities of H_2S , then SO_2 and S, and finally only SO_2 (Walz, C. N. 23, 245). The interaction of H_2SO_4 and copper has been fully examined by Pickering (C. J. [2] 18, 112; cf. Copper, vol. ii. p. 253).—8. H₂SO₄ reacts with the oxides of nitrogen (not with N₂O) to form NO₂(SO₂.OH) (v. NITROSULPHONIC ACID, p. 601).—9. Phosphoretted hydrogen reacts slowly with H₂SO₄ at the ordinary temperature, giving SO₂, S, and H₂PO₄ (Rose, P. 24, 189).—10. With sulphuretted hydrogen, H₂O₅, S, and SO₂ are formed.—11. Phosphorus pendoxies withdraws H₂O from hot conc. H₂SO₄, producing P₂O₃.H₂O and SO₃ (Barreswil, C. R. 25, 30; Evans, Ph. 8, 127); phosphorus pentachloride or trichloride forms SO₂ and POCl₃, also SO₂Cl₂ when PCl₃ is used (Williamson, P. M. [4] 7, 365; Schiff, A. 202, 111; cf. Michaelis, J. Z. 6, 233, 292), and also Cl(SO₂OH) (Müller, B. 6, 277; Geuther, B. 5, 925; Thorpe, C. J. 37, 358).—12. H₂SO₄ reacts with most self-to-form sulphotes and set free with most salts to form sulphates, and set free the acid of the salt if that acid is more volatile than H₂SO₄. In several cases secondary reactions occur between the acid set free and the H2SO4; thus, with bromides and iodides, Br and I, and SO, or H,S, are produced. The interaction of H2SO.Aq and KBr has been examined by Addyman (C. J. 61, 94), who found that the percentage of HBr decomposed varied almost directly with the quantity of H₂SO₄ used, and that when so much H₂SO₄ was present that the water produced in the reaction $2HBr + H_2SO_4 = 2H_2O + SO_2 + Br_2$ caused no sensible dilution of the H₂SO₄, all the HBr was decomposed; with dilute acid, c. 80 p.c. H₂SO₄, using excess of acid, scarcely a trace of HBr was decomposed (cf. Feit a. Kubierschki,

J. Ph. [5] 24, 159). Jackson (C. J. 43, 889) finds that the interaction of conc. H₂SO₄ and KI varies according as the acid is present in large excess, or in quantity just sufficient for the KI used. In the first case all the I of the KI is set free along with an equivalent quantity of SO₂; in the second case all the I is liberated with the equivalent quantity of H₂S. Many salts of acids containing much oxygen, e.g. dichromates and permanganates, are decomposed by hot conc. H_aSO_4 with evolution of O. The reaction of H_aSO_4Aq with $KMnO_4Aq$ has been examined by Gooch a. Danner $(Am.\ S.\ [3]$ 44, 301), who found that the quantity of O given off increased with (1) concentration of H2SO,Aq, (2) time, (8) temperature; no O was given off after a few hours with 20 p.c. acid at the ordinary temperature, but a considerable quantity of O was produced after five days' action, and c. the same result was obtained by heating to 80° for one and a half hours (cf. Jones, C. J. 33, 95).-13. Conc. H₂SO₄ reacts with many oxygenated organic compounds, withdrawing H₂O; with benzenoid hydrocarbons it forms sulpho- acids; many organic compounds, such as sugar, starch, &c., are charred by warm H.SO₄.—14. Pictet (C. R. 115, 708, 814) has examined the reactions of sulphuric acid at low temperatures, using acid of 89 p.c. H2SO4; no action occurred below -125° with such bodies as NaOH, NaCl, NH, Na₂CO₂, Na, K; in most cases action began suddenly at temperatures varying from -120° to -60°.—15. H.SO, combines with water; the H₂SO₄.H₂O₅ H₂SO₄.2H₂O₅ $H_2SO_4.8H_2O$ have been isolated as solids (v. infra).—16. Conc. H₂SO₄ combines with sulphur trioxide to form H₂SO₄.SO₅ and other compounds (v. Pyrosulphuric acid).

For the reactions of fuming H₂SO₄ v. Pyrosulphuric acid. (p. 625). The salts of H₂SO₄ are described under Sulphates (p. 567). Both amides of H₂SO₄, viz. SO₂OH.NH₂ and SO₂(NH₂)₂, have been isolated (v. Sulphamic acid., p. 567, and Sulphamide, p. 567); the imide SO₂.NH is also known, v. Sulphimide, p. 587; cf. also Sulphimide. NH(SO₂.NH₂)₂, p. 587, and Imido-amide NH(SO₂.NH₂)₂, p. 600.

Constitution of sulphuric acid.—The reactions of H₂SO₄ with Cl and with PCl₃, and the

Constitution of sulphuric acid.—The reactions of H₂SO₄ with Cl and with PCl₃, and the reactions of the products with water, also the formation of SO₂.OH.NH₂, SO₂(NH₂)₂, and SO₂.NH, show that H₂SO₄ contains two OH groups. The fact that only one compound, SO₂.OEt.OMe, can be formed from H₂SO₄ shows that the two OH groups in SO₂(OH)₂ are related to the rest of the molecule in the same way; hence the formula for the acid is to be written

 $S < _{0.0H}^{0.0H}$ or $O_2 S < _{0H}^{0H}$. Inasmuch as $SO_2 Cl_2$

is formed from SO₂(OH)₂ by the reaction of POl₃, as certain hydrocarbons interact with SO₂Cl₂ to form SO₂R₂ (where R = Me, Ph. &c.), and as these compounds SO₂R₂ are reduced to SR₂, where the radicles R are certainly in direct union with S, it is extremely probable that the OH groups in SO₂(OH)₂ are in direct union with

S, and hence the formula O₂S OH is to be pre-

ferred to S < 0.0H for sulphuric soid. It is

has the constitution O.S.O or S

HYDRATES OF SULPHURIC ACID. Two solid hydrates have been isolated; a liquid hydrate has probably also been isolated.

Monohydrate H2SO4.H2O (Tetrahydroxysulphuric acid SO(OH), Large, colourless, six-sided tablets; formed by mixing 1 part conc. H,SO, with rather less than one-fifth part water (by weight), and surrounding with a freezing mixture. The crystals melt at 7.5° (Pierre a. Puchot, A. Ch. [5] 2, 164). The hydrate readily remains liquid considerably below its m.p. S.G. $\frac{0^{\circ}}{70} = 1.7943$ (for the liquid) (Mendeléeff, B. 19, 380); 1.77806 at 15°, 1.77423 at 20°, 1.77071 at 25° (Perkin, C. J. 49, 777; determinations made with liquid, and referred to water at same temperatures). Thomsen gives H.F. from liquid H₂SO₄ and H₂O [H²SO⁴,H²O] = 6,379 (Th. 3, 54); Berthelot gives 6,120 (C. R. 78, 716). Heat of fusion = 3680 g. units (for 116 g.) H₂SO₄·H₂O (Berthelot, C. R. 78, 716). For vapour pressures from 143° to 204° v. Tate (P. M. [4] 26. 502)

Dihydrate H₂SO₄.2H₂O (Perhydroxysulphuric acid S(OH)₆). Formed by adding 368 parts H₂O to 1 part pure H₂SO₄, by weight. This hydrate has not been obtained as a solid; the S.G. of the liquid at $\frac{00}{40}$ is given by Mendeléeff (B. 19, 380) as 1.6655 (cf. Perkin, C. J. 49, 777).

It is stated (v. Mendeleeff, B. 19, 388) that the maximum contraction when H₂SO₄ and H₂O are mixed occurs when the compounds are present in the ratio H_2SO_4 :2 H_2O_7 , but Pickering (C.J.57, 84) says that the maximum does not correspond with the composition of any hydrate the existence of which is indicated in other

Tetrahydrate H.SO.4H.O. Obtained by Pickering (C. J. 57, 339) as a crystalline solid, melting at -25°, by cooling H₂SO₄Aq containing 57.6 p.c. H₂SO₄ (H₂SO₄.2H₂O corresponds with 57.66 p.c. H₂SO₄). The crystals are large and well defined, and resemble H₂SO₄.H₂O; the compound may be kept liquid at 20° to 25° below its freezing point.

Hydrates of sulphuric acid in aqueous solution.—According to Mendeléeff (B. 19, 380; cf. his Principles of Chemistry, vol. ii. p. 234), an examination of the connections between the concentration of H2SO, Aq and the S.G. indicates the existence of hydrates that have not yet been isolated. Pickering (C. J. 57, 64) comes to the same general conclusion from an extended examination of the connections between the changes of composition of H2SO, Aq and the changes of S.G., heat capacity, electrical conductivity, heat of solution, and expansion by heat of the solution. Pickering's reasoning is a special case of that explained in the article Solutions (part ii. p. 492); for the detailed argument, and criticisms of it, v. Pickering, C. J. 57, 64; Rücker, P. M. [5] 32, 304; 33, 204; P., P. M. [5] 38, 132, 463; cf. Mendeléeff, Z. P. C. 1, 275; and Crompton, C. J. 53, 116.

PERSULPHURIC ACID H.S.O.Aq. This acid is present in the solution obtained by elec- | PEROXIDE, p. 616. Vot. IV.

undecided whether the group SO₂ in O₂S.OH.OH | trolysing H₂SO₄Aq of certain concentrations;
O | Berthelot (C. R. 90, 269; 112, 1481) obtained a quantity of the acid equal to from 88 to 123 g. S,O, per litre by electrolysing H,SO,Aq containing water and acid in the ratio 10H, O:H, SO, (cf. SULPHURIC PEROXIDE, p. 616; and SULPHURIC ACID, Reactions, No. 2, p. 623). According to Traube (B. 22, 1518, 1528) a solution of H₂S₂O₈ in H₂SO₄ does not oxidise H₂C₂O₄, HNO₂, CO, or Cr₂(SO₁)₃; Richarz, however (B. 21, 1670), says that oxalic and nitrous acids are oxidised. The acid has not been isolated; by electrolysing at temperatures below 0°, and then adding BaOAq or KOHAq, the Ba or K salt is obtained. The formula H₂S₂O₈ is given rather than HSO₄, because of the results of Bredig's measurements of the conductivities of solutions of the K salt. For salts of this acid v. Persulphates, p. 582.

> PYROSULPHURIC ACID H2S2O, sulphuric acid). A solution of this acid in Nordhausen acid. This acid is obtained by adding rather more SO, to H SO, than the proportion SO,:H,SO,, cooling, and spreading out the crystals that form over nearly conc. H₂SO₄ under a bell-jar till the excess of SO₃ is removed (Schultz-Sellack, B. 4, 109). Forms large crystals, melting at 35°. Easily decomposed by heat to H₂SO₄ and SO₃. H₂S₂O₇ is dibasic, forming salts MHS₂O₇ and M₂S₂O₇ (v. PYROSULPHATES, p. 583). Funing sulphuric acid is a solution of H₂S₂O₂ in H₂SO₄; it is prepared by heating dry FeSO, in fireday vessels and collecting the oily, fuming liquid that distils over. The acid fumes strongly in the air, and gives off SO, when warmed. But few of the reactions described for pyrosulphuric acid, or for fuming sulphuric acid, can be set down as belonging to a definite compound, but rather to solutions of H.S.O, (and perhaps other compounds of H₂SO, and SO₂) in H₂SO₄. Finely-divided silver dissolves easily; the solution poured into water gives Ag, SO, and SO₂; mercury reacts similarly; spongy copper has a slight reaction, giving Cu, S and CuSO₄ (Divers a. Shimidzu, C. J. 47, 636). Distilled with phosphorus pentoxide in a stream of hydrogen chloride, Cl(SO2.OH) is formed; the same acid is formed by heating with PCl, and also by heating 'crystallised fuming sulphuric acid' with HCl (v. Chlorosulphonic acid, p. 509). Nitrogen dioxide passed into well-cooled pyrosulphuric acid gives NO₂(SO₂OH) (v. Nitrosulphunic Acid, p. 601). By saturating ordinary conc. H₂SO₄ with SO₂ (Jacquelain, A. Ch. [3] 30, 348), or by cooling fuming H₂SO₄ to -10° (Wacken-roder, Ar. Ph. [2] 87, 267), the compound H₂S₂O₂2H₂SO₄ (= 3H₂SO₄SO₃) is said to be ob-tained in transparent crystals. Schultz-Sellack (B. 4, 109) failed to obtain any compound of H₂SO₂ and SO₃, except H₂SO₂. Crystals of the composition H₂SO₂.3SO₂ (= H₂S₁O₁₂) melting at 8°-10° are said by Weber (P. 159, 313) to be formed by adding cooled conc. H,SO,, little by M. M. P. M. little, to SO.

> SULPHURIC ANHYDRIDE SO, v. Sul-PHUB TRIOXIDE, p. 615.

> Persulphuric anhydride S.O. v. Sulphuric 88

SULPHURIC-NITROSYL CHLORIDE

SO. NOCl v. NITROSULPHONIC CHLORIDE, p. 601. SULPHURIC OXIDE SO, v. SULPHUR TRIoxide, p. 615.

SULPHURIC PEROXIDE S2O, v. p. 616. SULPHUROSAMIC ACID SO.OH.NH, v. THIONAMIC ACID, p. 695.

SULPHUROUS HYPOSUL-ACID PHUROUS ACID.

SULPHUROUS ACID H2SO,Aq. The acid has not been isolated; a solution of SO₂ in water

reacts as a weak dibasic acid, forming salts of the types M'HSO₃, M'₂SO₃, and M'¹SO₃ (v. Sulphites, p. 587). H.F. [H',S,O',Aq] = 147,130; $[H^2O,S,O^2,Aq] = 78,780$ (from rhombic S; Th.

The methods of formation and preparation of H.SO.Aq are sufficiently detailed under Sur-PHUR DIOXIDE (p. 613); it is only necessary to

add water to the SO2 obtained.

Properties.—An aqueous solution of H.SO. readily decomposes, giving off SO2; on standing for some time in the air, and more quickly by warming, H₂SO₄Aq is formed with production of much heat; [H²SO³Aq,O] = 63,634 (Th. 2, 253). H2SO8Aq is altered by the action of light, so that after exposure to light for some time the solution ppts. Ag₂S from AgNO₃Aq, and reduces silver haloid compounds, forming some Ag S; the solution after exposure to light behaves like a dilute solution of a polythionic acid (Stas, Chem. Proport. 64). Löw (L'Institut, 1873. 67) says that dilute H,SO,Aq is changed to H,SO,Aq and S by long exposure to light.

Reactions.—1. Heated in a closed tube to 170°-180°, H2SO, Aq and S are formed, and after a time also H₂S (Geitner, A. 129, 350).—2. Decomposed by an electric current, giving H₂SO₄Aq at the positive pole and S at the negative; with a weak current H₂S₂O₄Aq is probably formed at first at the negative pole (Guéroult, C. R. 85, 225).-3. Phosphorus pentachloride produces SOCl, and POCl_s (Kremers, A. 70, 297; Schiff, A. 102, 111).—4. Sulphurous chloride with H₂SO₂Aq produces polythionic acids (v. Debus, C. J. 53, 345).—5. Vapour obtained by heating H2SO3Aq reacts with chlorine, when passed over red-hot Pt, to form Cl(SO2.OH) (v. CHLORO-SULPHONIC ACID, p. 599).-6. NO2(SO2.OH) (v. NITROSULPHONIC ACID, p. 601) is formed by the interaction of moist SO₂, i.e. H₂SO₂Aq, with nitric oxide or nitrogen dioxide.—7. H₂SO₂Aq is oxidised to H₂SO₄Aq by oxygen, the halogens, and many other oxidisers.—8. Heated with phosphorus in a closed tube to 200°, H2S is produced (Oppenheim, Bl. [2] 1, 163).—9. Reduced by sinc and dilute sulphuric acid to H₂S (v. Reinsch, D. P. J. 163, 286; 181, 832); with zinc alone, H2SO,Aq is partly reduced, probably giving H.S.O.Aq (v. HYPOSULPHUBOUS ACID, p. 627).-10. Many metals which evolve H from H2SO, Aq or HClAq react with H2SO, Aq to form sulphites and sulphides, with evolution of little or no H (v. Federoff, Chem. Zeitung, 5, 15). For a fairly full qualitative examination of the interaction of Al, Od, Mg, Ni, and Zn with H₂SO₂Aq, v. Schweitzer (C. N. 23, 293).—11. Many of the oxyacids of the halogens are reduced to hydracids by H₂SO₂Aq. — 12. Phosphorous acid gives H₂PO₂Aq and H₂S; then S, and thionic acids (Wöhler, A. 89, 252; Maquenne, Chem. Zeitung.

1890 [1] 511).—13. Phosphoretted hydrogen produces H₃PO₄Aq, and S (Cavazzi, G. 16, 169). 14. Hydrogen sulphide passed into H2SO3Aq forms a solution containing H₂SO₄, H₂S₃O₆, H₂S₄O₆, probably H₂S₅O₆, S in suspension, and a colloidal form of S in solution; if the passage of H.S is continued until reaction ceases the final products are S and H₂O (Debus, C. J. 53, 282; cf. Thionic acids, p. 698).-15. Many metallic salts are reduced, in aqueous solutions, by H2SO3Aq; salts of Ag and Hg to the metals, ferric to ferrous salts, &c. According to Divers a. Shimidzu (C. J. 49, 575), the reduction does not consist in the withdrawal of O by H,SO,Aq and the oxidation of that acid thereby to H.SO.Aq. The first step is looked on by D. a. S. as probably the formation of a sulphite of the metal of the salt, and this is followed by either the decomposition of this sulphite by water to metal and H2SO, Aq, or the reaction of the sulphite with some of the original, unchanged, salt to form metal and a persulphite (or -ic sulphite) of the metal.—16. With potassium thiosulphate H₂SO₃Aq forms K tri- tetra- and penta-thionates, and S (Debus, C. J. 53, 343).— 17. H2SO3Aq dissolves many freshly ppd. metallic sulphides, e.g. MnS, FeS, ZnS, forming thiosulphates (Schönbein, P. 104, 300; Heldt, J. pr. 83, 20).—18. Alkali nitrites give salts of nitrilosulphonic acid N(SO₂OH)₃ (q. v., p. 601), and also salts of oxy-imidosulphonic acid $N(OH)(SO_2OH)_2$ (q. v., p. 602). Using a dilute, ice-cold solution of H_2SO_3 (= 5 p.c. SO_2), and adding 40 g. KNO, in 1,000 c.c. water to 3 litres of this H,SO,Aq, Raschig (A. 241, 234) obtained $NH_2(SO_2.OK)$ (cf. Sulphamic acid, p. 567). Under certain conditions of concentration and temperature NO and N₂O, also NH₃ and sometimes NH₂OH, are produced. With nitrous acid the final products are NH₂, NO, N₂O, and H₂SO₂Aq; according to Raschig (l.c.), NH_.(SO_.OH) is always formed, and the gaseous products are to be looked on as resulting from secondary reactions between nitrous acid and NH2(SO2OH) or other nitrogenised sulphonic acids formed in the primary reaction between the nitrous and sulphurous acids. With silver and mercurous nitrites Divers a. Haga (C. J. 51, 659) found that sulphites were produced, along with some NO and H₂SO, and more or less NH OH.-19. For the reaction of H₂SO₃Aq with potassium platinochloride v. Platino-sulphonates, this vol., p. 285.

Neither the amide SO(NH2)2 (v. the heading THIONAMIDE), nor the imide SO.NH, of H.SO. has been isolated; and it is doubtful whether the acid SO.OH.NH₂ or SO₂.NH₂.H is known v. THIONAMIC ACID.

Constitution of sulphurous acid. The fact that many haloid derivatives of hydrocarbons interact with NaHSO, to form sulphonic acids $e.g. EtI + NaHSO_s = EtSO_tH + NaI$ —makes it probable that the Na in NaHSO, is in direct union with the S, because the S in sulphonic acids is almost certainly in direct union with the hydrocarbon radicle; hence it is likely that one H atom in H.SO, is directly combined with the S atom, and hence the formula for H₂SO₂ is probably H.SO₂.OH. This view of the constitution of H_2SO_2 is insisted on, and liustrated, by Divers (C. J. 47, 205).

Hydrates of sulphurous acid. Colourless, ice-like crystals are obtained by passing SO, into water surrounded by ice, by cooling saturated SO, Aq, or by passing moist SO, through a tube kept at a low temperature. After filtering quickly in a covered filter, and pressing between filter paper at 8°, the crystals have the composition H₂SO₂.xH₂O; Dopping (Petersburg Acad. Bull., 7, 100) found x = 10, Pierre (A. Ch. [3] 23, 416) found x=8, Schönfeld (A. 95, 22) found x = 14, Roozeboom (R. T. C. 3, 29, 59, 75, 84; 4, 65) found x=6, and this was confirmed by Geuther (A. 224, 218). Roozeboom (l.c.) gives S.G. of the crystals as 1.21 (referred to water at 4°); Geuther (l.c.) gives 1.147 as the S.G. after melting at 14°, and 14° as the m.p. with separation of some liquid SO, under the layer of SO,Aq. R. (l.c.) gives the following dissociation-pressures, which are independent of the quantity of the hydrate: 303 mm. at 0°, 754 mm. at 7.05°, 1177 mm. at 10°, 1773 mm. at 12·1°. According to R. the critical temperature of decomposition is 7.1° in an open vessel and 12.1° in a closed

HYPOSULPHUROUS ACID H₂S₂O₄Aq. This acid has not been isolated. An aqueous solution is probably formed by decomposing the Na salt by dilute H₂SO₄Aq or H₂C₂O₄Aq, and also perhaps by digesting H,SO,Aq with zinc clippings till the liquid is brownish red; but neither the aqueous solution of the acid, nor any of its salts, has been obtained free from impurities. For preparation of the sodium salt, and for references, v. Hyposulphites (p. 592).

M. M. P. M. SULPHUROUS ANHYDRIDE v. SULPHUR pioxide, p. 613.

SULPHUROUS OXIDE v. SULPHUR DIOXIDE,

p. 613.

SULPHURYLAMIDE v. SULPHAMIDE, p. 567.

SULPHURYL BROMIDE v. SULPHUR OXY-BROMIDES, p. 617.

Pyrosulphuryl bromide v. SULPHUR OXY-

BROMIDES, p. 617.

SULPHURYL CHLORIDE v. SULPHUR OXY-CHLORIDES, p. 618.

Nitrosulphuryl chloride v. Nitrosulphonic CHLORIDE, p. 601.

Pyrosulphuryl chloride v. Sulphur oxy-

SULPHURYL HYDROXYLCHLORIDE

CHLOROSULPHONIC ACID, p. 599. SULPHURYL HYDROXYLFLUORIDE

FLUOSULPHONIC ACID, p. 600. SULPHURYLIMIDE v. SULPHIMIDE.

587. SULPHURYL IMIDO-AMIDE v. SULPHIMIDO-

AMIDE, p. 587. SULPHYDRATES v. HYDROSULPHIDES, vol. ii.

p. 733.

SULPHYDRIC ACID v. Hydrogen sul-

PRIDE, vol. ii. p. 725. SULPHUVINURIC ACID C₄H₄N₂SO₂, i.e. S-C(NH₂) CH:C(CO₂H) N (Strende, A. 261, 26). Formed from di-bromo-pyruvic acid and a conc. solution of thio-urea (Nencki a. Sieber, J. pr. [2] 25, 74). Crystals (containing 2aq), sl. sol. cold water. Reduces Fehling's solution in the cold. FeCl. gives a deep violet colour.—Salts: MgA'2ZnA': minute needles.—HA'HCl.—HA'HBr.— HA'HNO, aq: needles, sl. sol. water.

SULPHYDRO-ACETIC ACID v. THIOGLY-COLLIC ACID.

SULPHYDRO-ACETONE CH,.CO.CH,.SH.

Formed from chloro-acetone and NaSEt (Autenrieth, B. 24, 165). Vialde a phase!

[36°]. Phenyl derivative PhS.CH, Ac. (266°). Formed from CH₃.CO.CH₂Cl and NaSPh in alcohol in the cold. Crystals. Yields a phenyl-hydrazide [87°] (A., cf. Delisle, A. 260, 250). Combines with KHSO, and NaHSO.

SULPHYDRO-ACETOPHENONE. Phenyl derivative C, H, SO, i.e. [53°]. C₆H₅.S.CH₂.CO.C₆H₅. Formed from ω-bromo-acetophenone and NaOPh (Delisle, B.

22, 309). Needles (from alcohol), v. sol. ether

and acetone

SULPHYDRO-ACETYL-PROPIONIC ETHER.

 $\begin{array}{l} P\,h\,en\,y\,l\,deriv\,a\,tiv\,e\,\,C_{13}H_{14}SO_3,\,\emph{i.e.}\\ C_6H_5.S.CH_2.CO.CH_2.CH_2.CO_2Et. \end{array} \eqno(19)$ (197° at 15 Formed from bromo-acetyl-propionic mm.). ether and NaSPh (Delisle, B. 22, 309)

SULPHYDRO-ALLYL-QUINAZOLINE. Di- $C_6H_4 < \stackrel{CH_r.N.C_3H_5}{\sim} .$ [91°]. Formed hydride from ω-oxy-tolyl-allyl-thio-urea and HClAq at 100° (Söderbaum a. Widman, B. 22, 1670). Needles, v. e. sol. alcohol and benzene.

Reference. — OXY-SULPHYDRO-ALLYL-QUINAZO-

DISULPHYDRO-DI-ALLYL-TRIAZOLE

NH CS.N.C₃H₅. Formed from allyl-thio-ures. and HNO₂ (Hector, J. pr. [2] 44, 505). Oil.—

B'2H.PtCl.: crystalline pp.
o-SULPHYDRO-BENZOIC ACID

C.H. (SH).CO.H. Thio-salicylic acid. Formed by reducing o-sulpho-benzoic chloride with zincdust and H.SO. (Delisle, B. 22, 2206). Amorphous powder, sl. sol. hot water, v. sol. alcohol. Not volatile with steam .- AgA'

Thio-amide CoH4(SH).CS.NH2. Formed by heating o-oxy-benzamide with P.S. (Spilker, B.

22, 2771). Minute dark-red crystals.

Phenyl derivative CoH, S.CoH, CO2H. Formed from o-diazobenzoic acid, phenyl mercaptan, and NaOHAq at 50° (Graebe a. Schultess, A. 263, 4). Plates (from alcohol) or needles (from HOAc); sl. sol. water.—NH,A.

—KA': needles, v. sol. water.—EtA'. [151°].

m-Sulphydro-bensoic acid C_sH,(SH).CO_sH.

[147°]. Formed by the action of tin and HClAq on C,H,(SO,CI).COCI (Frerichs, B. 7, 793).
Colourless plates, m. sol. water.—BaA', 2½aq.—HgA',.—PbC,H,SO, 3aq.—Cu(OH)A'.—AgA'.
a-SULPHYDRO-BUTYRIC ACID

CHEt(SH).CO2H. Formed from a-bromo-butyric acid and alcoholic KSH (Duvillier, Bl. [2] 30,

507). Sol. water, alcohol, and ether. Sulphydro-isobutyric acid HS.CMe. CO.H.

Formed from a-bromo-isobutyric ether by successive treatment with alcoholic K,S and alcoholic KOH (Lovén, J. pr. [2] 83, 109). a-SULPHYDRO-CINNAMIC ACID

C₆H₅.CH:C(SH).CO₂H. [119°]. Formed, together with sulphocyanic acid, by heating benzylidene-rhodanic acid C.H.:CH:C(SH).CO.S.CN with baryta-water (Ginsburg a. Bondzynski, B. 19, 128; M. 8, 850). Yellowish crystals. V. sol. alcohol, ether, benzene, and CS₂, nearly insol. water. By treatment with iodine in alcoholic solution it is oxidised to di-sulphido-di-cinnamic

acid S₂(C₂HPh.CO₂H)₂.

Reference.-NITRO-SULPHYDRO-CINNAMIC ACID. 8-SULPHYDRO-CROTONIC ACID. Ethyl CH₃.C(SEt):CH.CO₂H. [113°]. derivative Formed from \$\beta\$-chloro-crotonic acid [94.5°], mercaptan, and Na (Autenrieth, A. 254, 235; B. 20, 3189). Crystals, insol. water, v. sol. ether and alcohol. Gives a dark-green colour on warming with isatin and H₂SO₄. Yields CH₂.C(SEt):CH₂ (110°) on heating above its melting-point.— BaA'₂ aq.—AgA': silky needles, not decomposed by hot water. Ac.O forms the compound CH, C(SEt):CH.CO.OAc, a brownish oil.

Isomeride of the ethyl derivative CH₂.C(SEt):CH.CO₂H. [92°]. S. (alcohol) 20. Formed in like manner from the isomeric βchloro-crotonic acid (A.). Crystals, nearly insol. water. Forms a dark-green liquid on warming with isatin and H2SO4. When heated alone it yields the same ethyl-sulphydro-propylene (110°)

as its isomeride.—BaA', aq: thin plates (from Aq).

Phenyl derivative v. Phenyl-sulphydbo-

CROTONIC ACID

SULPHYDRO-ETHYL ALCOHOL

HO.C2H4.SH. Formed from glycolic chlorhydrin and alcoholic KSH (Carius, A. 124, 258). Oil, v. e. sol. alcohol. Oxidised by HNO, to isethionic acid-Hg(C2H5OS)2. Needles (from warm alcohol).

SULPHYDRO - ETHYL - QUINAZOLINE. Dihydride CeH4 CH2.NEt. [103°]. Formed by warming ω -tolyl-ethyl-thio-urea with HCl (Söderbaum a. Widman, B. 22, 2936). Needles.
—B'HAuCl₄. [118°].—B'₂H₂PtCl₆. [208°].

SULPHYDRO - METHANE DISULPHONIC ACID HS.CH(SO₂H)₂. Formed by boiling the tri-sulphonic acid with HOAc (Albrecht, A. 161, 134). Gives a blue colour with FeCl₃.—K₂A''½aq.

 $-\dot{P}b_s(CHS_sO_s)_2$ 8aq. Small needles.

Sulphydro-methane trisulphonic acid HS.C(SO,H), Formed by boiling CS₂ or CSCl, with K,SO,Aq. K,A''' 2aq: triolinic crystals. S. 2 at 12°. Gives a blue colour with FeCl,.-(NH.), A" aq. Thin plates.

SÜLPHYDRO-METHENYL-AMIDO-CRESOL $\begin{bmatrix} 4 & 1 \\ 2 \end{bmatrix}$ C_eH₂Me<^O_N>C.SH. [217°]. Formed by heating benzene-azo-p-cresol with CS₂ at 200° (Jacobsen a. Schenke, B. 22, 3235). Needles. SULPHYDRO - METHENYL - AMIDO - (a) -

NAPHTHOL $C_{10}H_6 < \stackrel{N}{\bigcirc} C.SH.$ [above 220°]. Formed by heating an alcoholic solution of $S_2(C_{10}H_a, NH_a[1:2])_2$ with CS₂ (Jacobson, B. 21, 2624). Needles. Oxidised by K_3 FeOy₆ to a crystalline disulphide.

Sulphydro - methenyl - amido - (0) - naphthol. [249°]. Formed by heating benzene-azo-(8)naphthol with CS₂ at 250° (Jacobson, B. 21, 414). Needles, sl. sol. benzene. HClAq at 170° yields amido- (β) -naphthol. An alcoholic solution of iodine forms crystalline (C₁₀H₀<\frac{N}{O}>C)₂S₂.

SULPHYDRO . METHENYL . a . AMIDO. WAPHTHYL MERCAPTAN C, H. CN C.SH. hove 240']. Formed by heating a-naphthyl

thiocarbimide with S at 225°, and got also by the action of CS, on di-amido-di-naphthyl disulphide (Jacobson, B. 21, 2625; 24, 1406). Oxidised by Minute rosettes, sol. alcohol.

 $K_{s}FeCy_{s}$ to $(C_{10}H_{s} < N > C)_{2}S_{2} - B'HgCl_{2}$. [210°].

Sulphydro - methenyl - amido - (β) - naphthyl mercaptan [232°]. Formed by heating (β) naphthyl thiocarbimide with S. Minute needles. Yields a methyl ether [74°], crystallising in needles.

SULPHYDRO - METHENYL - o - AMIDO-PHENOL C.H. COSH. Thiocarbamido .

phenol. [196°]. Formed by boiling an alcoholio solution of o-amido-phenol for several days with CS, in a current of H as long as H2S is given off (Dünner, B. 9, 465). Formed also by heating oxy-phenyl-thio-urea, and by heating o-oxyphenyl-allyl-thio-urea with HCl at 130° (Von Chelmicki, J. pr. [2] 42, 442; B. 20, 177). Obtained also by adding dilute alcohol to a mixture of potassium xanthogenate and amido-phenol hydrochloride (Kalckhoff, B, 16, 1825). Needles (from water), sl. sol. cold water. Sol. NH, Aq, crystallising therefrom unaltered.

Reactions.—1. Converted in o-amido-phenol by HClAq at 170°, and by NH, Aq at 200°.--2. Ac_2O yields $C_6H_4 < N > C.SAc [120°]. -3. BzCl$

forms benzenyl - amido - phenol and $C_{\rm e}H_{\star}({\rm NHBz}).{\rm OBz}.{--}4.$ An alcoholic solution of iodine, added to its solution in NaOHAq, forms $(C_0H_4 < N_0) C)_2S_2$ [110°], which is reconverted by hot dilute NaOHAq into sulphydromethenyl-amido-phenol. — 5. Boiling methylaniline forms a base (? C₆H₄<0>C.NPhMe) which yields B'2H2PtCls.

Salt.-AgC, H, NSO. Amorphous. Ethyl ether EtC, H, NSO. (c. 267°). Solid, melting below 35°; volatile with steam.

SULPHYDRO-METHENYL-AMIDO-PHENYL-MERCAPTAN $C_{s}H_{s} < N > C.SH.$

[179°] (H.);

[174°] (J. a. F.).

Formation.-1. By digesting o-amido-phenol with CS₂ (Hofmann, B. 20, 1789).—2. By heating C₆H₅.N₂.C₅H₅ with CS₂ at 265° (Jacobson a. Frankenbacher, B. 24, 1403).-3. By heating phenyl thiocarbimide with sulphur.

Properties.—Needles or plates, v. sol. alcohol and ether. Forms B'HgCl₂. Oxidised by K₂Cr₂O, to $(C_6H_4 < N_S)_2S_2$, which yields $B'K_2Cr_2O_4$

MeI yields $C_{\bullet}H_{\bullet} < N > C.SMe [52°]_{\bullet}$ [180°]. which forms B'2H2PtCl2, crystallising in plates.

SULPHYDRO - METHENYL - BENZENYL -THIO-AMIDOXIM $C_6H_6.C \stackrel{N.S}{\sim} C.SH.$ [162°]. Formed by the action of CS2 on an alkaline alcoholic solution of benzenyl-amidoxim (Crayen, B. 24, 388). Needles, nearly insol. water and cold alcohol. Conc. HClAq at 150° decomposes it into benzolo acid, CO₂, NH₂, H₂S, and S. NaOEt and Etl yield C₂H₂, Ce NS OEt [49].

while benzyl chloride forms, in like manner, $C_{\bullet}H_{\bullet}.C \stackrel{N.S}{\leqslant} C.CH_{\bullet}Ph$ [79°].

SULPHYDRO - METHENYL - p - TOLYL -METHENYL-THIO-AMIDOXIM C.H.N.S. i.e. $C_0H_1Me.C \stackrel{N.S}{\leqslant} C.SH.$ [166°]. Formed from CS₂ and tolyl-methenyl-amidoxim (Crayen, B. 24, 391). Crystals (from alcohol acidulated with HOAc), insol. water, sl. sol. ether. Forms salts with bases. p-Tolyl-methenyl-amidine forms the salt C₆H₄Me.C(NH₂):NHHA' [172°]. Boiling EtI and NaOEt yield $C_0H_4Me.C < N.S > C.SEt$ [87°].

SULPHYDRO - METHYL - GLYOXALINE $C_4H_6SN_2$ i.e. CH.NMe > C.SH. [142°]. (c. 280°). Formed by boiling CH(OEt)₂.CH₂.NH₂ with methyl thiocarbimide, alcohol, and dilute (30 p.c.) H₂SO₄ (Wohl a. Marckwald, B. 22, 1354). Plates, v. sol. water, sol. alcohol.—B'2PtCl,. B'AuCl₈.—C₄H₈AgSN₂.—B'MeI. [148°]. Needles.

derivative CH.NMe C.SMe. Methyl(225° un-Methylimidazolyl methyl sulphide. cor.). Formed by treatment with MeI and alkalis. Liquid, miscible with water, alcohol, ether. Boiling dilute HNO, and forms $C_sH_7(NO_2)SN_2$ [85°], which yields $B_2H_2PtCl_8$ [197°]. $C_sH_8SN_2MeI$. [173°]. Needles, v. sol. Aq.

Reference. — OXY-SULPHYDRO-DI-METHYL-GLY-OXALINE.

SULPHYDRO-DI-METHYL-PYRIDINE

 $CS < \frac{CH:CMe}{CH:CMe} > NH.$ [210°-215°]. Formed by heating oxy-di-methyl-pyridine with P2S, at about 160° (Guthzeit a. Epstein, B. 20, 2113). Needles, v. sol. hot water and alcohol.

SULPHYDRO - METHYL - QUINAZOLINE, $C_6H_4 < \stackrel{CH_2.NMe}{NH.CS}$ Dihydride [139°].

Formed by warming ω-oxy-tolyl-methyl-thiourea with HClAq (Söderbaum a. Widman, B. 22, 2935). Needles.— $B'_2H_2PtCl_6$. [195°].— $B'HAuCl_4$. [153⁶]. Minute needles.

SULPHYDRO-METHYL-QUINOLINE

 $C_{10}H_0NS$ i.e. C_0H_4 $\stackrel{CMe:CH}{\sim}_{N=C.SH}$. [253°]. Formed by heating (Py. 8,1)-oxy-methyl-quinoline with P_2S_5 (Roos, B. 21, 625). Small brown needles (from alcohol). Tastes bitter. Insol. cold water. Does not react with hydroxylamine and phenylhydrazine. H₂O₂ forms C₂₀H₁₈N₂S₂ [167⁵] crystallising from benzene in plates. EtI and alcohol at 100° form C_eH₄ CMe:CH_N which is oily and yields B'HI [240°] and B',H,PtCl, aq crystallising in reddish prisms. Sulphydro-methyl-quinoline

 C_0H , C_0SH): C_0H . [187°]. Got by heating (Py. 1,3)-oxy-methyl-quinoline with P_2S_a at 150° (Roos, B. 21, 629). Yellow prisms (from water). Etl and NaOEt yield C. H. EtNS [56°].—B'HCl. [205°-210°] (Conrad a. Limpach, B. 21, 1972). SULPHYDRO-METHYL-THIAZOLE.

hydride $CH_{a-N}^{CHMe.S}>C.SH.$ [82°]. Formed from β-bromo-propyl-amine hydrobromide, NaOHAq, and CS, (Hirsch, B. 23, 968). White needles, v. sol. hot water. Yields C.H., MeNS(SMe) (217°), C₃H₃MeNS(SEt) (229°), and the propyl ether C₂H₃MeNS(SC₃H₁) (247°).

 ψ -SÜLPHYDRO-(a) -NAPHTHYL - METHYL-BIAZOLE OCCH.NC10H. [86°]. Formed from acetyl (a)-naphthyl-hydrazine and CSCl₂ in CHCl₃ (Preund, B. 24, 4184). Yellow needles, v. e. sol. alcohol. The isomeride from (3). naphthyl-hydrazine melts at 109°.

SULPHYDRO-PENTHIAZOLE Dihydride CH₂ CH₂ S C.SH. [132°]. Formed by shaking γ -bromo-ethylamine hydrobromide with NaOHAq and CS₂ in the cold (Gabriel a. Lauer, B. 23, 92). Crystals, v. sol. hot water and alcohol, insol. acids, sol. NaOHAq. Bromine-water at 100° produces CH₂(NH₂).CH₂.CH₂.SO₂H.

 ψ - SULPHYDRO - PHENYL - AMIDO - (α) -NAPHTHYL-THIOBIAZOLE $C_{18}H_{13}N_{3}S_{2}$ S<C(NHPh):N. [255°]. Formed by mixing NHPh.CS.NH.NHC, H, with CSCl, in benzene (Preund, B. 24, 4192). Colourless crystals, v. sol. acetone, sl. sol. alcohol.

ψ-SULPHYDRO - PHENYL-METHYL-BI-AZOLE OCCS. NPh. [72°]. Formed from dichloro-phenyl-methyl-biazole dihvdride alcoholic ammonium sulphide (Freund a. Kuh, B. 23, 2837). Formed also from NHAc.NHPh and CSCl₂ (Freund a. Goldschmidt, B. 21, 2456). Prisms (from alcohol). Yields phenyl thiocarbimide on distilling with P2S

SULPHYDRO-PHENYL-GLÝOXALINE

 $C_9H_8N_2S$ i.e. $C_{N:C(SH)}>NPh$. [181°]. Formed by boiling NHPh.CS.NH.CH2.CH(OEt)2 with dilute H2SO, (Wohl a. Marckwald, B. 22, 569, Needles (from hot water), v. sol. NaOHAq, insol. NH3Aq.—Salts: C9H7AgN2S.— B'2H2PtCl_s: deep-red crystals.—B'MeI. [152°].

Methyl derivative N=C(SMe)>NPh. [54°]. Formed by treatment with alcoholic NaOH and MeI. White needles (from alcohol), insol. water.—B'C₂H₃N₃O₇.—B'HNO₃. Needles, v. e. sol. water.—B'MeI. [177°]. Crystalline rosettes (from water). Oxidised by dilute HNO. to a nitro- compound [116°

 ψ - SULPHÝDRO - PHENYL - DI - METHYL-PYRIDINE DICARBOXYLIC ETHER

 $CS < CMe:C(CO_2Et) > NPh.$ [245°]. Formed by heating CS CMe:C(CO₂Et) O with aniline CMe:C(CO₂Et) R 20 2112). and HOAc (Guthzeit a. Epstein, B. 20, 2112). Sulphur yellow crystals (from alcohol).

. SULPHYDRO - PHENYL - NAPHTHYL . BIAZOLE OCCS. NC .. H. [164°]. from NHBz NHC₁₀H, and CSCl₂ in CHCl₃ (Preund, B. 24, 4186). Needles, v. sol. alcohol. SULPHYDRO - PHENYL - QUINAZOLINE.

Dihydride C.H. CH, NPh. [197°]. Formed by warming ω-oxy-tolyl-phenyl-thio-urea with conc. HClAq (Söderbaum a. Widman, B. 22, 1671). Needles. — B',H,PtCl,. B'HAuCl,. [197°]. Needles. [219°]. -

Reference. -- OXY - SULPHYDRO'- PHENYL - QUIM-

AZOLINE.

DI-SULPHYDRO-DI-PHENYL-THIO-UREA. Di-methyl derivative CS(NH.C.H.,SMe),. [162°]. Formed by boiling [1:2] NH₂.C.H.,SMe with CS₂ and solid KOH (Hofmann, B. 20, 1794). Prisms, v. sl. sol. alcohol. Forms, when heated, SCN.C.H.,SMe (c. 270°).

B-SULPHYDRO-PROPIONIC ACID

CH₂(SH).CH₂.CO₂H. Formed from β-iodopropionic acid and KSH. Got also by reducing sulphido-dipropionic acid (Lovén, J. pr. [2] 29, 376). Liquid, miscible with water, alcohol, and ether. Easily oxidised by air. CuSO₄ gives a paleviolet pp., but if not in excess it forms a yellow pp. of CH₂(SCu).CH₂.CO₂H while S(C₂H₄.CO₂H)₂ remains dissolved.—Hg(S.C₂H₄.CO₂H)₂. Scales.—Bi(S.C₂H₄.CO₂H)₃.

Isomeride.—THIOLACTIC ACID.

SULPHYDRO - PROPYL - PHTHALIMIDE C₈H₄O₂:N.CH₂.CH₂.CH₂SH. [48°]. Formed by heating γ-bromo-propyl-phthalimide with KSHAq at 100° (Gabriel a. Lauer, B. 23, 88). Needles (from hot ligroïn). Fuming HClAq at 180° forms phthalic acid and γ-amido-propyl mercaptan.

(Py. 3)-SULPHYDRO-QUINOLINE

C_eH₄ CH:CH N:C.SH [174°]. Formed by heating (Py. 3)-oxy-quinoline with P₂S₅ at 140° (Roos, B. 21, 620). Yellow plates (from dilute alcohol), insol. cold water, sol. acids and alkalis. Oxidised by H₂O₅ to C₁₆H₁₂N₂S₂ [137°]. EtI and alcohol at 100° form C₉H₆(SEt)NHI [154°], which yields (C₆H₆(SEt)N)₂H₂PtCl₆ aq, melting, when anhydrous, at 190°.

SULPHYDRO - THIAZOLE. Dihydride CH_S CH_N > C.SH. [107°]. Formed from bromoethyl-amine, NaOH, CS₂, and alcohol (Gabriel, B. 22, 1152). Needles, sol. alcohol and hot water. Sol. alkalis. Bromine-water yields taurine. Alcohol, KOH and MeI form oily C₂H₄ S C.SMe (217°), insol. alkalis but sol. acids.

Reference .- OXY-SULPHYDRO-THIAZOLE.

SULPHYDRO-ACETONE. The compound CH₂.C(SEt)₂.CH₂.SEt (170°) is formed by passing gaseous HCl into CH₂.CO.CH₂.SEt mixed with mercaptan (Autenrieth, B. 24, 167).

ω-SULPHYDRO-o-TOLUIC ACID. Methyl derivative CH₂(SMe).C₆H₄.CO₂H. [138°]. Formed by the action of conc. HClAq at 100° on the nitrile CH₂(SMe).C₆H₄CN (278°), which is got by dissolving thio-phthalimidine in alcoholic KOH and adding MeI (Day a. Gabriel, B. 23, 2484). Light oil.

SULPHYDRO-ISOVALERIC ACID

CHPr(SH).CO.H. Formed from a-bromo-iso-valeric acid and alcoholic KSH (Duvillier, Bl. [2] 80, 507).

SUPER. Use of this prefix applied to inorganic compounds. The prefix super is sometimes employed to denote the presence in compounds of relatively more of the characteristic negative element, or elements, than is found in the normal compounds of the same form. The prefix per is, however, much oftener used than super. It has been proposed by Mendeléeff to designate the exides that belong to the same type as H₂O₈

superoxides, to distinguish them from other peroxides (v. Oxnes, vol. iii. p. 660).

M. M. P. M. SUPERBIN. A poisonous substance contained in the root of Gloriosa superba (Warden, Ph. [3] 11, 495).

SYCOCERYL ALCOHOL C₃₂H₃₄O. [114°]. Occurs as acetyl derivative in the resin of *Ficus rubiginosa* (De la Rue a. Müller, C. J. 5, 62; Rennie a. Goyder, C. J. 61, 916). Thin crystals (from alcohol), insol. water and alkalis. BzCl yields a crystalline benzoyl derivative.

Acetyl derivative C₂₄H₅₆O₂. [121°]. Flat six-sided tablets, v. sol. hot alcohol, ether, and

chloroform.

SYLVANE C₅H₈O. (63°). S.G. 887. Got by distilling pine-wood (Atterberg, B. 13, 879). Liquid, readily resinified. Oxidised by KMnO₄ to acetic acid. Not attacked by Na, alkalis, or Ac₂O. HCl passed into its ethereal solution forms a small quantity of C₁₅H₂₂O, a liquid boiling at 240°.

SYLVANE-ACETIC ACID v. METHYL-FUR-

FURYL-ACETIC ACID.

SYLVANE - CARBOXY-ACETIC ACID v.
METHYL-CARBOXY-FURFURYL-ACETIC ACID and METHBONIC ACID.

SYLVESTRENE v. TERPENES.

SYLVIC ACID C=78.7 p.c. $\mathbf{H}=9.7$ p.c. [c. 162°]. $[\alpha]_D=-53^\circ$. This acid is obtained from colophony by repeated crystallisation from dilute alcohol, and is perhaps identical with abietic acid (Liebermann, B. 17, 1884; Haller, B. 18, 2166). By heating with HI and P it is converted into a terpene $\mathbf{C}_{10}\mathbf{H}_{10}$. Sylvic acid appears also to be formed by the action of gaseous HCl on an ethereal solution of dextropimaric acid (Vesterberg, B. 19, 2173). A sylvic acid $\mathbf{C}_{20}\mathbf{H}_{30}\mathbf{O}_2$ is described by Maly (Sitz. W. 44, 121) as formed by adding dilute $\mathbf{H}_2\mathbf{SO}_4$ or gaseous HCl to a solution of abietic acid. Duvernoy (4. 148, 147) obtained a sylvic acid [129°] by distilling pimaric acid in vacuo.

Isosylvic anhydride $C_{10}H_{50}O_3$. (250° at 80 mm.). $a_D = +63^\circ$. Got by distilling rosin in vacuo (Bischoff, B. 23, 1921). Colourless, brittle, microcrystalline substance, insol. water, v. sol. alcohol and ether. Isosylvic acid [62°] is ppd. on adding dilute HOAc to its solution in

potash.

SYLVINOLIC ACID C₂₂H₂₆O₄? [130°]. Formed, together with sylvic acid, by saturating an alcoholic solution of abietic acid with HCl (Maly, Sitz. W. [2] 44, 121). Amorphous powder, v. sol. alcohol and ether.—CaA".—Ag₂A": pulverulent pp., insol. NH₂Aq.

SYMBOLS. Symbols are employed to express the composition, and, as far as possible, the interactions, of compounds; v. Equations, CHEMICAL (vol. ii. p. 433), FORMULE (vol. ii. p. 572), ISOMERISM (vol. iii. p. 79), and cf. OPTICAL METHODS (this vol. p. 253). M. M. P. M.

TICAL METHODS (this vol. p. 258). M. M. P. M.

SYNANTHEENE O. H. P. Phosens. Occurs
in crude anthracene (Zeidler, A. 191, 298; cf.
Barbier, A. Ch. [5] 7, 526). Yellowish-white
plates. Yields a di-bromo-derivative O. H. Br.
[175°], crystallising in minute yellow needles,
sol. alcohol (difference from di-bromo-anthracene). Di-bromo-synanthrene is oxidised by
CrO. and HOAc to a quinone (?) [240°-250°].

SYNANTHRIN . INVLIN.

SYNTHESIS. The building up of compounds from their elements, or from groups of elements. SYNTONIN v. PROTEÏDS.

SYRINGIN C₁,H₂,O₂. [191°]. Dimethoxyconiferin. Occurs in the bark of the lilac (Syringa vulgaris) and of the privet (Ligustrum vulgare) (Bernays, J. pr. 25, 121; Kromayer, Ar. Ph. [2] 108, 7; 109, 18, 216; 113, 19; Körner, G. 18, 215). White needles (containing aq), v. sl. sol. cold water, v. sol. alcohol, insol. ether. Its aqueous solution is slightly bitter, is levorotatory, and does not ppt. metallic salts. Conc. HNO₂ gives a blood-red solution. Conc. HClAq forms a colourless solution, turning blue, and finally giving a blue pp. Conc. H₂SO₄ gives a yellowish-green colour, changing to violetbrown and, on pouring into water, giving a bright-blue solution, which soon deposits an azure powder. Does not reduce Fehling's solution. Decomposed by emulsin into glucose and syringenin. KMnO₄ yields glucosyringic acid. CrO₂ forms glucosyringic aldehyde.

Syringenin $C_{11}H_{14}O_4$ i.e. [5:3:4:1]? $C_0H_2(OMe)_2(OH).C_0H_4.OH$. Formed as above.

Besembles coniferyl alcohol.

Glucosyringis acid $C_{15}H_{20}O_{10}$. [208°]. Formed as above. Needles or prisms (containing 2aq), sl. sol. cold water. Decomposed by dilute H_2SO_4 into syringic acid and glucose.

Syringic acid C₄H₄(OH)(OMe)₂.CO₂H. Dimethyl derivative of gallic acid. [202°]. Decomposes at 230° into CO₂ and the di-methyl derivative of pyrogallol.—BaA'₂8aq: four-sided tables.—MeA'aq. •[83·5°]. Crystals, sol. Aq.

Methyl derivative C₆H₂(OMe)₃.CO₅H. [168°]. Needles. Yields C₆H₃(OMe)₃ when distilled with lime. Identical with the tri-methyl derivative of gallic acid.—MeA'. [82·5°]. Needles.

Glycosyringic aldehyde $C_{15}H_{20}O_{\theta}$. [162°]. Colourless, silky needles. Decomposed by emulsin or dilute $H_{c}SO_{\phi}$ into glucose and syringic aldehyde. Yields a phenyl-hydrazide [156°] and a crystalline oxim.

and a crystanne cam.

Syringic aldehyde $C_pH_{10}O_4$ i.s. $C_eH_2(OH)(OMe)_x.CHO$. [111·5°]. Small needles, smelling like vanillin. Turns brown in air. Reacts with phenyl-hydrazine, and combines with NaHSO_{2*}.

 \mathbf{T}

TAIGUIC ACID v. LAPACHIC ACID.

TALOMUCIC ACID $C_sH_{10}O_s$. [158°]. [α]_D = +29·4° at 20°. Formed from talonic acid and HNO, (S.G. 1·15) (Fischer, B. 24, 3625). Minute plates, v. e. sol. cold water and warm alcohol; v. sl. sol. ether. Converted by HOl and HBr at 150° into dehydromucic acid. Yields a phenyl-hydrazide [185°–190°]. Pyridine at 150° forms mucic acid.—CaA": crystalline powder.

TALONIC ACID C.H., 20,. Formed by heating galactonic acid with pyridine at 150° (Fischer, B. 24, 3622). Purified by means of its brucine salt. Syrup, very soluble hot alcohol. Lævorotatory. Yields a phenyl-hydrazide C., H., O. (N., H., Ph.) [c. 155°].—CdA', 2aq. Needles (from alcohol), v. e. sol. water.—Brucine salt. [130°-133°]. Crystals, v. sol. water.

TAMPICIN C₂₄H₂₄O₁₄. [130°]. A neutral substance resembling convolvulin, occurring in Tampica Jalap (Spirgatis, N. R. P. 19, 452). Resin, v. sol. alcohol and ether. Converted by hot baryta-water into amorphous tampicie acid C₂₄H₂₆O₁₇. Boiling dilute acids resolve tampicin into glucose (8 mols.) and tampicolic acid C₁₆H₂₂O₃, which crystallises from dilute alcohol in minute needles and forms NaA' and EtA', both being crystalline.

TANACETIN C₁₁H₁₆O₄. A bitter substance obtained from the leaves and flowers of the tansy (Tanacetum vulgare) (Leroy, J. Chim. Med. 21, 357; Leppig, J. 1882, 1175). Amorphous, sol. water and alcohol.

TANACETOGENIC ACID C,H14O2. (114° at 15 mm.). Formed by the action of Br and (4 p.c.) NaOHAq on tanacetone (Semmler, B. 25, 8346). Oil, solidified below 0°.—AgA'.

TANACETONE C₁₀H₁₀O. (84.5° at 13 mm.). Occurs in oil of absinthe, thuja, and sage, con-

stituting the chief part of absinthol, salviol, and (8)-thujol (Semmler, B. 25, 3848, 3850). Got from ethereal oil of Tanacetum vulgare by shaking with NaHSO₂, decomposing the crystalline product with Na₂CO₃, and distilling with steam. Oil, v. sol. alcohol and ether. Does not reduce ammoniacal AgNO₃. Dextrorotatory; $\alpha = 38\frac{1}{3}^{\circ}$ in a 20 c.m. tube. S.G. $\frac{29}{3}$ -213. $\mu_{\rm D} = 1.450$.

Reactions.—1. Br and KOHAq yield bromoform, hence it probably contains CO.CH_s.—2. Reduced in alcoholic solution by Na to tanacetyl alcohol $C_{10}H_{10}O$, (93° at 18 mm.), S.G. 10 925, $\mu_D=1.4635$. This body does not combine with Br. With PCl_s it yields tanacetyl chloride (72° at 10 mm.).—3. Oxidised by KMnO. Needles [78°] (from ligroin) or plates [74·5°] (from water). Yields AgA'. Hydroxylamine yields $C_{10}H_{10}O_2(\text{NOH})$ [103] when prepared from the needles and [169°] when got from the plates. The ketonic acid is oxidised by Br and NaOH to tanacetogen dicarboxylic acid $C_9H_{10}O_4$ [142°]. Plates (from water). This acid yields Ag_2A'' while Ac_2O yields the anhydride $C_9H_{12}O_2$ [55°], which gives isopropyl succinic acid [114°] when fused with potash.

Oxim $C_{10}H_{1c}$:NOH. [52°]. (186° at 20 mm.). Reduced in alcoholic solution by Na to tanacetylamine $C_{10}H_{1}$ NH₂ (80°5 at 14 mm.), S.G. 12 ·874, μ_D 1·462, which yields B'HCl, which when heated gives tanacetene $C_{10}H_{1c}$ (68° at 14 mm.), S.G. 20 ·841, μ_D 1·476. Tanacetone oxim may be converted by warming with alcohol and dilute sulphuric acid into the cymidine $C_2H_1(C_2H_1)(NH_2)$ Me [1:8:4], whence nitrous acid forms carvacrol.

TANACETOPHORONE $C_aH_{12}O$. (90° at 18 mm.). S.G. 12 •938. μ_D 1·482. Got by distilling tanacetogen dicarboxylic acid with soda-lime

(Semmler, B. 25, 3350). An oil, smelling like camphor-phorone. Reacts with hydroxyl-amine. [182°]. TANGHININ C27H40O8 2aq. $[a]_{p} = -67^{\circ}$. Obtained from the kernels of

Tanghinia venenifera of Madagascar (Arnaud, C. R. 108, 1255; 109, 701). Crystals (from alcohol), v. sl. sol. water, sl. sol. ether. Cardiao Barytapoison. Resinified by dilute acids.

water at 180° forms C_nH₄₄O₁₀,
TANNIN. TANNIC ACIDS. Under the above names are included a large series of compounds, occurring in the vegetable kingdom, of different constitutions, many of which have been very imperfectly studied. Their chief characteristics are their astringent taste and property of giving a dark-blue or green colouration with iron salts and a precipitate with a solution of gela-They all reduce solutions of the noble metals, and absorb oxygen in presence of alkalis. They are all aromatic derivatives, and yield on fusion with potash either pyrogallol or pyrocatechin. In some cases they have been synthetically prepared from these phenols and their derivatives, but in many cases the tannin appears to exist in the plant as a glucoside and attempts at a synthetical production have proved futile. Schiff (B. 15, 2590) prepared some of the tannins synthetically, and concluded that they were anhydro-compounds of gallic and protocatechuic acids respectively. Etti (A. 186, 332) and others have also, for example, synthetically prepared catechutannic acid by heating pyrocatechin with water or alkalis. Gallotannic acid was formerly believed to be a glucoside, but subsequent investigation showed that the amount of sugar present was variable, and Schiff (A. 170, 43) finally proved that, although existing in the plant as a very unstable glucoside, when it was isolated it had the constitution of an anhydre- acid formed by the removal of one molecule of water from two molecules of a trioxy-benzoic acid. He considered that in gallotannic acid the anhydride was formed thus:-C₆H₂(OH)₃CO.O.C₆H₂(OH)₂COOH, whilst C. Etti, who has similarly investigated the tannins of the formulæ $C_1, H_{10}O_9$ and $C_{20}H_{20}O_9$ has proved that they also are not glucosides, and regards them as derivatives of a ketonic acid of the C_aH₂(OH)_a.CO.C_aH(OH)_aCOOH. would appear, then, possible to define the tannic acids as mono-carboxyl acids formed by condensation from two molecules of poly-oxy-benzoic acids and which exist as unstable glucosides in plants. The ease with which gallotannic acid is hydrolysed renders it probable that the difference in constitution between Schiff's and Etti's formulæ does actually exist, and renders it possible to subdivide the tannius into these two groups so soon as the different tanning have been re-examined from this point of view. The ketonic character of the tannins was established by Etti through noting that they reacted with phenyl-hydrazine and hydroxylamine.

The origin of tannin in plants has given rise to much debate. According to Waage (Ph. 1890; Phar. Centr. 1891, 247) its formation is similar to that of the conversion of glucose into starch. The plant removes one molecule H_.O from glucose for reserve purposes, and if under favourable circumstances three molecules H₂O can be removed, a body having the composition of a triketohexamethylene would be produced

CH_COCH_COCH_CO. This constitution corresponds to the secondary form of phloroglucin. It is probable that the phloroglucin combines with the CO₂ in the nascent state produced by the respiration of the plant, and is thus con-verted into a carboxylic acid (gallic acid); two molecules of this acid by the removal of another molecule of water would then form tannin. Light and chlorophyll are necessary for the production

Westermaier (B. B. 1887, 127-143) showed that with experiments made with Quercus pedunculata the tannin migrates downwards through the bark and the pith. Starch is always present, and it is probable that the starch only migrates in the form of tannin. The leaves of Rumex patentia and Rheum rhaponticum give both the starch and the tannin reactions. Schultze (A. a. 14, 525-526; B. C. 18, 137), by microchemical observations, also supports the view of Sachs and Heberland that the leaves of evergreens contain tannin as a winter reserve material. Tannin and starch only rarely occur simultaneously in the same cells. When the cells are rich in starch they contain a smaller amount of tannin, and vice versa.

In the case of leaves containing fatty oils and tannin, the cells which contain the oil are free from tannin, and cells containing no oil are free from starch. Kraus (B. C. 330-334) also notes that the formation of tannin in leaves depends on the presence of light and CO2, but believes that it acts more as a protecting agent either to prevent the plant from being eaten or rotting than as a reserve material. He points out that as fallen leaves contain as much tannin as they did during their best time of growth, the leaf tannin is of no value to the plant. Heckel a. Schlagdenhauffen (Ph. July 1892) note that the protecting leaf buds of certain species of gardenia contain a resinous substance which on analysis gives figures resembling cinchotannic acid. Both the gardenias and the cinchonas belong to the same order of plants (cf. Cross a. Bevan, C. J. 41, 106; Gardiner, Ph. [3] 14, 588; Braemer, Les Tannoides, Toulouse, 1890; Büsgen, C. C. 1890, 397).

Classification. -- No satisfactory classification of these bodies exists. Wagner (Fr. 5, 1) divides them into pathological and physiological tannins. The former are chiefly glucosides, and precipitate gelatin, yield gallic acid when hydrolysed with dense sulphuric acid, and when heated alone form pyrogallol. The physiological tannins, on the other hand, form leather, and on distillation yield pyrocatechin. The old classification into iron-blueing and iron-greening tannins is found now to be untenable, as the presence of acids and salts modifies the colour which the tannins produce with FeCl, and the purity of many of the tannins examined by the earlier investi-

gators cannot be guaranteed.

Reactions .- The oxidation of gallic and tannin acids by nitric acid of moderate dilution yields oxalic acid and two acids, probably tri-oxy-glutaric and trioxy-butyric. The acetyl oquercitannic acid yields the same acids on oxida-tion (Böttinger, A. 257, 248-252). Metallic Na does not reduce gallic acid in absolute alcohol, but tannic acid similarly treated undergoes simple hydrolysis, the yield of gallic acid amounting to 60 p.c. of the weight of tannin. No reduction products could be prepared (Böttinger, A. 258, 252-260). Ammonia and zinct dust at 60° convert both acids into benzoic acid. Dilute H₂SO₄ and Zn also form benzoic from gallic acid (Guignet, C. R. 113, 200-201).

The action of heat. Most tannins begin to blacken at 120°, and at 160° decomposition into metagallic acid, and either pyrogallol or pyrocatechin, takes place. Those which yield pyrogallol usually also form a 'bloom' on leather, and give a blue colour with FeCl₂. This class includes galls, sumac, chestnut, and oak bark. The pyrocatechin-yielding tannins do not form a 'bloom' on leather, give green compounds with FeCl₂, and include hemlock, catechu, rhatany, and mangrove (Trimble, The Tannins, 1892).

The products of the action of heat on the tannins are best obtained by allowing the heating to take place on the material suspended in glycerin (1 g. in 5 c.c. glycerin). After heating to 200° for 30 minutes, dilute with water, and extract with ether. The ethereal extract contains either pyrogallol or pyrocatechin. The tannin must be freed from gallic acid and catechin before subjecting it to this test by previous ether extraction of the aqueous solution.

The action of dilute acids. 2 p.c. absolute HCl at 100° in sealed tubes decomposes the tannins; insoluble ellagic acid, anhydrides, or phlobaphenes separate, and the filtrate contains gallic acid, glucose, and unaltered tannin. Glucose should be removed from the tannin by repeated lead acetate precipitation before this treatment, or the amount of glucose before and after hydrolysis may be determined by Fehling's solution (Wehmer a. Tollens, A. 243, 327).

The action of alkalis. The pyrocatechinyielding tannins on heating with alkali yield protocatechuic acid, and either phloroglucin or acetic acid. The pyrogallol-yielding tannins form gallic and ellagic acids (20 g. with 150 c.o. of KHO solution, sp.gr. 1.2 for 3 hours are convenient quantities (Trimble, The Tannins).

The tannins in the bark of oak, chestnut, horsechestnut, fir, &c., are precipitated in brominated condition by bromine water; tannin, sumach tannin, and oak wood tannin are not so precipitated (Böttinger, A. 240, 330). oak tannin, Böttinger, by treating with acetic anhydride, has formed an acetyl-o-compound of the formula C15H7Ac5O8, and from it two bromoderivatives $C_{18}H_{10}BrAcO_0$ and $C_{18}H_{18}Br_4AcO_0$ (B. 20, 761-766). A benzoyl-o-derivative has also been obtained from tannin by treatment with cone. NaOHAq and benzoyl chloride (Böttinger, A. 254, 370). The same author has attempted to prepare cyanhydrins, but without success. Well-dried tannic acid heated with anhydrous HCN in sealed tubes yielded only an amide of gallic acid. Treated with hydroxylamine no oxim was formed, but only gallic acid (A. 259, 182-136). With pure tannin phenyl-hydrazine, however, forms derivatives which do not crystallise, CO, and N being evolved at the same time. The tannins experimented upon were extracts of sumach, divi divi, oak wood, oak bark and pine bark; the dry precipitates contained phenyl-hydrazine derivatives of tannic and gallic acids, osazones of dextrose and lavulose, and caramel (A. 259, 125-132; A. 256, 342).

Tannin solutions are very unstable, undergoing hydrolysis into gallic acid on standing. The hydrolysis into gallic acid is almost quantitative when the acid is dissolved in conc. H.SO₄ and then water added (Böttinger, A. 254, 378). When heated with KHSO₄ and glycerin to 119°–200° and the melt extracted with water, a residue is left soluble in absolute alcohol which contains two new acids of the composition C₁₄H₁₄O₇Aq and called hydrotannic and isotannic acids respectively. The former, which is the less soluble in alcohol, forms with Ac₂O a tetra-acetyl-compound while the iso-acid forms a brown triacetyl-compound (C. J. 1892, 181).

Preparation of tannic acid for pharmaceutical purposes. Pelouze in 1834 first suggested the extraction of tannic acid from galls by percolation with ether, and this method is still in use. The powdered galls are placed in a closed percolator with commercial ether containing alcohol and water. The ethereal layer contains gallic and ellagic acids, together with resins and fats and a small portion of the tannic acid; the lower aqueous solution contains nearly pure tannic acid. The percolation is stopped when the lower layer ceases to increase in volume. The B. P. method consists in taking the damp galls, macerating with commercial ether, and expressing through muslin (Leconnet a. Dominé). The purest acid is obtained when 10 pts. powdered galls are allowed to stand two days covered in a percolator with a mixture of 12 pts. ether and 3 pts. alcohol. At the end of this time the percolation is commenced until 10 pts. are obtained. One-third its volume of water is then added, and the mixture well agitated. The aqueous layer contains nearly pure tannic acid, and is evaporated rapidly in vacuo. The commercial acid is known under the names of ether-tannin, alcohol-tannin, or water-tannin, according to the method of extraction adopted on a large scale.

Purification.—Petroleum ether percolation removes fat, wax, and some of the colouring matter. Solution in water removes resin, and, after decantation, sodium chloride precipitates the acid. Trimble adds lead acetate to the aqueous solution to precipitate colouring matter, filters, and extracts with acetic ether. The acid is re-dissolved in water, and extracted with ether to remove the last traces of gallic acid, and the aqueous solution finally dried in vacuo.

Detection.—1. Dilute H₂SO₄ or HCl precipitates conc. solutions.—2. FeCl₃, a blue or green precipitate destroyed by long boiling, sol. weak mineral acids.—3. Lime-water precipitates the Ca salt. 4. Cinchonine sulphate precipitates white cinchonine tannate.—5. Gelatin forms leather and albumen also gives a precipitate. Alum and NH₄Cl render the precipitation more complete. 6. An alcoholic solution of thymol, followed by conc. H₂SO₄, yields a rose-coloured turbid solution (pyrogallol violet; gallic acid gives no colour: Saul, Ph. [3] 17, 387).—7. Iodine in presence of neutral salts gives a purple colouration (Nasser). 8. Acetate of lead acidulated with acetic acid ppts. tannic and not gallic acid (Guyard, Bl. 2, 41, 386).—9. NH₄Cl and NH₆ give a white precipitate rapidly becoming red, gallic acid a red coloura-

tion only.-10. Cl water and NH, give a red colouration both with gallic and tannic acids .-11. K.FeCy, gives a similar reaction (Rawson, C. N. 59, 52-53).—12. In plants, Moll (A. a. 12, 496) detects tannins micro-chemically by treatment with 7 p.c. copper acetate for some days, then cuts sections, treats with a drop of 0.05 p.c. ferricacetate, washes with water and then alcohol to remove chlorophyll, and mounts in glycerin and looks for blue or green stain. 13. KCN gives a green colouration with tannin and none with gallic acid.—14. K₂Cr₂O₇ gives a brown pp. with most tannins.—15. Saturated solutions of NaCl, CaCl2, KOAc, and a number of other salts ppt. tannins from solution. -16. Conc. H.SO, dissolves the dry tannins with a yellow colour, and on heating forms dark-red rufigallic acid and then black metagallic acid.—17. HNO. forms a yellow colour, and finally oxidises them to oxalic acid.—18. As O heated with the dry tannins converts most into ellagic acid.— 19. Most tannins are ppd. by the alkaloids.

Estimation of tannic acid. The methods devised for the estimation of tannic acid in vegetable extracts are very numerous, and may be grouped under the following heads:-

1. Gelatin or hide-powder absorption.

2. Titration with permanganate.

3. Precipitation with metallic salts.

4. Methods not included in the above.

For details of these various processes, vide Proctor, Textbook of Tanning; Trimble, The Tannins; Thorpe's Dictionary of Applied CHEMISTRY, and Rideal, Chemical Notes on Modern Tanning; The Assay of Tanning Materials; Industries, vol. xi. pp. 19, 139, 451. The two methods most generally employed are the hidepowder method and Löwenthal's permanganate

The hide-powder method is conducted as follows. A weighed quantity of the material is extracted with water and made up to a known 100 c.c. of this solution is then evaporated, dried, and weighed to give the total solids present in the extract. Another portion of the same solution is agitated with, aspirated, or filtered through hide powder, and the total solids left in 100 c.c. of the solution again deter-The difference gives the quantity of mined. tannic acid or matter which combines with gelatin per 100 c.c. of the solution (Fr. 24, 271).

The modified Löwenthal's process, which is now official in Germany, requires a standard solution of permanganate whose indigo value is known. A measured volume of the tannin solution, together with a known volume of indigo solution, is then titrated with the permanganate solution. Another equal volume of the tannin solution is then agitated with hide powder or shaken with a gelatin and alum solution and filtered, and the filtrate, after indigo solution of known amount has been added, is titrated with the permanganate solution. The difference in the quantity of permanganate consumed is a measure of the tannic acid present. The various tannins of commerce have different permanganate values; it is, therefore, necessary to standardise the permanganate for different tannin materials (Löwenthal, Fr. 16, 83; Kath-reiner, Fr. 18, 118; Simaud, Fr. 22, 595; Schroeder, Fr. 25, 121).

Gallotannic acid C, H, O. Occurs in gall nuts and sumach.

Constitution. C₆H₂(OH)₃.CO.O.C₆H₂(OH)₂.COOH (Schiff, A. 170, 43; Paul a. Kingzett, C. J. 33, 217).

Solubility. - Sol. acetone, glycerin, and oils;

insol. CS₂, CHCl₂, petroleum, and C₆H₆.

Compounds.—With gelatin contains 16.5 p.c. nitrogen = 84 p.c. tannin (Böttinger, A. 244, 227). (For other nitrogen values of gelatin-tannin compounds in leather, v. Rideal a. Trotter, S. C. I. 1891.) Penta - acetyl - compound C₆H₂(OAcO),CO.Ó.C₆H₂(OAcO)₂COOH

(Böttinger, B. 17, 1504).

Action of heat.—At 215° it forms pyrogallol, CO₂, and a trace of metagallic acid C₆H₁O₂. Strong heating at 240°-250° forms chiefly metagallic acid (Pelouze, A. 10, 159). Cold HNO₃ forms oxalic, trioxyglutaric, and trioxybutyric acids (Böttinger, A. 257, 248). Boiling with KHO forms tannomelanic acid C,H,O,; cold KHO and air form tannoxylic acid C, H,O, (Büchner, A. 53, 873). Hydrolysed by dil. H.SO, to gallic acid. Ferments also form gallic acid (van Tieghem, C. R. 65, 1092)

Oak bark. Tannic acid C17H10O9

Preparation .- The bark is extracted with alcohol, and the extract agitated with acetic ether and ether. Evaporate off ether, separate the precipitate of ellagic acid, and from residue separate gallic acid from the tannic acid by acetic ether and ether.

Properties .- Reddish-white powder. Sol. alcohol and acetic ether. Sl. sol. ether and water. Heated to 130°-140°C. it forms an anhydride C₈₄H₃₀O₁₇ which is sl. sol. water; sol. alcohol and alkalis, and with conc. H2SO, gives a second anhydride C₈₄H₂₈O₁₆. Heated with conc. HCl it evolves CH₃Cl, gives also the iodoform reaction. On dry distillation it forms pyrocatechin. Fused with KOH yields protocatechuic acid, pyrocatechol, and traces of phloroglucin (Etti, M. 1880, 262-278). When boiled with ether the above anhydrides form a third anhydride C.H.O. (M. 1883, 512). Other anhydrides $C_{24}H_{28}O_{18}$ (M. 1883, 512). Other anhydrides analysed by Etti are $C_{40}H_{28}O_{17}$, $C_{10}H_{36}O_{16}$, C₄₀H₃₄O₁₃, and C₄₀H₃₂O₁₄. Forms two hydrates (Löwe, C. J. 40, 901).

Salts. With Ca, Ba, and Pb; analysed by

Etti and Löwe (l.c.).

Derivatives.—Oak-red C14H10O62aq (Grabow-Derivouries.—OBE-Ted $C_{14}H_{10}O_{2}$ â 4 (Grabow-ski, A. 145, 2); $C_{18}H_{20}O_{1}$, (Böttinger, A. 240, 341). Böttinger has prepared various bromo-, acetyl-, and benzoyl-derivatives (*l.c.*) (Rochleder, A. 63, 205; Böttinger, A. 202, 270; B. 14, 1598; Löwe, Fr. 20, 210; Etti, M. 4, 514; Grabowski, A 145, 2) Grabowski, A. 145, 2).

Fraxitannic acid C26H32O14. In ash-leaves. Forms an anhydride $C_{26}H_{30}O_{15}$, a benzoyl- compound $C_{26}H_{26}O_{16}(OB_8)_{4}$, an acetyl- compound $C_{26}H_{26}(OAc)_4O_{16}$, a bromacetyl- compound $C_{26}H_{26}(OAc)_4O_{16}$, and a similar nitrogram of the property of the second of the property of the compound. On heating with Ba(OH), it forms protocatechuic acid and other products. On heating to 100° it loses 1 mol. H.O, forming an anhydride C. H. O. (Gintl a. Reinitzer, M. 3. 745 et seq.).

The following table gives the source of the principal tannins which have been investigated. and references to the papers in which a descrip-

tion will be found.

Plant	Part	Иапе	Formula	Per cent.	References
Esculus Hippocastanum	Nearly all parts	Horsechestnut tannin	$C_{2a}H_{2d}O_{12}$	1	Rochleder, Sitz. B. 53 [2] 478;
Æsculus Hippocastanum Algarobilla	Juice of berries Pods	Rhamnotannic acid Ellagotannic acid	C ₂₅ H ₂ (O ₁₅ ,H ₂ O C ₁₄ H ₁₆ O ₁₆ ,i.e. C ₅ H ₂ (OH) ₂ .CO.O.C ₆ H ₂ (OH) ₂ .COOH?	68.38	Bochleder, Z. 1867, 84 Godeffroy, C. N. 40, 144; Löwe, Fr. 14, 35; Zölffel, Ar. Ph. 229, 123; Hurst, D. P. J. 562, 288; Eitner
Alnus glutinosa	Wood	Alder tannin	$G_{\mathbf{p}}\mathbf{H}_{\mathbf{p}}0_{11}$	ı	D. P. J. 244, 80 Dreykorn a. Reichard, D. P. J.
Anacahuita Asperula odorata Aspidium Filix-mas	Wood and bark Root	Aspertannic acid Filitannic acid	C,H,o,	111	446
Atherosperma Moschatum Bablak (acacia) Betula rubra	Bark Fruit Bark	Atherosperma tannin — Beech tannin	C ₁₀ H ₁₀ O ₃ C ₃₀ H ₂₂ O ₅	19.0	Zeyer, J. 1861, 769 Wagner, Fr. 6, 10 Hofstetter, A. 51, 78; Etti, M.
Caffea arabica	Berry	Caffetannic acid	Ch.H.sO.	1	4. 66, Hasi-
Calandra granaria (corn]	Fracticornitannin	1	i	wetz, Sitz. B. 9, 286 Villon, C. N. 56, 175
Calluna vulgaria	Plant without	Callutannic acid	G,4H,10,	1	tz. B.
Canaigre	Bark	Canaigre tannin	C 58-10 per cent. H 5-33 ".	i	D. P. J. July 1893
Castanea vesoa	Bark	Chestnut bark tannin or	(same group as mangrove tannin) $C_{1}H_{1}O_{2}^{1}$	1	Trimble, C. N. Jan. 6, 1898
Catechu Cephalis Ipecacuanha	Boot 1	Catechutannic acid Ipecacuanhic acid	C,HH,O,? C,HH,O,? C,HH,O,?	1 1	Guignet, C. R. 113, 200 Willigt, A. 76, 342
Chirococca racemosa	Root	Caffetannic scid	C1122010 + \$1120 C13H18O	11	Rochleder, A. 84, 354; Sitz. B.
Cinchona	Bark	Quinotannic or cinchons- tannic acid	C, H, O,	ı	Schwartz, Sitz. B. 7, 250; Rembold, 4.143, 270; Ber-
	15				Pelletier a. Caventou, A. Ch. 55, 337

Plant	Part	Name	Formula	Per cent.	References
Cinchons nova	Bark	Quinovatannic scid	C, H ₁₈ O,	ı	Hasiwetz, 4. 79, 129; Bem-
Divi-divi	ı	Ellagotannic acid	C,H,OH),CO.O.C,H,OH),COOH?	ı	Zölffel, Ar. Ph. 229, 213; Löwe, Fr. 14, 35; Hurst, D. P. J.
Elm Erythroxylon Coes	Leaves in autumn Leaves	Xanthotannic acid Cocatannic acid	G ₂₈ H ₃₆ O ₄₁ ,3H ₂ O? G ₁₇ H ₂₂ O ₁₆ + 2H ₂ O?	1 1	Ferrian, J. 1858, 463 Niemann, J. 1860, 368; Warden,
Euphrasia officinalis	Green parts	1	$G_{x_2}H_{x_2}O_{x_2}$	i	Enz, Vierteljahrsch. Ph. 8,
Filix-mas Fraxinus excelsior	Leaves	Filitannic acid Fraxitannic acid	G.H.O.	11	Gintl and Beinitzer. M. 8. 745
Galium verum Hemlock	Bark	Galitannic acid	C1,41,20,41,0	1 1	Schwartz, A. 83, 57 Böttinger R 17, 1041
Норв	1	Hop tannin	0°4420	1	Bissell, C. J. 34 328; Ph. [3]
Nex paraguayensis	I	Caffetannic scid?	$G_{13}H_{18}O_8$?	ı	Rochleder, 4. 66, 39; J. 1856,
Krameria triandra	Bark of root	Rhatany tannin	G ₃₀ H ₂₀ O₂	ı	Wittstein, J. 1854, 656; Basbe,
Larch Laurna canatica	Bark	Larch tannin	1 D	1 1	J. 1880, 1060 Stenhouse, P. M. 23, 336 C. J. 34, 986 · C. J. 40, 602 ·
		ı	-11-11		Arata, An. Soc. Cient. Argent.
Mangrove	Bark	ı	C_16 H24 O 18	1	10, 193 Trimble, Ph. Feb. 4, 1893; S. C. I. 1893, 364; Sten-
Myrobalans	ı	Ellagotannic acid	C,(H,0),	I	house, Pr. 11, 405 Löwe, Fr. 14, 44; Zölffel, Ar.
Nux vomica Pinus sylvestris	Bark -	Igasuric acid Cortepinitannic or pinicor-	C ₂₇ H ₃₁ O ₁ ,	1 1	Fn. 229, 123 Ludwig, Ar. Ph. [3] ii. 137 Kwalier, Sitz. B. 11, 363
Pinus sylvestris	Needles	tannic acid Tannopinic acid	$(=2C_{10}H_{11}O_{2}+3H_{2}O)$ $C_{20}H_{10}O_{18}$?	1	Kwalier, Sits. B. 11, 856
Potentille Pormentilla	Root	Oxypinitannic acid			Dombold 4 145 F
Punica Granatum	Root bark	Pomegranate tannin	Carta Con Ha O13	1 1	Rembold, A. 143, 285
Unebracho Colorado	Gum or wood	Vuebrachitannic acid	C 52.52 per cent. H 5·11 " O 42·37 "	I	Jean, Arata, An. Soc. Cient. Argent. Feb. 1879; Bl. 28, 16

Guerous	Bark, wood, and Quercitannic acid	Quercitannic acid	C ₁₈ H ₁₂ O ₅ + 2H ₂ O (from wood) C ₁₈ H ₁₆ O ₁₆ C ₁₄ H ₁₆ O ₁₆	1	Etti, M. 10, 650; M. 1, 264; Stenhouse, P. M. 22, 425; Böttinger, A. 202, 270:
	Bark and gall- nuts	gall- Gallotannic acid	С,Н,О, <i>i.е.</i> С,Н,(ОН),СООС,Н ₂ (ОН),СООН	ı	Lowe, Fr. 20, 210 Berzelius, Lehrb. 3 Aufl. 6, 213; Löwe, Fr. 11, 378;
Quereus Ægilops	ı	ı	ı	19-26.7	Wagner, Fr. 5, 10; Stenhouse,
Quercus pubescens Quercus robur Red wine Rhododendron ferrugineum Rhorberh	Leaves Prot	Quercitannio acid Quercitannio acid Enotannin Rbdotannio acid Photografio acid	CmH.O. C1.H.O. 4C1.H.O. + 3H.O	1111	F. M. 22, 424 Etti, B. 17, 1823; M. 1, 262 Etti, B. 17, 1829; M. 1, 262 Gautier, Bl. 27, 496 Schwartz, Sitz, B. 9, 298
	Leaves and twigs	Gallotannic acid	CAH, O		A. 6,
Rhus Coriaria Rhus Cotinus Rubia tinctoria St Impetius beans	Leaves Leaves and stalks Leaves		26,4H ₂ O ₁ +H ₂ O	3.4 13.26	213; Schiff, 4. 170, 43 Lidoff, J. R. 20, 607 Lidoff, J. R. 20, 607 Willigt, 4. 82, 340
Salix triandra and S. undulata	Leaf, stems, and green twigs	- Rasulto solu	i 1	I I	Johanson, Ar. Ph. [3] 13, 103; E. Stenhouse, Pr. 11, 403; P. Stenhouse, Pr. 11, 403
Sorbus aucuparia	Juice of ripe	Sorbitannic acid	i	I	Vincent a. Delachanal, Bl. Z
Spruce Strawberry	Bark Root	Spruce bark tannin Fragarianin	G11H2010 P	11	
•	ı	Queroitannio acid	C,,H,O,	10	Stenhouse, P. M. 23, 332; O. Rochleder, A. 63, 205; Etti, C. B. 17, 1823; M. 1, 262; W. Hasiwetz & Malin, J. Ph.
Thuja occidentalia Unacaria Gambier Walnut	Green parts Leaves Episperm	Pinitannio acid Mioitannio acid	G,H,O, Besembles spruce bark tannin	111	101, 109 Rochleder, Sits. B. 29, 20 Böttinger, B. 17, 1129 Phipson, C. N. 20, 116

S. B.

TANSY OIL. The essential oil, obtained by distillation of the tansy (Tanacetum vulgare contains 1 p.c. of a terpene C10H16 (1550-1600), 26 p.c. of an alcohol $C_{10}H_{18}O$, and 70 p.c. of tanacetyl hydride $C_{10}H_{18}O$ (Bruylants, J. Ph. [4] 26, 893; B. 11, 449; cf. Persoz, C. R. 8, 483). Tanacetyl hydride $C_{10}H_{16}O$ (195°–196°), S.G. 4 918, V.D. 5·1, is converted by H₂SO₄ into cymene. It reduces ammoniacal AgNO₃, form-NaHSO, forms crystalline ing a mirror. C10 H13 NaSO3. Split up by water into the parent substances.

TANTALATES v. TANTALUM, ACIDS OF, AND

THEIR BALTS, p. 639.

TANTALUM. Ta. At. w. c. 182 (exact value not known). Mol. w. not known. Ta has not

been isolated.

Occurrence.—Tantalates occur in a few rare minerals, generally associated with niobates; tantalite contains from 85 to 75 p.c. Ta,O, columbite from 13 to 36 p.c., yttrotantalite from 6 to 47 p.c., and a mineral from Western Australia, recently analysed and called stibiotantalite, c. 52 p.c., Ta₂O₅ (Goyder, C. J. 68, 1076).

History.—A short account of the researches that led to the identification of two distinct oxides in tantalite is given in the article Nio-BIUM (vol. iii. p. 505). Marignac (C. R. 60, 234, 1355) gave the formula Ta₂O₅ to oxide of Ta, which had been represented by Rose as TaO₂. Deville determined the V.D. of the chloride, and deduced the molecular formula TaCl, (C. R. 56, 891). By heating Na, TaF, with Na, H. Rose (P. 99, 69) obtained a black powder, probably a mixture of Ta and oxides of Ta. Berzelius (P. 4, 6) also obtained very impure Ta; Marignac (Ar. Sc. 1868) failed to isolate approximately pure metal.

Preparation of impure Ta.—Very finely-powdered tantalite is fused with three times its weight of KHSO, in an iron or platinum crucible, until completely dissolved. After cooling, the fused substance is powdered and treated with boiling water, whereby sulphates of K, Fe, and Mn are dissolved out; the insoluble portion is washed, and then digested with yellow ammonium sulphide, wherehy sulphides of Sn and W are dissolved, and FeS remains mixed with Ta₂O₅ and Nb₂O₅. The residue is washed thoroughly, and digested with HClAq to remove FeS, and the portion insoluble in HClAq is thoroughly washed with boiling water until it is white. The mixture of Ta2O5 and Nb2O5 thus obtained is dissolved in HFAq in a platinum dish, the solution is heated to boiling, and a quantity of KHF, is added equal to one-fourth of the weight of mixed Ta₂O₅ and Nb₂O₅ used; the liquid is evaporated until 1 g. of the mixed oxides is present in about 7 c.c. and allowed to cool; the prismatic crystals of K. TaF, that separate are washed with cold water until the washings give a yellow pp., without any shade of red, after standing for two hours with tincture of galls. The K, TaF, thus obtained is heated, in a platinum dish, with rather more than its weight of pure conc. H₂SO₄, whereby KHSO₄, HF, and Ta₂O₅ are formed; the HF is removed by heat, and the KHSO₄ by repeated washing with water (Berzelius, P. 4, 6; cf. H. Rose, P. 144, 64, 72).

Lawrence Smith (Am. 5, 44) recommends to warm 5 g. of very finely powdered tantalite, dried at 150°, in a platinum basin with a little water and 8 to 10 c.c. very conc. HFAq, to filter when reaction is completed, add a little water to the filtrate and evaporate nearly to dryness, to warm with excess of conc. pure H2SO4, and, after the acid has been almost wholly removed by heating, to place the residue in c. 500 c.c. dilute HNO₃Aq (L. M., C. N. 51, 289, 304), and boil until Ta₂O₅ and Nb₂O₅ are ppd. By dissolving in HFAq, adding KHF, and proceeding as described above, Ta2Os is obtained free from

Nb₂O₅.

The pure Ta₂O₅ obtained by one of the preplatinum vessel, the solution is heated to boiling, and 70.3 pts. by weight of KHF₂ are added for 100 pts. Ta₂O₃ used. The solution is evaporated and allowed to cool; the crystals of K.TaF, are washed with a little cold water, dried, mixed with c. half their weight of potassium, and heated in an iron crucible, the mixture of K.TaF, and K being covered with KCl. The contents of the crucible, after cooling, are added, little by little, to water; the black powder that separates is washed with water, and then with dilute alcohol and dried (Berzelius, P. 4, 6). The black powder is probably a mixture of Ta and oxides of the metal. Berzelius found that 100 pts. took up 17 pts. O when heated in air, 100 pts. pure Ta require 22 pts. O to form Ta2O3. H. Rose (P. 99, 69) obtained a black powder-probably Ta mixed with oxides—by reducing Na, TaF, by heating with sodium.

Properties and Reactions.—The black powder obtained by Rose was a good conductor of electricity; it glowed when heated in the air, and slowly formed Ta₂O₅; it was insoluble in acids, except in HFAq, by which it was slowly dissolved; heated in a stream of Cl it burnt to TaCl. Oxidation was effected by molten alkalis or alkali carbonates. The S.G. of a specimen containing c. 40 p.c. acid sodium tantalate was

c. 10·8.

The at. w. of Ta was determined by Marignac (A. 140, 153; Suppl. 4, 351 [1865]) by deccmposing K, TaF, and (NH4), TaF, by H, SO4, and determining the quantities of Ta₂O₃ and K₂SO₄ produced. The values obtained varied from 180·1 to 185·2. H. Rose in 1856 (P. 99, 80) analysed TaCl, by decomposing by water, ppg. Ta,O, xH,O by NH,Aq, and estimating Cl in the filtrate. His results showed marked discre-

pancies.

Ta is metallic in its physical properties, so far as may be judged from the impure specimens that have been prepared. Ta₂O₃ dissolves in HFAq, probably forming TaF₃. No salts are known to be formed by replacing the H of an oxyacid by Ta. In all the salts of Ta that have been isolated, other than the haloid compounds, Ta forms part of the negative radicle. the fifth member of the even-series family of Group V.; it is closely related to Nb, and less closely to N, P, V, As, Sb, Di, Er, and Bi. The only compound of Ta whose V.D. has been determined is TaCl_a; in this molecule the atom of Ta is pentavalent (v. Nitrogen Group of Elements, vol. iii. p. 571).

Detection and Estimation.—Tantalum com-

pounds form potassium tantalate when fused with KOH, and the fused mass dissolves in water. The product obtained by fusion with NaOH dissolves in water to a clear liquid only after NaOH has been removed by washing, as sodium tantalate is insoluble in much NaOHAq. Addition of acid to an aqueous solution of an alkali tantalate, followed by boiling, ppts. Ta₂O₈.xH₂O more or less completely. By adding HClAq to a solution of an alkali tantalate, and placing zinc in the solution, no blue colour is produced (cf. Nionium, detection of, vol. iii. p. 506). According to Levy (C. R. 103, 1074), an amethyst colour is produced by adding a very small quantity of Ta2O3 to a solution of resorcin in H₂SO₄Aq. Ta is estimated as Ta₂O₅; the process is sufficiently indicated under Preparation.

Tantalum, acids of, and their Salts. Ta₂O₅ reacts with molten alkalis, and alkali carbonates, to form salts which may be regarded as derived from various hydrates of Ta₂O₅. Two hydrates are obtained, Ta₂O₅.2H₂O and 2Ta₂O₅.3H₂O, by decomposing TaCl₅ by a little water, and by fusing Ta₂O₅ with KHSO₄ and washing with water (v. Tantalum, oxides and Hydrates of, p. 640). The first of these hydrates may be called pyrotantalic acid H₄Ta₂O₇, corresponding with H₄P₂O₇; and the second may be formulated H₄Ta₄O₁₂. The tantalates that have been examined do not seem to be derived from either of these compounds, but from the hypothetical acids HTaO₃, corresponding with HPO₅ and HNO₅, and H₅Ta₆O₁, (=3Ta₂O₅.4H₂O). Tantalates are not produced by neutralising Ta₂O₅.xH₂O, but by fusing Ta₂O₅ with alkalis, or by double decomposition from alkali salts.

Tantalates. The tantalates belong to the form xTa, 0, yMO, where $M = (NH_4)_2$, Mg, Hg, Kg, Ag2, or Na2; of those that have been fairly fully examined, some correspond with the metaphosphates and meta-niobates $M^{\text{I}}XO_8$, and others belong to the more complex form M^{I}_8 Ta, 0_{19} . Some of the alkali tantalates are insoluble. Fluotantalates and fluoxytantalates are also known (v. infra). The tantalates have been investigated chiefly by H. Rose (P. 100, 417) and Marignac (Bl. [2] 6, 111, 118).

Ammonium tantalate. A pp. is obtained by adding NH₁Cl to solution of Na₂Ta₂O₁₀.xH₂O, but the composition of the pp. is not known with certainty.

Magnesium tantalate. By adding MgSO, Aq to solution of Na, Ta, O_{12} , xH₂O a crystalline pp. was obtained which, after drying at 100°, had the composition Mg, Ta, O_{11} , 9H₂O. A crystalline Mg tantalate was obtained by Joly (O. R. 81, 266, 1266) by fusing Ta₂O₃, with MgCl₂.

Mercurous tantalate. A brown, amorphous compound, perhaps Hg.Ta.O., xH2O, is formed by adding HgNO.Aq to solution in water of

Na, Ta, O19. xH2O.

Polassium tantalates. (1) K₈Ta₆O₁₉.16H₂O; this salt is obtained by fusing Ta₂O, with KOH, dissolving in water, and crystallising. (2) KTaO₃: this salt, potassium metatantalate, is formed by heating the other tantalate, alone or with (NH₃)₂CO₃₉ and then treating with water. Silver tantalate. The yellowish white pp.

obtained by adding solution of a salt of Ag to solution of Na₈Ta₈O₁₀.xH₂O, and drying at 100°, has the composition Ag₈Ta₈O₁₀.3H₂O.

Sodium tantalates. The meta-salt

Sodium tantalates. The meta-sati NaTaO₃ is obtained by fusing Ta_2O_5 with Na₂CO₃, and washing with water. A salt of the composition Na₈Ta₅O₁₉.xH₂O (x = 25 and 30) is prepared by fusing Ta_2O_5 with NaOH, dissolving in water, and crystallising.

FLUOTANTALATES. (Tantalifluorides.) These salts may be regarded as compounds of TaF₃ with metallic fluorides, or as metallic derivatives of the hypothetical acid H₂TaF₇; they are prepared by dissolving Ta₂O₃ in HFAq, adding metallic fluorides, and crystallising; some of them are formed by dissolving Ta₂O₄ and a metallic oxide in HFAq (Marignac, A. Ch. [4] 9, 276; Berzelius, P. 4, 6).

Ammonium fluotantalate (NH₄)₂TaF, is obtained by evaporating a solution of NH₄F in solution of TaF, in HFAq, and evaporating; the salt is crystalline and easily soluble in

water.

Copper fluotantalate CuTaF, forms blue, deliquescent, rhombic prisms; it is prepared by dissolving CuO and Ta₂O₅ in excess of HFAq,

and evaporating.

Potassium fluotantalate K₂TaF, forms white needles by dissolving KHF₂ in solution of Ta₂O₃ in HFAq, evaporating, and crystallising from hot water. The salt is very slightly soluble in cold water, but dissolves easily in hot water. On long boiling, K₂TaF,Aq gives a white pp., probably having the composition K₄Ta₄O₅F₁₄ (=2TaF₅.Ta₂O₅.4KF). By dissolving K₂TaF, in warm 4 p.c. H₂O₂Aq with a little HFAq and allowing to cool, Piccini (Zeit. flur anorg. Chemie, 2, 21) obtained crystals of the fluo xy tantalate K₂TaO₂F₅.H₂O (= TaO₂F₅.2KF.H₂O).

Sodium fluotantalate Na, Taf, H₂O; obtained, as white crystals, by dissolving Na, Ta₂O₁₉ in HFAq, evaporating, and drying at 100° the

salt that separates.

Zinc fluotantalate ZnTaF, 7H₂O; a deliquescent salt, obtained by dissolving ZnO and Ta₂O, in excess of HFAq and crystallising.

Tantalum, alloys of. By heating K₂TaF, with Al, and washing with HClAq, Marignac (P. 100, 145) obtained a grey powder, S.G. 7.02. By heating to whitness, in a carbon crucible, a mixture of Ta₂O₃ and iron filings, an alloy of Ta and Fe was obtained resembling pig iron (Gahn, Berzelius a. Eggertz, S. 16, 437).

Tantalum, bromide of, TaBr_s. A yellowish, crystalline compound, obtained by heating a mixture of dry Ta₂O₅ and C in vapour of Br, and removing excess of Br by a long-continued stream of dry CO₂; decomposed by water to HBrAq and Ta₂O₅.xH₂O (H. Rose, P. 90, 456; 99, 75). The conditions of preparation are similar to those in making TaCl₂(q. v.).

Tantalum, carbide of. By heating TaN (v. Tantalum nitrade) with C to the melting-point of steel, N is given off and the nitride is partly converted into brass-yellow coloured

TaC₂ (Joly, Bl. [2] 25, 206).

Tantalum, carbonitride ef. Joly (Bl. [2] 25, 206) obtained a substance, to which he gave the formula 10TaC.TaN, by very strongly heating a mixture of Ta₂O₂ with C and soda.

Tantalum, chloride of, TaCl, Mol. w. c. 359 (not determined with great accuracy, as at. w. of Ta is doubtful). V.D. 185 at 360° (Deville a. Troost, C. R. 64, 294). Melts at 211°, and boils at 242° at 753 mm. pressure (D. a. T., l.c.). Only one chloride of Ta has been isolated.

Preparation.—About 5 g. pure dry Ta2O, is mixed with a considerable excess of dry sugar or starch, and the mixture is completely charred by heating in a closed crucible; the charred mass is broken into small pieces, which are heated to redness and placed, while red hot, in a rather wide tube of hard glass that is quickly heated to redness while a stream of thoroughly dried CO₂ is passed through it as long as any trace of moisture is given off from the contents of the tube; the tube is then allowed to cool, the current of dry CO2 being maintained; when the tube is cold a stream of dry Cl is passed through it, and when every part of the apparatus is filled with Cl (but not until then) the contents of the tube are heated. TaCl, forms immediately behind the carbonaceous matter, as a pale-yellow solid; when the reaction has entirely ceased and CO is no longer evolved, the TaCl, may be sublimed into another part of the tube, a plentiful stream of dry Cl being maintained during the process. If sublimation is attempted before the whole of the Ta,O, has been chlorinated, a part of the TaCl, is decomposed by the CO, with re-formation of Ta,O₅. If there should be any air in the tube during the process a white sublimate is formed which is probably an oxychloride (H. Rose, P. 90, 456; 99, 75). Should the Ta₂O₅ used contain any WO₂, the sublimate obtained is reddish; by gently heating, the greater part of the red WOCl, may be removed, as it is more volatile than TaCl,.

Demarcay obtained TaCl, by passing vapour of CCl, over Ta₂O₃ heated to redness (C. R. 104,

Properties and Reactions.—Pale-yellow prismatic needles, melting at 211°, and boiling at 242° under the pressure of 753 mm.; begins to vaporise at 144°; V.D. 185 at 360° (Deville a. Troost, C. R. 64,294). Decomposes in ordinary air, giving off HCl and becoming covered with a crust of crystalline Ta₂O₃. Decomposed entirely by water to HOlAq and Ta₂O₃.xH₂O. Conc. H2SO, causes evolution of HCl, and forms somewhat turbid solution from which Ta₂O₃.xH₂O separates on boiling; by adding water to this solution, and boiling, the whole of the Ta is precipitated as Ta,O,xH,O. Conc. HClAq reacts similarly to H.SO., but only a portion of the Ta₂O₂.xH₂O is ppd. on adding water and boiling. TaCl, is partly dissolved by heat-ing with KOHAq. TaCl, is soluble in absolute alcohol; H2SO, Aq does not ppt. Ta2O, xH2O from this solution.

No double compounds of TaCla with KCl or NaCl-similar to those of TaF, have been obtained.

Tantalum, fluoride of. Ta2O3.xH2O dissolves in HFAq; neither boiling, nor adding H₂SO₄Aq to the solution, ppts. Ta₂O₄. The solution in HFAq gives off vapours on evaporation that probably contain TaF,; by evaporating at a low temperature, Rose obtained crystals that dissolved easily in water, and were partly vaporised when heated, leaving some Ta₂O₃ (P. 90, 456; 99, 75). By evaporating in vacuo a solution of Ta₂O₃.xH₂O in HFAq, Marignac obtained a white amorphous mass—probably an oxyfluoride -and then small crystals that were likely

TaF₅.
TaF₅ combines with some metallic fluorides,

forming fluotantalates (q.v., p. 639).

Tantalum, haloid compounds of. The only haloid compounds of Ta that have been isolated are TaCl, and TaBr. The former has been gasified, and the formula is molecular. TaF, gasified, and the formula is molecular. also probably exists in solution of Ta2O, xH2O in HFAq. There is no reaction between I and Ta₂O₄ mixed with C even at a very high temperature. TaOl, and TaBr, are readily decomposed by water, with formation of Ta₂O₅.xH₂O and HXAq. Double compounds of TaF, with some metallic fluorides are known; they belong to the form M'2TaF, (v. Fluotantalates). Oxyhaloid compounds probably exist, but none has been isolated with certainty.

Tantalum, nitrides of. TaCl, absorbs NH, at the ordinary temperature; by heating the product in NH₃, Joly (Bl. [2] 25, 206) obtained an amorphous, yellow solid, to which he gave the formula Ta₃N₅. By heating this yellow solid to redness in very dry NH_s, a black solid was obtained which, after washing with water and drying, had the composition TaN. This black solid conducts electricity; heated in air it burns to Ta₂O₅; NH₃ is given off by the action of molten KOH; it is insoluble in acids, except in a mixture of HFAq and HNO₃Aq (Joly, *l.c.*; cf.

H. Rose, P. 100, 166).

Tantalum, nitrocarbide of; v. TANTALUM CARBONITRIDE, p. 639.

Tantalum, oxides and hydrated oxides of. The oxide Ta₂O₅ has been isolated, and also a hydrate of this oxide Ta₂O₅.2H₂O, and probably also another hydrate 2Ta,O, 3H,O. The existence of another oxide, TaO2, is doubtful.

TANTALUM PENTOXIDE Ta,o,. (Tantalic anhydride. Tantalic oxide.) The preparation of this compound from tantalite is described at the the beginning of this article under Preparation of tantalum (p. 638). Ta₂O₅ is also obtained by decomposing TaCl, by water, or by adding H,SO,Aq to solution of a tantalate and boiling, and then heating the Ta₂O₃.xH₂O thus ppd. Ta₂O₃ is a white powder; it has not been fused; by heating with boric acid or microcosmic salt in a porcelain oven it is obtained in rhombic prisms (Ebelmen, A. Ch. [3] 33, 34; Nordenskjold a. Chydenius, J. 1860. 145). S.G. c. 7·6 (Marignac, Ar. Sc. 1868; Deville, C. R. 56, 894). Ta₂O, is insoluble in acids except HFAq; after being very strongly heated, it is insoluble in HFAq. Ta,O, dissolves in molten alkalis, also in molten KHSO₄ (v. Tantalates, p. 639).

HYDRATES OF TANTALUM PENTOXIDE. By decomposing TaCl, by water, Ta2O, xH2O is obtained, which, after very thorough washing with water, then with NH, Aq to remove adhering HCl, and finally with water, and drying at 100°, gives off from 6 to 7.8 p.c. water when heated to redness. The formula 2Ta,O,.3H,O requires 5.7 p.c. H,O, and the formula Ta,O,.H,O requires 7.5 p.c. H₂O. It is doubtful whether one or more than one definite hydrate exists in this pp. When SO, is passed into a solution of Na, Ta,O, a pp. of Ta,O, xH,O is obtained, which, when thoroughly washed and dried at 100°, is said to be $Ta_2O_s.2H_2O = H_4Ta_2O_7$ (H. Rose, P. 100, 417). Strong acids, such as H_2SO_4 , HNO₃, and HCl, ppt. $Ta_2O_s.xH_2O$ apparently in combination with the acid used as precipitant from solutions of alkali tantalates; weak acids ppt. insoluble alkali tantalates.

TANTALUM DIOXIDE TaO₂. (Tantalum tetroxide [Ta₂O₄].) The isolation of this oxide is doubtful. Berzelius obtained a brown powder by heating Ta₂O₅, to redness in a carbon crucible; it was not dissolved by any acids, not even by a mixture of HNO₈Aq and HFAq; when strongly heated in

air it gave Ta_.O₅ (P. 4, 20).

Tantalum, oxyfluoride of. The white powder formed by decomposing K₂TaF₇ by boiling water may be an oxyfluoride, according to

Marignac (A. Ch. [4] 9, 276).

Tantalum, salts of. No salt has been isolated by replacing the H of acids, except HCl and HF, by Ta. When alkali tantalates are decomposed by strong acids such as HCl, HNO₃, or H₂SO₄, the ppd. Ta₂O₃.xH₂O retains some of the acid used as precipitant; compounds of Ta₂O₅ with these acids are perhaps formed.

Tantalum, sulphide of. By strongly heating Ta₂O₅ in CS₂ vapour, also by heating to redness a mixture of vapour of TaCl₃ and H₂S₄ a yellowish black solid is obtained, which has the composition TaS₂ according to Berzelius (P. 4, 6; v. also H. Rose, P. 99, 575). Roasted in air, this solid gives Ta₂O₅; heated in Cl it produces TaCl₅ and S₂Cl₂; conc. HNO₅Aq oxidises it to Ta₂O₅.xH₂O and H₂SO₄Aq. TaS₂ does not combine with alkali sulphides (R., l.c.); fused with KOH, it forms K₂S and potassium tantalates. M. M. P. M.

TARCHONYL ALCOHOL $C_{50}H_{102}O$ (?). Obtained from the leaves of Tarchonanthus camphoratus by extracting with alcohol (Canzoneri a. Spica, G. 12, 227). Silvery scales, insolwater and ether. Converted by PCl₃ into a chloride [70°] crystallising in small plates (from

alcohol)

TARCONINE v. vol. iii. p. 496. TARNINE v. vol. iii. p. 497. TARTARIC ACID C₄H₆O₆ i.e.

CO.H.CH(OH).CH(OH).CO.H. Dextro-tartaric acid. Di-oxy-succinic acid. [135°]. S.G. 1-764 (Schiff, A. 113, 189); 1-74 (Buignet, J. 1861, 15); $\frac{1}{4}$ 1-7594 (Perkin). S. 115 at 0°; 132 at 15°; 343 at 100° (Leidie, Fr. 22, 269; C. R. 95, 87; cf. Schiff, J. 1859, 41; Maisch, J. 1865, 392). S. (alcohol) 25-6 at 15°. S. (ether) 4 at 15° (Bourgoin, Bl. [2] 29, 244; A. Ch. [5] 13, 400). [a]_D = 15·06 - 131p in a p per cent. solution (Landolt, B. 6, 1075; Lippmann, B. 24, 3300). H.F. 372,000 (Von Rechenberg). Heat of solution: -3454 at 9° (Pickering, C. J. 51, 376).

Occurs free or as K or Ca salt in grape-juice, tamarinds, unripe mountain-ash berries, madderroot, potatoes, Jerusalem artichokes, sorrel, gherkins, mulberries, pine-apples, black pepper, the leaves of Chelidonium majus, the bulbs of Scilla maritima, in beet-juice, and in many other

plants.

Formation.—1. Mostly together with racemic acid, by oxidation of saccharic acid, dextrose, cane-sugar, milk-sugar, starch, gum arabic, and sorbin by nitric acid (Dessaignes, A. Suppl. 2, 242; Hornemann, J. 1863, 381; Kiliani, A. 205, 175.—2. By the action of sodium-amalgam on Vol. IV.

an alcoholic solution of oxalic ether (Debus, A. 166, 124; C. J. 24, 376). The product is probably inactive tartaric acid.

Preparation.—Powdered chalk is added to a boiling solution of cream of tartar; the filtrate is ppd. by calcium chloride, and both pps. decomposed by the proper quantity of boiling dilute H₂SO₄. The filtrate from CaSO₄ is evaporated to crystallisation. It may be purified by saturating with CaCO₂, digesting with ZnCl₂, and decomposing the zinc salt by H₂S (Ficinus, Ar.

Ph. [8] 14, 310).

Properties. - Monoclinic prisms, with hemihedral facets; a:b:c = .785:1:.805; $\beta = 79^{\circ}$ 48'. Dextrorotatory, the rotation being dependent on the concentration of the solution, and being also greatly affected by the presence of alcohols, acids, and other substances in the solution (Landolt, B. 13, 2329; Biot, J. 1850, 169; Pribram, M. 9, 485; B. 22, 6; Long, Am. S. 36, 351; Thomsen, J. pr. [2] 35, 145). Sodium molybdate in the proportion of Na, MoO, to 2C, H, O, increases the rotation 37 times, while ammonium molybdate (1 mol.) increases the rotation of tartaric acid (3 mols.) 56 times (Gernez, C. R. 104, 783; 105, 803). Lithium and magnesium molybdate also increase the rotation. Sodium tungstate also increases the rotation, the maximum effect being 22 times (Gernez, C. R. 106, 1527; 108, 942). Crystals of tartaric acid are strongly pyro-electric. An aqueous solution of tartaric acid is ppd. by baryta, lime, and lead acetate. Potassium salts, when the solution is not too dilute, form a crystalline pp. of KC4H5O6; the formation of the pp. being promoted by rubbing the sides of the vessel with a glass rod and by the addition of alcohol. CaCl₂ ppts. CaA'₂ from neutral solutions of tartrates; the ppn. is hindered by ammonium salts. Calcium tartrate dissolves in KOHAq and is re-ppd. in gelatinous form on boiling. Reduces ammoniacal AgNO₂, forming a mirror; the solution then contains oxalic acid (Claus, B. 8, 950). Tartaric acid prevents the ppn. by alkalis of the oxides of Al, Bi, Ni, Co, Cr, Cu, Fe, Pb, Pt, and Zn, even on boiling (Aubel a. Ramdohr, A. 103, 33; Grothe, J. pr. 92, 175). 1 mol. acid prevents the ppn. of 1 mol. Cu(OH)2. A solution of tartaric acid, coloured by a drop of K2CrO4, gradually becomes colourless (difference from citric acid) (Salzer, B. 21, 1910). A saturated solution of potassium bichromate is slowly turned coffee-brown by tartaric (but not by citric) acid (Cailletet, C. C. 1879, 14). Boiling alkaline KMnO₄ is turned green, and finally brown, by tartaric acid, but only green by citric acid. Decolourises KMnO, in acid solution. When H SO, is present, twice as much KMnO, must be added to produce a permanent red colour as when it is absent, for in the latter case MnC, H,O, is formed (Fleischer, B. 5, 850). Tartaric acid gives an odour of burnt sugar when heated till carbonised. A crystal added to conc. H2SO, containing 1 p.c. resorcin gives a red colour on warming (difference from citric and maleic acids) (Mohler, Bl. [8] 4, 728).
To detect small quantities of tartaric acid in

To detect small quantities of tartaric acid in presence of citric acid. Add about 1 g. citric acid to a 20 p.c. solution of molybdate of ammonia (1 c.c.), then two or three drops of a dilute solution of H₂O₂, and heat for three minutes on the water-bath. If the citric acid is pure the

solution remains yellow. If 0.1 p.c. or more of | tartaric acid is present, the liquid become blue

(Crismer, Bl. [3] 6, 23).

Reactions.—1. Melts at 135°, changing to the isomeric metatartaric acid. The metatartaric acid may be reconverted into tartaric acid by boiling with water (Grosjean, C. J. 43, 334). 2. At 140°-150° water is eliminated and ditartrylic acid C₆H₁₀O₁₁ is formed. On further heating soluble tartaric anhydride (tartrelic acid), and then insoluble tartaric anhydride, are formed. On further heating the products of distillation (from 250 g.) are pyruvic acid (9 g.), pyrotartaric acid (2 g.), formic acid (4 g.), resins, aldehydes and volatile acids (2 g.), and tarry bodies (3 g.) (P. Liebermann, B. 15, 428). 'Dipyrotartracetone' C_aH₁₂O₂ (230°), a colourless liquid, with aromatic odour, is among the products of the distillation of tartaric acid. It combines with bromine (Bourgoin, C. R. 86, 674). - 8. By heating tartaric acid (10 pts.) with water (1 pt.) at 175° it is converted into racemic acid and inactive tartaric acid. At 165° there is a greater yield of the inactive acid (Jung-fleisch, J. 1872, 515). Heated with more water at 150° it yields CO₂ and pyrotartaric acid (Wedard, C. C. 1888, 889).—4. Potash-fusion yields acetic and oxalic acids. -5. Oxidation by CrO₂, KMnO₄, PbO₂ or MnO₂, and H₂SO₄ yields CO₂ and formic acid. The rate of oxidation has been studied by Krutwig (Z. P. C. 2, 787). By slow oxidation by HNO, tartronic acid may be obtained.-6. Reduced by HI and P to malic and succinic acids. - 7. PCl, forms chloro-fumaric chloride.—8. FeSO₄ at 100° yields iso-arabic acid C_aH₁₀O₃, a thick dextrorotatory syrup, [a]_D = 20°, yielding the salts CaA', 2aq, Ca₂OA', 2aq, PbA', and Pb₂A', O₂ (Ballo, B. 22, 750).—9. Chloral forms C₈H₄Cl₆O₆ [124°], crystallising from chloroform in small needles, insol. water, sol. warm alcohol and ether (Wallach, A. 193, 46).-10. On submitting tartaric acid to electrolysis and treating the syrupy mass with phenyl-hydrazine, glyoxal-osazone [160°] and glyoxal-carboxylic osazone [218°] are obtained (Friedel a. Combes, Bl. [8] 8, 770).—11. On adding powdered tartaric acid to fused glucose tartaric glucoside is obtained (Guyard, Bl. [2] 41, 291; cf. Berthelot, A. Ch. [3] 54, 78).—12. m-Amido-benzoic acid at 160° forms 'tartrylo-dibenzamic'acid C18H18N2O8 i.e. $(CO_2H.C_6H_4.NH.CO)_2C_2H_2(OH)_2$, a colourless powder, insol. water, sol. hot alcohol. alkaline salts form yellow solutions in which Cu(OAc), ppts. C₁₈H₁₄(CuOH),N₂O₅. On heating to 190° it yields the anhydride C₁₈H₁₄N₂O₇, a greenish-yellow powder, converted by an alcoholic solution of aniline into tartranil dibenzamic

CO₂H.C₆H₄.N<CO_{CH.CH}(OH).CONHPh [200°] (Schiff, A. 282, 156; G. 16, 28). Another product of the action of tartario acid on mamido-benzoic acid is 'tartryl-benzamic' acid

CO₂H.CH(OH).CH(OH).CO.NHC₄H₄.CO₂H, which crystallises from water in pale-yellow aggregates, yields an acetyl derivative and an anhydride 'tartranbenzamic' acid C₁₁H₂NO₂, crystallising from water in green plates, forming C11H,BaNO, and C11H,CuNO, and converted at 210° in 'benzamtartridic' acid by dehydration. A warm alcoholic solution of aniline converts 'tartranbenzamic' acid into 'tartranilbenzamic'

acid [245°], which yields an acetyl derivative [1980]. 'Tartryl-benzamic' acid heated with dry amido-benzamide at 140° forms the compound C₂H₂(OH)₂(CO.NH.C₆H₄.CONH₂)₂, a white powder, insol. water, sl. sol. alcohol, and yieldring C₁₈H₁₅CuN₃O₇ aq.—13. Glycerin at 100° yields the compounds C₃H₃(OH)₂·O.C₄H₃O₅ and C₃H₅(OH)(OC₄H₅O₅)₂ (Desplats, J. 1859, 500). Glycerin at 140° gives C₁₁H₁₅O₁₂ and C₁₈H₂₅O₁₉.—14. Emitting to 100° forms of H. O. Third the strong o 14. Erythrite at 100° forms $C_{12}H_{18}O_{14}$, which gives Ca₃(C₁₂H₁₃O₁₄)₂ 3aq (Berthelot, A. Ch. [8] 54, 84). 15. Quercite at 120° forms C₂₂H₃₂O₇, which gives Ca₃C₂₂H₂₆O₇ 2aq. — 16. Pinite at 120° forms Ca₂C₂₂H₂₆O, 2aq. — 10. Finite Bt 120 Iorians $C_{30}H_{36}O_{45}$, which gives $Ca_{3}C_{30}H_{30}O_{35}$.—17. Mannite at 120° forms $C_{30}H_{30}O_{35}$, which yields $Mg,O_{4}(C_{30}H_{30}O_{35})$ 30aq and $Ca_{3}C_{30}H_{30}O_{35}$ 6aq.—18. Dulcite forms $C_{14}H_{20}O_{15}$, which gives $Ca(C_{14}H_{19}O_{15})_{2}$ 4aq.—19. Phenyl hydraxine (2 mols.) at 140° forms $C_{16}H_{18}N,O_{4}$ i.e. (2 mols.) at 140° forms C₁₆H₁₈N₁O₄ i.e. C₂H₁O₂(CO.N₂H₂Ph)₂[226°] (Bülow, A. 236, 196); [c. 240°] (Fischer a. Passmore, B. 22, 2734), crystallising from alcohol in plates.

Estimation.—Methods for estimating tartaric acid have been described by Scheurer-Kestner, C. R. 86, 1024; Berthelot, Fr. 8, 216; Vogel a. Braun, Fr. 7, 149; Kissel, Fr. 8, 409; Fleischer, Fr. 12, 328; Oliveri, G. 14, 453; Piccard, Fr. 21, 424; C. Schmitt a. Hiepe, Fr. 21, 539; Amthor, Fr. 21, 195; Nessler a. Barth, Fr. 21, 60; 22, 159; Kayser, Fr. 22, 123; 23, 29; Musset, Fr. 24, 279; Ferrari, Fr. 24, 279; Warington, C. J. 28, 25; Grosjean, C. J. 35, 841; Klein, Fr. 24, 379; Bornträger, Fr. 25, 827; 26, 699; Gantter, Fr. 26, 714; Von Lorenz, Fr. 27, 8; Philips, Fr. 29, 577; Goldenberg, Fr. 22, 270; Heidenhain, Fr. 27, 681; Weigert, Fr. 23, 357; Haas, C. C. 1888, 1045; Ward, Ph.

[3] 19, 380.

Salts (Berzelius, P. 19, 305; 36, 4; A. Ch. [2] 67, 303; Werther, J. pr. 32, 383; Dumas a. Piria, A. Oh. [3] 5, 353; De la Provostaye, A. Ch. [3] 3, 129).—(NH₄)₂A". [a]_D = 33·7° in a 1·2 p.c. solution (Sonnenthal, M. 12, 603). Efflorescent monoclinic prisms, v. sol. water; $a:b:c=868:1:1\cdot244\alpha=88^\circ9'$. Giveoff NH_2 in air. Yield succinic acid onfermentation (Koenig, G.11, 180). -(NH₄)HA". [α]_D = 25.7° (Landolt). Minute laminæ, sl. sol. cold water. Forms with ammonium malate (NH₄)HA"(NH₄)C₄H₅O₅, crystallising in prisms, S. 9 (Pasteur, J. 1853, 415).—K, A" aq. [a]_D = 25.5° in a 2 p.c. solution. S. 133 at 2°; 158 at 28° (Osann). V. sl. sol. hot alcohol. Monoclinic crystals; a:b:c = .402:1:1.109; $\alpha = 75^{\circ} 12'$.—KHA". Cream of tartar. $[\alpha]_D = 22^{\circ}$ in a 4 p.c. solution. 100 g. solution contain $^369 + 000569t^2$ grammes at t° (Blarez, C. R. 112, 434, 808). S. ·25 at 0°; ·4 at 10°; ·55 at 20°; 1.13 at 40° (Chancel, C. R. 60, 408; cf. Alluard, C. R. 59, 500). Obtained by crystallisation of argol, which is deposited during vinous fermenargot, which is deposited during vincous termientation. Trimetric crystals; a.b.c = .712:1:.737. Insol. alcohol.—Na,A" 2aq. [a]_b = 31° in a 1 p.c. solution (Sonnenthal; cf. Thomsen, J. pr. [2] 34, 80; 35, 145). S. 29 at 6°; 44 at 24°; 66 at 42.5° (Osann). Trimetric prisms; a.b.c = .770:1:.937. V. sol. hot water, insol. alcohol. -NaHA". $[\alpha]_p = 24.3^\circ$ in a 1.3 p.c. solution (S.). —NaHA"aq (Dumas a. Piris, Å. 44, 80). S. 11 in the cold; 55 at 100°. Minute crystals, with hemihedral facets. Insol. alcohol.—Li₂A". $[a]_D = 87.5^{\circ}$ in a .7 p.c. solution (S.). Deliques-

cent.—LiHA"1\frac{1}{2}aq. Small crystals, v. sol. water (Dulk, Schw. J. 64, 180, 193; A. 2, 47).—LiHA"aq.—Li4H_A", TeO 2aq. Needles (Klein, A. Ch. [6] 10, 118; C. R. 102, 47).—Hydroxylamine salt (NH₄O)₂A". Very thin plates (Lossen, A. Suppl. 6, 233).—NH₄NaA"4aq. Trimetric crystals, with hemihedral facets, isomorphous with KNaA" 4aq; a:b:c = .823:1: 420. [a]_j = 26°. [a]_b = 32.7° (Landolt). S. 26 at 0° (Pasteur, J. 1849, 309). — LiNaA" 2aq. — Na₂A" 2TeO 2aq. — K_2A "TeO aq. — (NH₄)KA". Monoclinic crystals, isomorphous with K_2A ". Give off NH₃ in air. V. sol. water. [a]_D = 31° (L.).—KLiA"aq.—KNaA"4aq. Rochelle salt. S. 30 at 3°, 66 at 26° (Osann). Heat of solution —6290 at 15°. The heat of solution of KNaA" is -2989 (Pickering, C. J. 51, 351). Large trimetric prisms, with hemihedral facets. Melts at 70°-80° in its water of crystallisation, and becomes anhydrous at 215° (Fresenius, A. 53, 234). [a]_D = 29.7° (L.). Molecular rotation: Long, Am. S. 36, 351.—K₂A"₂TeO (dried at 100°).-RbHA". Trimetric prisms, isomorphous with the cosium salt; a:b:c = .726:1:.695 (J. P. Cooke, Am. S. [2] 37, 70). S. 1·18 at 25°; 11·7 at 100° (Allen, Am. S. [2] 34, 367).—NaRbA" 4aq. Isomorphous with Rochelle salt (Piccard, J. 1862, 125).—Cs.,A". Trimetric prisms. S. 10 at 25°; 100 at 100°.—Tl.,A" \(\frac{1}{2}\) aq. \(\alpha \)] = 4.8° in a 5 p.c. solution at 20°. Sl. sol. water and alcohol.— Solution at 20°. St. sol. water and account TIHA''. [α] = 12° in a 1 p.c. solution at 20°. Small flat prisms (De la Provostaye, A. Ch. [3] 3, 129; Kuhlmann, J. 1862, 188).—TINAA"4aq. At 20° [α] = 0° in a 5 p.c. solution and 6.5° in α a 20 p.c. solution.—TlLiA"aq. [a] = 9.5° in a 5 p.c. solution at 20° .—TlKA". [a] = 10° in a 5 p.c. solution at 20° .—Tl(NH₁)A". [a] = 10° in a 5 p.c. solution at 20° .—BaA". White amorphous pp., becoming crystalline. S.G. 214 2.973 (Clarke, Am. 2, 174).—BaA"aq (Dulk). S. (amorphous) 1.2; (crystalline) 077 (Vogel a. Reischauer, J. 1859, 288). K,BaA", 2aq. Powder, sl. sol. water.—Na,BaA", 2aq. Needles, sl. sol. water, more soluble in a solution of Rochelle salt. Ppd. on mixing a solution of Rochelle salt with BaCl₂.—BaA"TeO (dried at 100°).— SrA" 4aq. Monoclinic prisms. S. 7 at 16°. S.G. 1966 (Clarke).—SrA" 3aq (Marignac, Ann. M. [5] 15, 280).—(NH₄)₂SrA"₂ 12aq. Thin plates.-K₂SrA" 2aq.—Na,SrA" xaq.—CaH₂A"₂. S. 71 at 15.6°. Trimetric crystals.—CaA" 4aq. Occurs in grapes. Ppd. as white crystalline powder on adding $CaCl_2$ to a solution of K_2A'' . Small trimetric prisms. a:b:c = .872:1:.908 (Anschütz, A. 226, 191). S. .016 at 15°: .3 at 100° (Mohr, J. 1865, 893). Forms, with calcium malate, the double sait CaA"CaC,H,O, 6aq, crystallising in double salt CaA"CaC,H,O, 6aq, crystallising in needles, S. 1-25 at 17° (Ordonneau, Bl. [3] 6, 262).—MgA"4aq. S. 8 at 16°. [α]_p=36°.—Mg,C,H₂O, 3aq. S. 924. Ppd. by adding NH, to a solution of MgA" (Mayer, A. 101, 166).—MgNa,A".210aq.—MgK,A"8aq.—MgH,A". S. 2 at 16°.—BeA"3aq (Atterberg, Bl. [2] 21, 162).—BeK,C,H,O, 3aq. Prisms (Toczynski, Z. 1871, 277).—K(SbO)A" $\frac{1}{2}$ aq. Tartar emetic. Prepared by boiling Sb₂O₃ (3 pts.) with cream of tartar (4 pts.) and water. Octahedra; a:b:c (4 pts.) and water. Octahedra; a:b:c = 956:1:1·054. [a] = 141° in a 4 p.c. solution at 20°. S.G. 2·6. Efflorescent. At 200° it becomes K(SbO)C4H2O5. S. 5 at 8°; 8 at 21°; 52

at 100°. . 05 to .1 g. produce vomiting. Its solution reddens litmus, and gives a crystalline pp. of K(SbO)A" with alcohol. Its solution is ppd. by mineral acids, by alkalis, and by H.S. HgCl₂ is reduced by it to calomel. Heat of formation: Guntz, C. R. 104, 699. Volumetric estimation of Sb in tartar emetic (Dunstan a. Boole, Ph. [3] 19, 385). — K(SbO)A"H₂A" 2½ aq. Efflorescent prisms. Alcohol added to its aqueous solution ppts. tartar-emetic, leaving tartaric acid in solution (Knapp, A. 32, 76).—K(SbO)A"3KHA".

Mammellated groups of needles, v. sol. water.— (K(SbO)A"), NaNO3 (Martenson, J. 1869, 539).— Na(SbO)A" ‡aq. Trimetric, hygroscopic prisms; a:b:c=:922:1:1.08. — (Na,A"),2Sb(OH), 8aq. Amorphous, v. sol. water (Clarke a. Evans, B. 16, a:b:c = .922:1:1.08. — 2385). — NH₄(SbO)A'' ½aq. — NH₄(SbO)A'' 2½aq (Berlin, A. 64, 359).—Rb(SbO)A'' ½aq. Isomorphous with tartar-emetic (Grandeau, A. Ch. [3] 67, 155).—Tl(SbO)C,H₁O₄ aq. [a] = 100° in a 2 p.c. solution at 20° (Long, Am. S. [3] 38, 264).
—Ag(SbO)A" aq. Crystals (J. P. Cooke, Am. S. [3] 19,393).—Ba(SbOA"),2 aq.—Cd(SbOA") 2aq. Crystallises from a solution of tartar-emetic mixed with calcium nitrate in trimetric forms (Marignac, Ann. M. [5] 15, 280).—Sr(SbOA")2 Hexagonal crystals. — Sr₂(SbOA")₂(NO₃)₂ 12aq. Very soluble crystals (Kessler, P. 75, 410).— Pb(SbOA'), 4aq. — Be, 3b2, (C, H, 2O,). — Sb(HA''), 4aq. Needles, v. sol. water (C. a. E.). — Sb2A'', 6aq. — (SbO)HA'' aq (Clarke a. Stallo, B. 13, 1788).— (SbO)HA'' (Guntz, C. R. 104, 1768).— (SbO)HA'' (ShO)HA'' (S White granular pp., 850). - C, H, O, Sb, O, aq. formed by adding alcohol to a solution of Sb.O in aqueous tartaric acid (Berzelius). Converted by heat into SbOC,H₂O₅,-(C,H₅O₁₎,Sb₂O₅6aq (Péligot, A. Ch. [3] 20, 289).-(C,H₂O₄)₂Sb₂O₅ (dried) (P.). - (C₄H₄O₄)₂Sb₂O₅H₂O. Crystals (Gunts, A. Ch. [6] 13, 396).—Berberine salt $C_2H_{18}NO_4(SbO)A''$ (Stenhouse, J. 1863, 452).—Brucine salt $C_2H_{28}N_2O_2(SbO)A''$.—Quinine salt $C_2H_{28}N_2O_2(SbO)A''$. (Clarke, R. 15, 1540).— Atropine salt C₂H₂M¹O₂(SbO)A'' (Marke, A. 10, 1840).—
Aniline salt C₃H₂N(SbO)A''. Long white prisms.
S.G. 1·89 at 11°.— NH₄(AsO)A'' ½aq. Large efflorescent trimetric crystals; a:b:c = .876:1:.694. Got by dissolving As₂O₃ in NH,HA" (Mitscherlich; Werther, J. pr. 32, 409).—K(AsO₂)A" 23aq. Got by dissolving arsenic acid in a solution of cream of tartar (Pelouze, A. Ch. [3] 6, 63).— (Sr(AsOA'')₂)₂NH₄NO₃12aq. Large trimetric crystals (Marignac).—Bi₂A'', 6aq. Small crystals, decomposed by water (Schneider, P. 88, 55). - K(BiO)C, H₂O₆ (Schwarzenberg, A. 61, 244). -Bi₂K(BiO)(C,H₂O₆)₂ (Frisch, J. 1866, 401).--K(BO)A" (dried at 100°). Potassium borotartrate. Amorphous mass, v. sol. water, insol. alcohol. At 100° it becomes K(BO)C,H,O, Purgative. • [a] = 58° in a 20 p.e. solution at 20°.

- K₂(BO)C,H₂O₆. - K₂(BO)₂C,H₂O₆. - Na(BO)A".

- Na₂(BO)C,H₂O₆. - Na₂(BO),C,H₂O₆. (Duve, J. 1869, 540). - K₂(BO)₂C,H₂O₆. BA''.

- Ra₂(BO)C,H₂O₆. - Ra₂(BO)₂C,H₂O₆. BA''.

- Ra₂(BO)C,H₂O₆. - Ra₂(BO)A''.

- Ra₂(BO)C,H₂O₆. BA''. Ba(BO)C,H,O,.—Ba(BO),C,H,O,.—PbA". S.G. 172 4.012 (Clarke, Am. 2, 174). White crystalline pp., v. sol. HNO₃.—Pb₂C₂H₂O₃ aq (Erdmann, A. 21, 14). Insol. water, v. sol. KOHAq. — Pb₃(C₂H₂O₃)₂. — Ce₂A''₃ 9aq (?). — Th₂K₂A''₃. — CuA'' 3aq. S. 06 in the cold; ·8 at 100°. Light-

green powder. - CuA"4NH, (Schiff, A. 123, 46). —(H(NH,)A"),(HgO),3H,O (Harff, Brande Arch. 5, 259; Burckhardt, ibid. [2] 11, 257). Brandes' Efforescent, apple-green crystalline powder (Fabian, A. 103, 248).—CoCl(HA''), 5NH, 2½ aq (Jörgensen, J. pr. [2] 18, 289).—ZnA'' 2aq (Schiff, A. 125, 146).—Zn₂C, H₁O₆ aq. Powder, insol. water.—Cr(OH)A''. Violet flakes, turning dark blue over H SO (Schiff, A. 125, 146). dark-blue over H₂SO₄ (Schiff, A. 125, 145). At 220° it yields Cro, H₂O₂.—K(CrO)A" 3 aq (Malaguti, C. R. 16, 457).—(UrO₂)A" 3aq (Péligot, A. 56, 231).—K₂(UrO₂)A"₂ (dried at 200°).—(SbO)₂(UrO₂)A"₂ 8aq.—SnA". Minute crystals.—K(FeO)A"4 or 5aq. Prepared by digesting cream of tartar with water and Fe(OH)_s. Brown scales, red by transmitted light. Acids ppt. from its solutions a basic salt, sol. excess. — NH₄(FeO)A"4 (or 5) aq. — Fe₂A"₂. — FeA". Formed by boiling tartaric acid with water and iron wire (Méhu, N. J. P. 40, 257). Minute crystalline powder. S. 877 at 15.6° (Dulk, A. 2, 62).—K(MnO)A"4aq (Descamps, Z. 1870, 317).-Ag,A". Amorphous curdy pp. got by adding AgNO, to a cold solution of Rochelle salt. Crystallises from a warm solution in white scales. Nearly insol. water. Its ammoniacal solution deposits Ag on warming (Liebig a. Redtenbacher, A. 38, 132; Erdmann, J. pr. 25, 504).
—AgHA" (Perkin, C. J. 51, 869). Monoclinic tables.

Ethylene-diamine salt

C₄H₄O₄(N₂H₆C₂H₄). Dextrorotatory. Leaflets, v. sol. water. The acid salt is sl. sol. water, S. 8 at 15°, and the solution is dextrorotatory (Colson, Bl. [3] 7, 808).

Mono-methylether MeHA". Formed by boiling tartaric acid with MeOH (Guérin, A. 22,

248). Prisms.—KMeA".— BaMe₂A"₂aq (Dumas a. Péligot, A. Ch. [2] 61, 200; [3] 5, 373).

Di-methyl ether Me₂A". [48°]. (280° i.V.). S.G. (liquid) ¹⁵ 1°340. [a]_D = 2°454 at 18° (Anschütz a. Pictet, B. 13, 1176; 18, 1399); = 2°14 (Phillips a. Guye, C. R. 110, 714). Yields are certified derivative (1024 f. 1). an acetyl derivative [103'] $[a]_{D} = -14^{\circ}$ and a dibenzoyl derivative $[a]_{D} = -88^{\circ}$.

ether EtHA". Mono-ethyl Formed by dissolving tartaric acid in boiling alcohol (G.; Marian, N. J. T. 13, 2, 43; Trommsdorff, N. J. T. 24, 1, 11; Guérin, A. Ch. [2] oorn, N. J. T. 24, 1, 11; Guérin, A. Ch. [2] 62, 57). Very deliquescent prisms.—NaEtA'. Powder, sl. sol. alcohol (Mulder, B. 8, 370).—KEtA'. S. 106° at 23·5°. Colourless trimetric prisms; a:b:c='288:1: '417.—Ca(EtA'')_2 5aq.—Ba(EtA'')_2 2aq. S. 38 at 23°; 128 at 100°.—Pb(EtA'')_2—Cu(EtA'')_2 6aq. Blue silky efflorescent needles.—AgHA''. Prisms.

Diethylether EtA''. (280°; V.) (232°

Di-ethyl ether $\text{Et}_2\text{A}''$. (280° i.V.); (233° cor. at 197 mm.). S.G. $\frac{1}{18}$ 1·2097; $\frac{2}{28}$ 1·2019. $[a]_D=9$ at 18° (Anschütz a. Pictet, B. 13, 1176); 7·22 at 12°; 8·02 at 20° (Perkin, C. J. 51, 863). M.M. 8·766 at 14·8°. Liquid miscible with water (Demondesir, A. 80, 301). Sodium CO, Et.CH(ONa).CH(OH).CO, Et forms CO.Et.CH(ONa).CH(ONa).CO.Et, the latter being converted by EtCl into a deposit and a latter liquid, part of which is soluble in alcohol and ether and part insoluble in alcohol and ether (Mulder, R. T. C. 8, 375; 9, 238; 10, 171; cf. Cohn, B. 20, 2003). The soluble product mixed with alcoholic CuOl, and poured into water forms a pp. crystallising in light-green needles, sol. alcohol and ether, insol. water and $CuC_{10}H_{24}O_{11}$. Tartaric ether is converted by treatment with gaseous HCyO into allophanyl-tartaric ether NH₂,CO.NH.CO.O(C₄H₃Et₂O₅) [188°], which is m. sol. hot water and alcohol.

Acetyl derivative of the ether CO₂Et.CH(OAc).CH(OH).CO₂Et. Formed from the ether and AcCl, the mixture becoming hot (Perkin, C. J. 20, 145). Heavy oil, sl. sol. water, insol. saline solutions. Neutral to litmus.

Di-acetyl derivative of the ether CO₂Et.CH(OAc).CH(OAc).CO₂Et. [67°]. (292°) (Pictet, B. 14, 2790); (280° at 100 mm.). Formed by heating tartaric other with AcCl at 100° (Perkin; Wislicenus, A. 129, 187). Prisms (from water), v. sl. sol. water. Slowly saponified by KOHAq.

derivative of the Acetyl-benzoyl ether CO, Et. CH(OAc). CH(OBz). CO, Et. Formed by heating the mono-benzoyl derivative of the

ether with AcCl at 150° (Perkin). Heavy oil.

Mono-benzoyl derivative of the Mono-benzoyl derivative of the ether CO₂Et.CH(OH).CH(OBz).CO₂Et. [64°]. Formed by heating the ether with BzCl at 100°. Prisms, sl. sol. hot Aq. Saponified by alcoholic potash yielding C₂H₂(OH)(OBz)(CO₂Et).CO₂K, C₂H₂(OH)(OBz)(CO₂K)₂, and finally, potassium tartrate. C₂H₂(OH)(OBz)(CO₂Et).CO₂H forms tufts of needles, sl. sol. water.

Di-benzoyl derivative of the ether

[a]_D = -60° (Phillips a. Guye, C. R. 110, 716).

Di-n-propyl ether Pr₂A". (303° i.V.).
S.G. 11 1:1392. [a]_D = 13:77 at 18° (A. a. P.); = 12.44 (P. a. G.).

Di-isopropyl ether Pr_2A'' . (275°). S.G. 20 1·130. $[a]_D = 14\cdot89$ at 20° (Pictet, J. 1882, 856). Liquid.

 $[a]_{D} = 19.9^{\circ}$ (P) Di-isobutyl ether (Pr.CH2)2A". (324°). S.G. 100 1.015.

Yields a dibenzoyl derivative $[\alpha]_D = -42^\circ$. Isoamyl ether $(C_5H_{11})HA''$. Formed by digesting tartaric acid (150 pts.) with isoamyl alcohol (88 pts.) at 130° (Balard, A. Ch. [3] 12, 309; Breunlin, A. 91, 314). Nodular mass, with very bitter taste.— $KC_bH_{11}A''$ aq. Crystalline.— $NaC_aH_{11}A''$. Nodules. — $Ba(C_bH_{11}A'')_2$ 2aq: pearly plates (from water).— $Pb(C_3H_{11}A'')_2$.— $Ca(C_5H_{11}A'')_2$.— $AgC_5H_{11}A''$. Tufts of needles.

Di-acetyl derivative C₈H₁₀O₈ i.e. CO₂H.CH(OAc).CH(OAc).CO₂H. $[a]_D = -23.14$ (Phillips a. Guye, C. R. 110, 716). Formed slowly from the anhydride by the action of water. Hard deliquescent crystals (containing 3aq), melting at 58°. V. sol. water, alcohol, and ether, sl. sol. benzene. Lævorotatory. Its salts form lævorotatory solutions (Colson, Bl. [8] 7, 806; C. R. 114, 417). Converted into tartaric acid by hot KOHAq.—KHA": crystalline powder.— BaA": slender deliquescent needles.—CaA".—CuA".—Ag₂A": white crystalline mass.— Ethylene-diamine salt CoH,Os(N2HC2H4).

Les vorotatory. V. sol. water.

Anhydride of the di-acetyl derivative C.H.O., [127]. Formed by heating tartaric acid with AcCl (Ballik, Sits. W. 29, 26; Pilz, Sitz. W. 44 [2] 7; Perkin, C. J. 20, 149). Slender needles (from benzene). Dextrorota-

Benzoyl derivative C, H, O, i.e. H.CH(OH).CH(OBz).CO,H. Formed CO₂H.CH(OH).CH(OBz).CO₂H. heating tartaric acid with benzoic acid at 1500 (Dessaignes, J. Ph. [3] 32, 47). Minute crystals, v. sol. water, sl. sol. alcohol.—Ag2A": white pp.

Di-benzoyl derivative C₁₈H₁₄O₈ i.e. H.CH(OBz).CH(OBz).CO₂H. [90°].

CO₂H.CH(OBz).CH(OBz).CO₂H. [90°]. [α]_D = -116° in alcohol. Formed by dissolving its anhydride in hot water. Needles (containing aq). Converted by heat into a crystalline sub-

stance melting at 132°. Anhydride $C_{18}H_{12}O_7$. [174°]. [α]_D=143° in accone at 18°. Formed by heating tartaric acid with BzCl. Small needles, insol. cold water, sol. alcohol, benzene, and alkalis. Dextro-

rotatory.

Di-nitroxyl- derivative

CO₂H.CH(NO₃).CH(NO₃).CO₂H. Nitrotartaric acid. Formed by dissolving tartaric acid (1 pt.) in HNO₂ (4½ pts.), and stirring with an equal volume of H₂SO₄ (Dessaignes, A. 82, 362; 89, 339). The crystals are dried and crystallised from ether (Kekulé, A. 221, 245). Silky prisms, decomposed by hot water, forming oxalic and tartronic acids, CO₂, and NO. Alcohol containing HNO₂ forms di-oxy-tartaric acid. Ammonium sulphide reduces it to tartaric acid.-(NH₄)HA".—Ag₂A" aq. V. sol. water.—Et₂A". [46°]. S.G. $\frac{16}{2}$ 1.278. Formed by dissolving tartaric ether in a mixture of HNO, and H.SO. and ppg. with water.

Amide C2H4O2(CO.NH2)2. Formed by passing NH, into an alcoholic solution of the ether, or of its di-acetyl derivative (Demondesir, A. 80, 303; Pasteur, C. R. 35, 176; Grote, A. 130, 202; Ruhemann, B. 20, 3366). Trimetric crystals (from water), v. e. sol. water. Dextrorota-

tory.

Amic acid C,H,NO, i.e.

CO₂H.CH(OH).CH(OH).CO.NH₂. Formed by passing NH3 over tartaric anhydride moistened with alcohol (Laurent, Compt. Chim. 1845, 153). Formed also, together with tartaric acid, by heating tartaric ether with NH, Aq at 100° (Grote). Syrup. - CaA'₂ 6aq: tetrahedra, v. sol. water. -- Pb₂(C₄H₄NO₅)₂. -- BaA'₂ 8aq. -- EtA'. Anilide C₂H₄O₂(CO.NHPh)₂. [250°].

Formed by boiling aniline with tartaric acid (Polikier, B. 24, 2959). Pearly scales, insol. water, sl. sol. other solvents. Yields mono-, di-, and tri-acetyl derivatives, melting at 227°, 216°,

and 137° respectively.

Phenyl-amic acid C10H11NO, i.e. CO,H.CH(OH).CH(OH).CO.NHPh. Formed by boiling the phenylimide with NH,Aq (Arppe, A. 93, 352). Plates, v. sol. water and alcohol.—BaA'2: spangles, m. sol. hot water.-AgA': white powder, v. sl. sol. water.

Phenylimide C10H2NO, i.e.

CH(OH).CO NPh. [280°]. Formed, together with the anilide, by heating tartaric acid with aniline at 150°. Pearly plates, v. sol. water.

Di-o-toluide C.H.O.(CO.NH.C.H.Me)...
3°]. White plates, insol. water (Bischoff, B. 23, 2049). Ac₂O at 140° yields a di-acetyl

compound [222°]

C2H4O2(CO.NH.C4H4Me)2. Di-p-toluide [264°]. Got by heating p-toluidine with tartario acid at 185°. Silky needles (from alcohol). Converted by Ac₂O into a di-acetyl derivative [202°].

Hydraside C₂H₂(OH)₂(CO.NH.NH₂), [198°]. Formed from tartaric ether and hydrazine hydrate (Von Rothenburg, B. 26, 2057). Needles, sl. sol. alcohol. Converted by benzoic aldehyde into C.H. (OH)2(CO.N2H:CHPh)2 [225°], crystallising in yellowish plates (from alcohol). Lævo- tartaric acid

CO₂H.CH(OH).CH(OH).CO₂H. Anti-tartaric acid. [170°]. When a solution of racemic acid is divided into two equal parts and one is neutralised by ammonia and the other by soda, and the solutions mixed and allowed to evaporate, hemihedral crystals of Na(NH4)C4H4O4 are obtained. Half of these crystals have the hemihedral faces oppositely situated to those of the other half; and on carefully selecting the two kinds of crystals (the solutions of which exhibit equal and opposite action on polarised light), ppg. their solutions by lead nitrate, and decomposing the lead salts by H2S or H2SO4 solutions are obtained which on evaporation yield crystals, the one of dextro- and the other of lævo-tartaric acid (Pasteur, A. Ch. [3] 28, 56). If in a solution of sodium ammonium racemate

of S.G. 1.26 a crystal of the dextro- salt be put on

one side and one of the lævo- salt be put on the

other side of the vessel, the two salts will crystallise separately (Jungfleisch, J. Ph. [5] 5, 346). The separation of sodium ammonium

racemate into the dextro- and lævo- tartrates

does not take place if the solution is heated to 118° in a sealed tube and allowed to crystallise without access of air (Bichat, Bl. [2] 46, 54). According to Wyrouboff (C. R. 102, 627), the splitting up of the sodium ammonium racemate depends upon strength of solution and tempera-

Lævo-tartaric acid is exactly like dextro tartaric acid in solubility and S.G. Its crystals are like the reflection of those of the dextroacid. The two acids are oppositely pyro-electric. Solutions of the lævo- acid exhibit an effect on light equal and opposite to that produced by solutions of the dextro- acid of like strength. When conc. solutions of equal quantities of the dextro- and lævo- acids are mixed, heat is given out and racemic acid separates. The salts of the levo-tartaric acid have the same amount of water of crystallisation, and the same crystalline form as the salts of dextro-tartaric acid. They exhibit opposite hemihedry, opposite rotation, and are oppositely pyro-electric. Differences are observed between the compounds of the two acids with optically active substances. Acid ammonium (active) malate combines with acid ammonium dextro- (but not lævo-) tartrate. Cinchonine lævo- tartrate C18H22N2OC4H6O6 con-

tains aq, while the dextro-tartrate contains 4aq.

Di-methyl-ether Me,A". [48°]. (160°
at 16 mm.). Lævorotatory (Anschütz, B. 18, 1398; A. 247, 112).

Di-acetyl derivative of the dimethyl ether CO₂Me.CH(OAc).CH(OAc).CO₂Me. [103°].

 $[\alpha]_D = -1.3$ (in a 3.6 p.c. solution in alcohol of S.G. 826). Formed by heating Me₂A" with AcCl. Monoclinic plates (from benzene); a:b:c=1.052:1:1.007; $\beta=87^{\circ}$ 54'. On mixing with the corresponding dextro-compound it forms the isomeric racemic compound [86°].

Amide C,H,O,(NH,)2. Combines with the amide of active malic acid, forming needles, S.

more than 83.

Racemic acid $C_1H_0O_6$. Paratartaric acid. [206°] (Bischoff, B. 22, 1812). S.G. (of H_1A'') $\frac{1}{4}$ 1.778; (of H_2A'' aq) 1.687 (Perkin, C. J. 51, 866). S. (of H_2A'') 8.16 at 0°; 18 at 20°; 187.8 at 100°. S. (of H_2A'') aq) 9.23 at 0°; 20.6 at 20°; 184.9 at 100° (Liedie, C. R. 95, 87). Heat of solution at 9°: -5675 (for H_2A''); -7065 (for H_2A'' aq) (Pickering, C. J. 51, 367). H.C. (for H_2A'') 1,851,190 (Ossipoff, C. R. 108, 1105). H.F. (for H_2A'' aq) 278,400. Acid potassium racemate either occurs sometimes in crude argol, or is produced therefrom in the purification processes (Kestner; Jungfleisch, C. R. 85, 805). On crystallisation of cream of tartar, acid potassium racemate remains in the mother-liquor.

Constitution.—Assuming racemic acid to be a compound of the dextro- and lævo-taríaric acids, its formula must be at least $(C_4H_4O_8)_2$.

Formation. - 1. By mixing solutions of dextro- and lavo- tartaric acids .- 2. By heating tartaric acid (30 g.) with water (4 c.c.) for 30 hours at 175° (Jungfleisch, Bl. [2] 18, 203).— 3. By heating the dextro- or levo- tartrate of quinine or cinchonine at 170° (Pasteur, A. 72, 164; 84, 157; 88, 211).—4. By heating tartaric ether .- 5. By heating inactive tartaric acid at 200° or by boiling it with HClAq (Dessaignes, Bl. 1865, i. 34).—6. By oxidation of fumaric acid by aqueous KMnO, (Kekulé a. Anschütz, B. 18, 2150; A. 226, 191; cf. Tanatar, B. 12, 2293). 7. By the action of HNO, on lævulose, mucic acid, and inulin; and, together with tartaric acid, by the action of HNO, on milk sugar, cane sugar, saccharic acid, dulcite, and mannite (Carlet, Rep. Chim. pure, 2, 345; 4, 17; Hornemann, J. pr. 89, 283).-8. Together with inactive tartaric acid by boiling silver di-bromo-succinate with water (Perkin a. Duppa, C. J. 13, 102; Pasteur, A. Suppl. 2, 242; Jungfleisch, Bl. [2] 19, 198) .- 9. By boiling silver iso-di-bromo-succinate with water (Demuth a. Meyer, B. 21, 268). 10. By boiling glyoxal with dilute HCy and HCl (Schöyen, A. 132, 168; Strecker, Z. [2] 4, 216).-11. By the action of zinc-dust (2 mols.) and dilute HOAc (3 mols.) on glyoxylic acid (1 mol.) at 100° (Genviesse, Bl. [3] 7, 225; C. R. 114, 555).—12. From piperic acid, sorbic acid, and CHO.CH.CH.CH.CH.CO.2H by careful oxidation with dilute KMnO, below 4° (Doebner, B. 23, 2374).—13. By heating a solution of desoxalic acid (Löwig, J. 1861, 605; Klein, J. pr. [2] 20, 157).

Preparation.—Sodium ammonium racemate, got from the mother-liquor from which cream of tartar has separated, is treated exactly in the same way as cream of tartar in the preparation of tartaric acid. The small opaque needles of racemic acid can be separated by handpicking from the large transparent crystals of tartaric acid.

Properties. — Triclinic crystals (containing aq); a:b:a=484:1:806; $a=120^\circ$; $\beta=96^\circ$ 13°; $\gamma=76^\circ$ 5' (De la Provostaye, A. Ch. [3] 3, 131). Slightly efflorescent. Gives off aq at 100° . Less soluble than tartaric acid. In dilute solutions of racemic acid the lowering of the freezing-point is the same as with an equal weight of tartaric acid (Racult, Z. Ph. 2, 186). Absorption of a solution of racemic acid by filter paper does not separate the dextro- and lævo- acids (March-

A solution of racemic lewski, B. 26, 983). acid is ppd. by CaSO, (unlike tartaric acid). Calcium racemate is insol. acetic acid (unlike calcium tartrate). NH,Cl does not hinder the ppn. of calcium racemate. When heated above 200° racemic acid yields the same products as tartaric acid. By various ferments (e.g. Penicillium glaucum) racemic acid, when free or as acid ammonium salt, is converted into tartaris acid (Pasteur; cf. E. Mulder, R. T. C. 1, 231). A solution in HNO, (S.G. 1.52) poured into half its volume of H.SO, deposits the crystalline $C_2H_2(O.NO_2)_2(CO_2H)_2$ (Dessaignes, J. Ph. [3] 32, 46). AcCl converts racemic acid into di-acetylracemic anhydride C₂H₂(OAc)₂C₂O₃ [123°]. Racemic acid and its salts are inactive to light.

Salts (Fresenius, A. 41, 1; 53, 230).—
(NH₄)₂A". Trimetric prisms; a:b:c=:509:1:*847.
V. sol. water, nearly insol. alcohol. Give off NH₄ in air.—(NH₄)HA". S. 1 at 20°; less sol. boiling water. Monoclinic prisms. Reddens litmus.—K₄A" 2aq. Six-sided tables. S. 103 at 25°. Nearly insol. alcohol.—KHA". S. :55 at 19°; '7 at 100°. Insol. alcohol.—K(NH₄)A". Striated prisms.—Na₂A". S. 38 at 25°. Trimetric prisms. Insol. alcohol.—NaHA"aq. S. 9 at 19°. Monoclinic prisms (from water). Ppd. from aqueous solution by addition of alcohol.—Na(NH₄)A"aq. S.G. 1:740 (Wyruboff, A. Ch. [6] 9, 229).—NaKA" 4aq. S. 47 at 6° (Fresenius, A. 53, 230).—NaKA"3aq. Monoclinic crystals (Wyruboff, A. Ch. [6] 9, 232).—RbA".—CaA"4aq. Acicular prisms (Anschütz, A. 226, 191).—BaA"2½aq. S. :05. Crystalline pp.—SrA"4aq.—TlA" (De la Provostaye, A. 126, 79).—LiTlA"2aq (Wyruboff).—NaTlA"2aq. S. 25 at 25°.—MgA"5aq. S. :85 at 19°.—CdA" (Schiff, A. 104, 326). PbA" (dried at 100°).—Pb₂C₄H₂O₆ (Krug, J. 1861, 367).—CuA"2aq. Pale-blue needles, sol. alkalis and solution of Na₂CO₃.—Na₂CuC₄H₂O₆ 4aq. —MnA" aq. — NiA"5aq. —NH₄(AsO)A"½aq. S. 94 at 15°. Efflorescent crystals.—K(AsO)A"½aq. S. 94 at 15°. Efflorescent crystals.—K(AsO)A"½aq. S. 7 at 19°. Non-efflorescent white crystals (Werther, J. pr. 32, 385).—K(SbO)A"½aq. Monoclinic prisms. — Ag₂A". Shining scales, less sol. Aq than silver tartrate.

Mono-methyl ether MeHA". Formed by evaporating racemic acid with MeOH (Guérin-Varry, A. Ch. [2] 62, 77; A. 22, 252). Rectangular prisms, v. e. sol. cold water. Decomposed by hot water into the parent substances.—KMeA" aq. Prisms.—Ba(MeA") 4aq. Monoclinic prisms.—AgMeA". Pp.

clinic prisms.—AgMeA". Pp.

Di-methyl ether Me₂A". [85°]. (282°
i.V.). Formed by methylation of racemic acid, and also by mixing equal weights of the dimethyl ethers of the lavo- and dextro-tartaric acids (Anschutz a. Pictet, B. 13, 1178; 18, 1897; A. 247, 116). Monoclinic crystals; optically inactive. Yields a di-acetyl derivative C₂H₂(OAc)₂(CO₂Me)₂ [86°] in trimetric crystals; a:b:c='8095:1:'6728. This compound, examined by Racult's method, has the same molecular weight as the di-acetyl derivative of di-methyl tartrate.

Mono-ethyl ether EtHA". Deliquescent prisms, v. sol. alcohol. Inactive to light.—KEtA" aq.—Ba(EtA"), 2aq.—AgEtA". Prisms.

Di-ethyl ether Et.A". (238° cor. at 197 mm.) S.G. § 12098; § 12021. M.M. 8759 at 1556 (Perkin, C. J. 51, 364). Yields an oily acetyl derivative and a di-acetyl derivative C₂H₂(OAc)₂(CO₂Et)₂, [51°] (230° at 100 mm.). BzCl forms C₂H₂(OH)(OBz)(CO₂Et)₂ [57°]. The compound C₂H₂(OEt)₂(CO₂Et)₂ (147° at 15 mm.) is formed from di-bromo-succinic ether [58°] and NaOEt (Michael, J. pr. [2] 46, 234). – Ag_2A'' . AgHA". Monoclinic, a:b:c = .721:1: .869; $\beta = 70^{\circ}$

Inactive tartaric acid

CO₂H.CH(OH).CH(OH).CO₂H. Mesotartaric

acid. [140°-143°]. S. 125 at 15°.

Formation. - 1. By heating cinchonine tartrate or racemate for several hours at 170° (Pasteur, A. 88, 212). The product is mixed with CaCl₂, which ppts. calcium racemate, and the filtrate slowly deposits calcium inactive tartrate.—2. By boiling tartaric or racemic acid for 400 hours with HClAq (Dessaignes; Pasteur, Bl. 1862, 107).—3. By heating tartaric acid (30 g.) or racemic acid with water (4 c.c.) for 2 days at 165°. Separated by means of its very soluble acid K salt (Jungfleisch, Bl. [2] 19, 101).—4. A product of the action of HNO₃ on sorbin (Dessaignes, C. R. 55, 769), erythrite (Przybytek, B. 17, 1412; Bl. [2] 43, 126) and glycerin.—5. By oxidation of lævulose (Kiliani, B. 14, 2530).-6. By oxidation of phenol by alkaline KMnO, at 0° (Doebner, B. 24, 1755).—7. A product of the action of moist Ag₂O on di-bromo-succinic acid. 8. By the action of nitrous acid on di-amidosuccinic acid (Lehrfeld, B. 14, 1819).—9. By oxidation of maleic acid by alkaline KMnO₄ (Kekule a. Anschütz, B. 14, 713).

Properties.—Prisms (containing aq). Loses its aq at 100°; at 200° it yields pyrotartaric and racemic acids. Prolonged boiling with HCl aq converts it into racemic acid (Dessaignes). The conversion is also effected by heating the acid with a little water at 175°. ether is converted by AcCl into C₂H₂(OAc)₂(CO₂Et)₂ [48°], crystallising in silky

needles.

Salts KHA". Needles. S. 12·5 at 19°.— RbHA" jaq. — Rb₂A"aq. — RbNaA" 2jaq. — CaA" 3aq. S. 17 at 100°. S. (HOAc) 028 at 18°; '085 at 100°.—PbA" aq: amorphous pp. — ZnA" 2aq (dried at 110°). — TlHA". — Tl₂A".—NaTlA" 2 aq. — Ag₂A" aq: white pp.

Metatartaric acid (C4H6O6)x. A deliquescent, glassy mass, v. sol. water, obtained by melting tartaric acid (Braconnot, A. Ch. [2] 68, 299; Erdmann, A. 21, 9; Laurent a. Gerhardt, A. 70, 848). It is sometimes formed when a solution of tartaric acid is evaporated over a water-bath (Grosjean, C. J. 43, 334). Converted by boiling water into tartaric acid.

Salts .- (NH4)HA". Minute needles. More soluble than the acid ammonium tartrate. Its solution is ppd. by CaCl, and NH2Aq only when highly concentrated. CaA" 4aq. Minute lenticular grains, sol. dil. HClAq, converted by

boiling water into calcium tartrate.

Ditartaric acid CaH10O11. An anhydride of tartaric acid, got by heating tartaric acid at 170° till it is no longer ppd. by lime-water (Fremy, A. 19, 197; 29, 142; 78, 297). Amorphous and very deliquescent. Sol. alcohol, insol. ether. Its salts are amorphous, v. sol. water, and ppd. by alcohol. They are converted into metatartrates, and finally into tartrates, by boiling water.—CaA". - BaA".—PbA".—CuA".— Ag₂A". V. sol. water (Schiff, A. 125, 129).

Ethyl ether Et.A". Fat-like solid.

Tartrelic acid C.H.O. Soluble tartaric anhydride. Formed by quickly heating small quantities of tartaric acid till the fused mass swellsup. Yellowish deliquescent mass, sol. water, forming an acid solution, from which CaCl, ppts. CaA'₂ as an amorphous mass, which becomes crystalline on adding alcohol. Tartrelic acid is converted into tartaric acid by boiling water.

Insoluble tartaric anhydride Formed by heating tartaric acid for some time at 150° and exhausting the product with cold water. Powder, insol. water, alcohol, and ether. Converted by hot water into tartaric acid.

TARTRONIC ACID C3H4O5 i.e.

CH(OH)(CO,H). Oxy-malonic acid. Mol. w. 120. [187°]. Heat of solution -4331 (Gal a.

Werner, Bl. [2] 46, 803).

Formation.—1. By spontaneous decomposition of nitro-tartaric acid in aqueous solution (Dessaignes, C. R. 34, 731; 38, 44).—2. By the reduction of mesoxalic acid by sodium-amalgam (Baeyer, J. 1864, 641).-3. By the action of Ag₂O on bromo-malonic acid (Petrieff, B. 11, 414).—4. By boiling chloro-malonic ether with KOHAq, the yield being 25 p.c. of the theoretical amount (Conrad a. Bischoff, A. 209, 222) .-5. From di-bromo-pyruvic acid and barytawater at 40° (Grimaux, B. 10, 903).-6. From glyoxylic acid by treatment with KCy, the resulting nitrile being saponified (Böttinger, B. 14, 729).-7. A product of the action of alkaline cupric solution on glucose (Claus, Z. [2] 5, 152; A. 147, 114).—8. By slow oxidation of glycerin (Sadtler, B. 8, 1456; Campani a. Bizzari, G. 12, 1).-9. By heating sodium carboxy-tartronate (Gruber, B. 12, 514).

Preparation.—1. By boiling the di-nitrate of tartaric acid (20 g.) with alcohol (60 g. of S.G. 925) (Demole, B. 10, 1778).—2. By warming tri-chloro-lactic ether CCl3.CH(OH).CO2Et with a 10 p.c. solution of NaOHAq (5 mols.), neutralising with HOAc and ppg. with BaCl, (Pinner

B. 18, 752, 2852).

Properties.—Prisms (containing aq), v. sol. water and alcohol, sl. sol. ether. After drying at 100° it loses its water of crystallisation and becomes v. sol. ether. Massol (C. R. 114, 422) obtained anhydrous crystals [155°]. Decomposed by fusion, yielding CO2 and glycolide C2H2O2.

Reduced by HIAq to malonic acid. Salts.—Na₂A" (dried at 100°). water. — KA" aq. — (NH₄)_A". Needles. —
*(NH₄)HA". Yields glycollic amide on heating.
—BaA"2aq. Micro-crystalline powder, v. sl. sol. water (Pinner).—BaA" aq (G.).—CaA" 2½ aq (P.).—CaA" aQ (G. a. B.).—CdA".—PbA" (dried at 120°).—MnA".—Ag₂A". Needles (from hot Aq).

Ethyl ether Et₂A". (c. 224°). Heavy oil (Freund, B. 17, 786; Pinner).

The acetyl derivative CH(OAc)(CO₂Et).

(c. 161° at 60 mm.), S.G. $\frac{19.5}{16}$ 1.131, is formed from bromo-malonic ether and KOAc in EtOH at 50° (Conrad a. Brückner, B. 24, 2997). Br converts it into mesoxalic ether C(OH)2(CO,Et). [57°].

Amids CH(OH)(CO.NH₂)_x [196°]. Six-

sided plates, m. sol. hot water.

Amic acid CO.H.CH(OH).CO.NH. [160°]. Formed by boiling sodium dialurate with water (Menschutkin, A. 182, 82). Needles, sl. sol. cold water. Converted by nitrous acid into glycollic acid.—KA'aq.—BaA', aq.—PbA', ½aq. Needles, v. sol. hot water.—AgA'. Needles.

Phenyl derivative CH(OPh)(CO2H)2. Formed by saponification of its ether CH(OPh)(CO₂Et)₂ (230°-240° at 60 mm.), which is got from bromo-malonic ether, phenol, and NaOEt (C. a. B.). At 196° it forms CH₂(OPh).CO.H.
TARTRONYL-UREA v. DIALURIO ACID.

TAURINE C2H, NSO, i.e.

CH₂(NH₂).CH₂.SO₃H. Amido-ethane sulphonic acid. Mol. w. 125. S. 6.5 at 12°. H.F. 185,700 (Berthelot a. Matignon, A. Ch. [6] 22, 182). Obtained by the action of acids and alkalis on taurocholic acid (Gmelin, Tiedemann a. Gmelin's Die Verdauung, 1, 43; Demarcay, A. 27, 286; Dumas a. Pelouze, A. 27, 292; Redtenbacher, A. 57, 170; Strecker, A. 65, 132; 67, 34; 91, 101). Occurs in the free state in putrid bile and in the muscles of the oyster, in shark's blood, in the liver, spleen, and kidneys of the ray, and in the lungs of oxen (Valenciennes a. Fremy, C. R. 41, 735; Städeler a. Frerichs; Clöetta, A. 99, 289). Formed also by heating β-chloro-ethane sulphonic acid with NH3Aq at 100° (Kolbe, A. 122, 33), and by the action of 80, on vinylamine (Gabriel, B. 21, 2667). Prepared by evaporating ox-bile with dilute HCl at 100°, decanting from a resin, allowing to crystallise, and re-crystallising from water. It may be purified by means of the lead salt.

Properties. - Transparent crystals, m. sol. water, insol. alcohol. Neutral to litmus. Not melted below 240°. Converted by nitrous acid into isethionic acid (Gibbs, Am. S. [2] 25, 30).

Decomposed by potash-fusion.

Salts.—NaA'. Very deliquescent (Lang, Bl. [2] 25, 180).—CaA'₂. Slender needles, v. e. sol. water.—PbA'₂. Needles, v. e. sol. water. water.—PbA'₂. Needles, v. e. sol. water.— Pb₂A'₄(OH)₂—CdA'₂.—HgA'₂.—Hg₂A'₂O (Engel, Bl. [2] 23, 532).—AgA'.

Benzoyl derivative NHBz.C.H. SO.H.

Formed by heating μ -phenyl-thiazole tetrahy-dride with HCl and bromine-water (Gabriel a. Heymann, B. 23, 159). Syrup.—AgA'. Spheri-

cal groups of plates.

Phthalyl derivative (C,H,O,):N.ČH,CH,SO,H. Got by oxidising di-phthalimido-di-ethyl-disulphide with HNO. (S.G. 1.25) (Gabriel, B. 24, 1116). Crystals (containing 1½aq), sol. EtOAc. Tastes bitter.

Imide C₂H₄ NH SO₂. [45°-50°]. Prepared 8 - chloro - ethane - sulphonic chloride CH2Cl.CH2.SO2Cl by treatment with ammonia (James, C. J. 49, 491). Solid, resembling gum arabic; bitter in taste. V. al. sol. cold, v. sol. hot, water, insol. alcohol and ether. Ppd. HgO appears to form the compound (C.H.SO.N).Hg.

Ethyl-taurine v. ETHYL-AMIDO-ETHANE SUL-

PHONIC ACID.

Methyl-taurine v. Methyl-amido-ethane SULPHONIC ACID.

TAUROCHENOCHOLIC ACID O.H. NSO. Occurs as K and Na salts in goose-bile (Heints a. Wislicenus, P. 108, 547; Otto, A. 149, 192). Amorphous, sol. water and alcohol. Decomposed by boiling with baryta or potash into taurine and chenocholic acid.—NaA'aq (dried at 110°). Small tables. Its aqueous solution gives pps. with CaCl2, AgNO2, and lead subacetate, but not with lead acetate.

TAUROCHOLIC ACID C28H48NSO7. Occurs as Na salt in the bile of most animals (Strecker, A. 67, 30; 70, 169; Hüfner, J. pr. [2] 19, 302; Copeman a. Winston, J. Physiol. 10, 213). Obtained by ppg. with lead acetate, filtering from lead glycocholate, and ppg. lead taurocholate by lead subacetate. Deliquescent silky needles, v. sol. water and alcohol, m. sol. ether. Dextrorotatory, [a]p = 25° (Hoppe-Seyler, Arch. Path. Anat. 15, 126). Boiling water splits it up into taurine and cholic acid. Antiseptic, and ppts. albumen completely from its solutions (Maly, M. 4, 89). Salts.—NaA' (dried at 120°). Ppd. by alco-

hol in an amorphous state, rendered crystalline by ether .- KA'. Needles, resembling wavellite.

TAUROCYAMINE C. H.N. SO. i.e.
NH:C(NH₂).NH.CH₂.CH₂.SO.₂H. [226°]. S. 4 at
21°. Formed by heating taurine with symamide (1 mol.) and water at 110° (Engel, B. 8, 1597; Dittrich, J. pr. [2] 18, 63). Small white prisms, v. sol. water, insol. alcohol and ether. From cold water it crystallises with aq.

TAXINE C₃₇H₃₂NO₁₀. [82°]. Obtained from the leaves, twigs, and fruit of the yew (*Taxus baccata*) by extracting with ether (Lucas, Ar. Ph. [2] 85, 145; Van der Harst, R. T. C. 3, 279; Marmé, Bl. [2] 26, 417; Hilger a. Brande, B. 23, 464). The ethereal solution is quickly shaken with acidulated water, and the solution ppd. by NH₂Aq. Crystalline. Gives off aromatic fumes on melting. V. sl. sol. water, v. sol. alcohol and ether, insol. benzene. Conc. H2SO4 gives a deep purple; Fröhde's reagent a reddish-violet colour. Gives yellow pps. with sodium phosphomolybdate, I in KI, and potassium bismuth iodide. Insol. alkalis. The salts crystallise with difficulty.—Salts: B'HCl.—B'2H2SO4.—B'2H2PtCl6. -B'HAuCl₄.—B'EtI.

TECTOQUINONE. [171°]. Obtained from teak resin or by the distillation of teak wood (Romanis, C. J. Proc. 4, 116). Amber-yellow crystals, sol. alcohol. Reduced by zinc-dust and HOAc to C18H22. Yields a nitro-derivative and

a bromo-derivative [165°].

TELLURATES. Salts of telluric acid, H₂TeO₄ (q.v., p. 649). The normal tellurates of monovalent metals have the composition M₂TeO₄.xH₂O; there are also acid tellurates MHTeO, xH2O; and two classes of tetratellurates, (1) M₂Te₁O₁₃.xH₂O, which are soluble in water when M is an alkali metal, and (2) M2Te4O18, which are insoluble in water. The hydrated tetratellurates may be represented as acid ditellurates derived from the hypothetical acid $H_2Te_2O_7$; thus $K_2Te_4O_{13}.4H_2O=2KHTe_2O_7.3H_2O$. Some acid salts may be represented as derived from hypothetical ditelluric acid, e.g.

PbTe₂O₇ (=PbTeO₄.TeO₃).
The normal alkali tellurates are obtained by saturating H₂TeO₄ with MOHAq or M₂CO₂Aq and crystallising. Alkali acid tellurates, and hydrated tetratellurates, are obtained by interactions of acids with the normal tellurates; anhydrous (insoluble) tetratellurates of the alkali

metals are formed by heating the acid tellurates; thus $6KHTeO_4 = K_2Te_4O_{12} + 2K_2TeO_4 + 3H_2O_4$ Most of the other tellurates are obtained from the alkali salts by double decomposition.

The alkali tellurates, other than those of the form M2Te4O1, are soluble in water; most of the other tellurates are insoluble or sparingly soluble in water. Tellurates generally dissolve easily in HClAq; the solutions are not ppd. by water, but on boiling the solutions become yellow, give off Cl, and then yield a pp. of H2TeO2 on addition of water. Acidulated solutions of tellurates are slowly decomposed by H2S, probably with ppn. of TeS₃ (v. Tellurium sulphides, p. 655). Solutions of tellurates are slowly reduced by SO2, more quickly on heating, with separation of Te. When tellurates are heated to redness they give off O and form tellurites; heated with K₂CO₂ and charcoal, they give K₂Te.

The tellurates have been examined chiefly by Berzelius (P. 32, 577); and some of them

also by Oppenheim (J. pr. 71, 267).

Potassium tellurates. The normal salt K₂TeO₄.5aq is obtained by dissolving H₂TeO₄ and K₂CO₃, in the ratio H₂TeO₄:K₂CO₃, in hot water and crystallising; also by dissolving H. TeO, in excess of KOHAq, heating, and allowing to cool slowly. Crystallises in rhombic prisms, isomorphous with K2SO4 (Handl, J. 1861. 266). The crystals become moist in air and form KHTeO4. xaq and K2CO3. The acid salt 2KHTeO₄. 3aq separates on adding to K_2 TeO₄Aq sufficient H_2 SO₄Aq, or other strong acid, to remove half the potassium; the salt is crystallised from hot water. This salt is slightly soluble in cold water, but much more soluble in hot water; when heated it is converted into K₂TeO₄ and K₂Te₄O₁₃. The tetratellurate K₂Te₄O₁₃ is obtained, as a yellow powder, by heating 2KHTeO. 3aq below redness and washing out K2TeO. in water. K2Te.O. is insoluble in water, also in HClAq or HNO. Aq except after prolonged boiling. The hydrated tetratellurate K2Te4O13.4H2O-which may also be regarded as an acid ditellurate 2KHTe₂O₇.3H₂O - is formed by adding H2SO4Aq, or another strong acid, to $K_2\text{TeO}_4\text{Aq}$ in quantity to remove 6K from $4K_2\text{TeO}_4$ ($4K_2\text{TeO}_4\text{Aq} + 3H_2\text{SO}_4\text{Aq} = K_2\text{Te}_4\text{O}_4\text{Aq} + 3H_2\text{O} + 3K_2\text{SO}_4\text{Aq}$), and crystallising. The salt is fairly soluble in cold water; it is partly decomposed to KHTeO, and H2TeO, Aq by repeatedly crystallising in aqueous solution.

The following tellurates have been described (Berzelius, l.c.; Oppenheim, l.c.): Ammonium salts (NH₄)₂TeO₄, (NH₄)₂Te₂O₇, (NH₄)₂Te₄O₁₂, obtained by ppg. solutions of the corresponding salts of K or Na with NH₄Cl. Barium salts BaTeO₄, 3aq,BaH₂(TeO₄)₂, 2aq, BaTe₄O₁₈, obtained by ppn. Cadmium salt CdTeO₄, obtained by ppn. Calcium salt CaTeO, obtained by ppn. Lead salts PbTeO₄, PbTe₂O₇, PbTe₄O₁₈, by ppn; a basic salt is also said to exist. Silver salts Ag₂TeO₄, and several basic salts xAg₂O₄/TeO₃. Sodium salts Na₂TeO₄, xaq, NaHTeO₄, xaq, NaHTeO₅, xaq, and Na₂Te₄O₁₈, obtained similarly to the potassium salts. Tellurates of Al, Cr, Co, Cu, Fe, Li, Mg, Mn, Hg, Ni, and Sr have also been described, but only very superficially.

M. M. P. M. TELLURHYDRIC ACID H.Te; v. Hydrogen TELLURIDE, vol. ii. p. 727.

TELLURIC ACID H. TeO.; crystallises from

solutions as H₂TeO₄.2H₂O.

Formation.—1. By oxidising Te by aqua regia.-2. K.TeO, is formed by the interaction of Cl with K2TeO2Aq, also by fusing TeO2 with

KNO, or KClO.

Preparation.—1. A mixture of TeO2, KClO2, and KOH, in the ratio TeO2:2KClO2:2KOH is thoroughly fused; the fused mass is dissolved in water and BaCl, Aq is added; the pp. of BaTeO. is purified by recrystallisation from hot water, and is then dried at 100°; the dry BaTeO, is mixed with its own weight of water and one-fourth its weight of conc. H2SO4, and heated till decomposition is complete, when BaSO, is removed by filtration, and the filtrate is evaporated at 100° to the crystallising point; the crystals that separate are dried, powdered, washed repeatedly with alcohol to remove H2SO4, and recrystallised from water (Oppenheim, J. pr. 71, 267). Berzelius (P.28, 392; 32, 1) fused TeO_2 with an equal weight of K2CO3 or Na2CO3, dissolved in water, added KOH equal to the weight of TeO, used, passed in Cl until the liquid smelt strongly thereof, ppd. by BaCl2Aq, and proceeded as described above. -2. TeO2 is dissolved in HNO, Aq, the solution is boiled with PbO, and filtered, Pb is ppd. from the filtrate by H₂SO₄Aq, the filtrate from PbSO, is evaporated to dryness at 100°, the residue is digested with alcohol and ether to remove H2SO4, and is then crystallised from hot water in vacuo (Becker, A. 180,

Properties and Reactions.-H2TeO42H2O is obtained by slow crystallisation in white, monoclinic crystals, resembling gypsum, S.G. 29999 at 25.5° (Clarke, Am. S. [3] 16, 401). When obtained by rapid evaporation H, TeO, 2H,O appears as a glassy solid. Heated to c. 130° H₂TeO₄ is obtained; decomposition begins at a little above 160° to TeO, and H₂O, and at c. 180° TeO, begins to decompose to TeO, and O, this decomposition being complete at c. 850° (Wills, C. J. 35, 704). H₂TeO, has S.G. 3.425 at 18.8° (Clarke, l.c.). H₂TeO, $2H_2$ O is slightly soluble cold water; it is dissolved fairly easily by boiling water; soluble in acid and alkali solutions; almost, if not quite, insoluble in alcohol. H_2 TeO₄ dissolves very slowly in boiling water. Aqueous solutions of H_2 TeO₄ have a metallic, not a sour, taste; they very slightly redden litmus. Thomsen (Th. 2, 277) gives the following H.F.: [Te,O²,Aq] = 98,380; [TeO²Aq,O] = 21,200. H_2 TeO₄ dissolves gradually in hot, fairly conc. HClAq, with evolution of Cl and formation of H2TeO,Aq. H2TeO,Aq is not reduced by passing H.S into it, but if the solu-tion is saturated with H.S and kept in a warm place, in a closed flask, for some time, the liquid becomes brown, and flakes, probably of TeS, (v. Tellurium sulphides, p. 655), are deposited.

H2TeO Aq reacts with alkalis to form salts M₂TeO, and MHTeO; salts of the forms MHTe₂O, and M₂Te₄O; are also known (v. Tel-M. M. P M. LURATES, p. 648)

TELLURIDES. Any compound of Te with a single element more positive than itself is called a telluride. Tellurides of the heavier metals are generally formed by melting the metals with Te, also by passing H_zTe into solutions of salts of the metals; tellurides of K and Na are formed by heating TeO₂ with K₂CO₃ or Na₂CO₃ and charcoal; many tellurites and tellurates give tellurides when heated in a current of H. ppts. tellurides of Cu, Ag, and some other metals, by boiling solutions of salts of these metals with Te (Parkmann, C. C. 1862. 814). Tellurides of K and Na dissolve in water with a red colour; this colour is due either to polytellurides or to the presence of traces of oxides according to Demarçay (Bl. [2] 40, 99). Berthelot a. Fabre (C. R. 105, 92) obtained colourless solutions of alkali tellurides by passing pure H2Te into solutions of alkalis; a trace of O produced a red colour and separated Te. Tellurides react with HClAq giving off H.Te; they are oxidised to tellurites by HNO, Aq, and to tellurites and tellurates by aqua regia. Warmed in Cl tellurides give metallic chlorides and TeCl,.

Tellurides of Sb, As, Bi, Au, Fe, K, Ag, and Na are mentioned under these metals (cf. Brauner, C. J. 55, 388). M. M. P. M.

TELLURITES. Salts of tellurous acid H₂TeO₃ (q. v., p. 656). The normal and acid tellurites of monovalent metals belong to the forms M₂TeO₃ and MHTeO₃. Some salts are also known containing relatively more of the acidic radicle; these may be formulated as M₂O_xTeO₂ where x is 2 and 4; or they may be regarded as M₂TeO₃,xTeO₂, where x is 1 and 3; or finally these hyperacid salts may be looked on as M₂Te₂O₃ and M₂Te₄O₉—that is, as derivatives of the hypothetical acids H₂Te₂O₃ and H₄Te₄O₉.

Tellurites are generally formed by heating together basic oxides and TeO₂ in the proper proportions; some tellurites of the heavier metals are obtained by ppn. from solutions of alkali tellurites. Tellurites of the alkali metals dissolve readily in water; those of the alkaline earth metals are less soluble; the others are insoluble in water. Most tellurites dissolve in HClAq, forming yellow solutions which do not give off Cl on heating; this reaction distinguishes them from tellurates (v. p. 649). Solutions of tellurites in a little HClAq deposit H₂TeO₃ on dilution. Most tellurites are fusible, and crystallise on cooling; the hyperacid alkali tellurites form colourless glass-like solids after fusion. By strongly heating with charcoal and K₂CO₃ most tellurites give K₂Te.

Tellurites of potassium. The normal salt K₂TeO₂, and the acid salt K₂TeO₃.TeO₂

K_TeO₂, and the acid salt K_TeO₃.TeO₂ (= K_Te₂O₃ or K_2O.2TeO₂) are prepared by fusing TeO₂ with K_2CO₃ in the proper proportions. K_TeO₄ melts at a red heat; dissolves slowly in cold water, more quickly in boiling water; the aqueous solution has an alkaline taste and reaction, and is decomposed by the CO₂ in the air. K_2O.2TeO₂ melts below redness, and solidifies again to a transparent, colourless, crystalline mass; dissolved by boiling water, and the solution on cooling deposits K_O.4TeO₂. The tetratellurite or hyperacid tellurite K_0.4TeO₂.4H_2O is prepared by boiling H₂TeO₂Aq with K₂CO₂ and filtering at 100°; the salt is deposited on cooling in microscopic six-sided prisms and tablets. This salt reacts with cold water to give a solution of K_TeO₂ and K_Te₂O₃ and a gelatinous pp. of H₂TeO₂; when heated 4H₂O is given off, and K₂O.4TeO₂ remains as a salt which melts at a

moderate red heat and solidifies on cooling to a colourless glass.

The tellurites have been examined chiefly by Berzelius (P. 32, 577; v. also Oppenheim, J. pr. 71, 267). Salts of the following metals have been isolated, but many have been studied in very superficial way: Al, NH,, Ba, Cd, Ca, Cr, Co, Cu, Fe, Pb, Li, Mg, Mn, Hg, Ni, Ag, Na, Sr, and Zn.

M. M. P. M.

TELLURIUM. Te. At. w. c. 125 (v. infra). Mol. w. c. 250 (v. infra). Melts at c. 455° (Carnelley a. Williams, C. J. 37, 125). Boils at c. 1390° (Deville a. Troost, C. R. 91, 83). S.G. 6.2549 at 0° (after compression); 6.2322 at 0° (uncompressed) (Spring, B. B. 5, 854); 6.2549 at 18.2° (after melting in H) (Přiwoznik, C. C. 1892. ii. 962; for further data v. Clarke's Table of Specific Gravities [2nd ed.] 10). V.D. at c. 1390° to c. 1440° = 130 (Deville a. Troost, C. R. 56, 891). S.H. (21° to 51°, crystalline) ·0475 (Kopp, T. 1865, 71); (at 98°, crystalline) ·0483; (at 98°, crystalline) ·0483 after distillation in SO₂) 0518 (Fabre, A. Ch [6] 14, 101). C.E. (linear, at 0°) ·00001675 (Fizeau, C. R. 68, 1125). E.C. ·000777 at 19·6 (Ag at 0° = 100) (Matthiesen, P. 103, 428); conductivity increases slightly when light falls on Te; the increase is very much less than in the case of Se (v. Adams a. Day, Pr. 24, 163; 25, 113). E.C. decreases slightly to c. 100°-140°, and then increases to 200°; on cooling after fusion E.C. decreases regularly and markedly; the following values are given by Exner (P. 158, 625) .0035 at 200°, .00286 at 20° (after fusion), .000435 at 20° (after very slow cooling from fusion). crystallises in rhombohedra; a:c=1:1.3298; it is isomorphous with S and Se (v. Muthmann, Z. P. C. 8, 396). For emission-spectrum v. Plücker a. Hittorf (Pr. 13, 153); Thalen; Salet (A. Ch. [4] 28,51); Ditte (C. R. 73, 622); Hartley a. Adeney (T. 1884 [1] 117). For absorptionspectrum of vapour of Te v. Gernez (C. R. 74, 1190). H.C. [Te,O²,Aq] = 98,380 (Th. 2, 278).

Historical — Examining a mineral containing gold from the Seven Mountains in 1782 Müller von Reichenstein suspected the presence of a new metal (Abhand. einer Privatgesellschaft in Bühmen). He sent a specimen of the mineral to Bergmann, who said that it contained a metal which was not the same as that in blende. In 1798 Klaproth received some of the mineral from M. von R., and isolated the characteristic element, to which he gave the name tellurium (from tellus = the earth) (K., Beiträge zur chem. Kenntniss der Mineralkörper, 3, 1; cf Crell's Anntniss der Mineralkörper, 3, 1; cf Crell's Antonis of Te were more fully examined by Berzelius (S. 6, 311; 34, 78; P. 8, 411; 28, 392; 32, 1,

Occurrence.—Te is found native, in small quantities, and in a few localities—chiefly in Hungary and Transylvania, and in Virginia, U.S.; it generally accompanies Au and Ag; specimens have been obtained with 97-98 p.c. Te. Tellurides are also found in small quantities in Hungary, Mexico, California, Bolivia, and some other places; in considerable quantities in Colorado; these are generally compounds of Te with Bi, Ag, Pb, Bi and S, Ag and Au, Au, Pb and Sb; tellurite (or tellurium cochre) containing TeO₂ is found in small quantities in the Seven Mountains (v. G. Rose, P.

18, 64; Rammelsberg, J. M. 1874. 30; Burkart,
J. M. 1873. 476; Silliman a. Marvine, Am. S.
[3] 8, 25; Mathewson, J. M. 1866. 93; Genth,

J. M. 1875. 188, 314).

Formation.—1. By passing SO₂ into H₂TeO₃Aq; H₂TeO₂Aq + 2SO₂ + H₂O = 2H₂SO₄Aq + Te. Also by ppg. solution of TeCl₄ in dilute HClAq, or of a Te compound in H₂SO₄Aq, by an alkali sulphite. 2. By reducing a boiling solution of $(NH_4)_2$ TeO₄Aq by grape sugar (Stolba, J. 1873. 214).—3. By decomposing K₂TeAq by air or oxygen (v. Oppenheim, J. pr. 71, 267; 81, 308; Himly, J. 1877. 213).—4. By ppn, by means of zinc, from solutions of Te compounds in H₂SO₄Aq <math>(v. Löwe, J. pr. 60, 168).

Preparation.—There are several methods of preparing crude Te from the minerals that contain it. Native bismuth telluride is made into a paste with oil and K2CO3, and gradually heated to whiteness in a covered crucible; K,Te is extracted from the fused mass by boiling water, and the solution deposits Te on standing in air. Or the mineral containing Te is powdered and treated with dilute HClAq; the insoluble matter is gradually added to three times its weight of conc. H2SO4, and heating is continued until SO2 ceases to be given off; the solid that forms on cooling is dissolved in water containing a little HCl; the solution is filtered from AgCl, Te is ppd. from the filtrate by zinc, and undissolved zinc is removed by digestion with dilute HClAq. For details and other methods v. Berzelius (P. 8,411; 28,392); Löwe (J. pr. 60, 163); Wöhler (4.84,69); Oppenheim (J.pr.81,308); Schnitzer (D. P. J. 211, 484, 492); Schwarz (D. P. J. 186,

To prepare pure Te, Brauner (C. J. 55, 382; cf. M. 10, 411) recommends to digest the crude material with HClAq in a large flask, and to oxidise by HNOsAq added little by little till the solid matter becomes white, to evaporate several times with conc. HClAq (to remove HNO₃), and then to dilute cautiously with water, taking care not to add enough water to ppt. H. TeO, along with any PbCl₂ which may come down. He then adds some saturated SO₂Aq, warms to 60°-70°, and ppts. Te by passing SO_2 into the warm solution. The Te thus obtained generally contains a little copper and lead selenides. The pp. is dried and fused in small portions with KCy; the fused mass, which contains K2Te, is dissolved in water, from which air has been removed, in a stoppered flask; the liquid is allowed to clear (there is generally a small pp. of tellurides of heavy metals mixed with a trace of Te) and is then poured off, and a current of air is passed through it, whereby Te is ppd. The Te is dried, placed in a porcelain boat, which is introduced into a wide tube of hard glass surrounded by wire gauze, and distilled in a current of pure dry H. (For a method of purification based on the ppn. of Te from $(NH_4)_2$ TeO₄Aq by grape sugar, v. Stolba, J. 1878. 214.)

Properties.—As prepared by sublimation in H, tellurium is a white, shining, crystalline, very brittle solid, unchanged by exposure to air. During distillation in H, Brauner (l.c.) noticed that some H₁Te was formed, and was decomposed with deposition of needle-shaped crystals of Te, some of which were 1 centim. in length. The crystals of Te are hexagonal rhombohedra, iso-

morphous with S and Se (v. Muthmann, Z. P. C. 8, 396). By allowing K. TeAq to decompose in air, H. Rose obtained six-sided prisms with rhombohedral summits (P. 21, 443; cf. Fabre a. Berthelot, C. R. 104, 1405). Amorphous Te is obtained as a brown powder by ppg. H2TeO3Aq by SO2; on warming, the amorphous variety changes to the crystalline, giving off a large quantity of heat (F. a. B., l.c.). Te that has been melted and cooled quickly seems to contain both the crystalline and the amorphous varieties. According to Fabre (A. Ch. [6] 14, 100), both varieties have practically the same S.H. at c. 100°. Te melts at c. 455° and boils at c. 1390°; the vapour is golden yellow. Heated in air it burns with a brilliant blue-green flame to white clouds of TeO2, which are poisonous; the smell, like that of decaying radishes, that is noticed on burning impure Te is due to the oxidation products of Se, which is commonly present in specimens of crude Te. Te dissolves in very conc. boiling KOHAq, forming a red liquid that contains K₂TeO₃ and K₂Te; on allowing to cool, or on adding water, the whole of the Te separates. HClAq is without action on Te; conc. hot H2SO4 dissolves it slightly, but the Te separates on dilution; when heated strongly with conc. H₂SO₄ Te dissolves to H₂TeO₃ with evolution of SO₂. Te dissolves in SO₃, forming TeSO₃. Conc. HNO₈Aq forms a solution of H,TeO,; aqua regia forms H2TeO, and some H₂TeO₄. Molten KNO₃ produces K₂TeO₄.

In its physical properties Te approaches the metals, and especially resembles Sb. In its chemical properties Te is, on the whole, nonmetallic; but the oxide TeO₂ combines with certain anhydrides of strong acids to form saltlike compounds, and the oxide TeO is said to dissolve in H₂SO₄Aq, forming Te(SO₄)₂. Te is closely related to Se, and less closely to S; it shows distinct analogies with Cr, Mo, W, and U (v. Oxygen group of elements, vol. iii. p. 705, and Chromhumgroup of elements, vol. iii. p. 168). The atom of Te is divalent in the gaseous molecules TeH₂ and TeCl₂, and tetravalent in

the gaseous molecule TeCl4. Atomic weight of tellurium. The at. w. of Te was determined by Berzelius in 1812, 1818, The at. w. of and 1832 by oxidising Te by HNO, Aq and weighing the TeO2 produced; he obtained values varying from 127.9 to 128.5 (P. 8, 24; 28, 395; 32, 16). In 1857 von Hauer analysed K₂TeBr, and obtained values from 126.5 to 128.1 (W. A. B. 25, 135). In 1879 Wills determined the weight of TeO2 obtained by oxidising Te by HNO, Aq and by aqua regia, and he also made analyses of K2TeBrs; his results varied from 126·1 to 129·3 (C. J. 35, 704). In 1888 Brauner obtained numbers varying from 124.94 to 125.4, by oxidising Te to TeO2, by converting Te and TeO2 into (TeO2)2SO2, and by synthesising Cu2Te (J. R. 1883. [1] 433; abstract in B. 16, 3055). In 1889 Brauner made a very careful revision of the at. w. of Te (C. J. 55, 382):—(i.) by oxidising Te to TeO₂ he obtained values varying from 124.48 to 127.28; (ii.) by converting TeO₂ into (TeO₂)₂SO₃, the numbers varied from 124.7 to 129 (one result gave 135.9); (iii.) by conversion of Te into (TeO₂)₂SO₃, the numbers varied from 1243 to 129.7; (iv.) by syntheses of tellurides of Ag, Cu, and Au, the numbers varied from 125.7

to 129.9; (v.) by analysis of TeO₂ the value 127.2 was obtained, but only one satisfactory analysis was made; (vi.) by analyses of TeBr. the numbers varied from 127.27 to 127.31. The results obtained by analysing TeBr, were more concordant than any obtained by the other methods; but the value 127.3 is almost .75 units greater than the at. w. of iodine; hence it seems as if the at. w. of Te is greater than that of I. But if this is so, Te must be separated from S. Se, and the chromium metals, which are placed in Group VI. in the periodic arrangement of the elements; and Te must be placed either with the halogens in Group VII. (I being put into Group VI.), or this element must find a place in Group VIII. (I remaining in Group VII., and an empty place being left in Group VI., The chemical properties of Te Series 7). show that it belongs to the same group as S and Se; and the chemical properties of I make it impossible to place that element elsewhere than with F, Cl, and Br.

Is tellurium a homogeneous body? Rather than doubt the applicability of the periodic law, Brauner made the hypothesis that tellurium is not a homogeneous body, but that it contains an admixture of foreign substances. He dissolved Te, and ppd. in fractions by SO₂, fused the fractions with KCy, dissolved in water, and ppd. fractionally by a current of air; he then distilled each fraction in H, and converted it into TeBr, and analysed each fraction. The values thus obtained for the at. w. of Te were almost identical, the mean being 127.33. Pure TeO2 was then dissolved in HClAq, and the solution was ppd. in eight fractions by ammonia; the filtrate from the eighth fraction was ppd. by SO2 in presence of HClAq; after boiling off SO₂ from the filtrate from the ninth fraction, the liquid was saturated with H₂S, and the TeO₂ in solution was obtained, converted into Te, and this converted into TeBr4; the at. w. thus obtained (in three experiments) was 127.32, the three experi-Te was then ments agreeing extremely well. fused, but not distilled, in a current of H; it was then converted into TeBr4, which was freed from TeBr₂ by fractional sublimation in vacuo, without converting the TeBr, into vapour, and analysed; the values obtained for the at. w. of Te differed very much, ranging from 128.5 to 187.4. From these results Brauner concluded that 'if tellurium, obtained by any process of fractionation whatever, be subjected to distillation in a current of hydrogen, and the tetrabromide be prepared from this by sublimation in a vacuum, its analysis always gives the same atomic weight for tellurium. . . . If, however, the tellurium be only fused in a current of an indifferent gas, and the tetrabromide be not sublimed, higher numbers will be obtained. . . . It is therefore very probable that on distilling tellurium in a current of hydrogen, and the tetrabromide in a vacuum, one constituent of tellurium escapes partly, the presence of which in the last cases renders the atomic weight higher.' Brauner prepared TeBr. by fusing TeBr, with excess of Te, and subliming either in vacuo or in a current of CO2; the analyses of different specimens of TeBr, showed percentages of Br varying from 54.658 to 55.14; the percentage of Br calculated for Te = 127.32 is 55.6. Brauner concluded that 'tellurium is

not a simple substance.' This conclusion is still held by B. to be correct. In a private communication (January 1894) he says: 'I consider Te to be a mixture of two elements that cannot be separated by ordinary chemical means, not even by fractionation. It cannot, however, be shown that the at. w. of Te is about 125, for all attempts I have made in this direction have failed completely. All atomic weight determinations made by me, and others, giving values higher or lower than 127.8 are due, partly to impurities, but in most cases to some undetected source of error.'

Molecular weight of tellurium. Deville a. Troost (C. R. 56, 891) found the V.D. of Te between c. 1890° and c. 1440° to be 180; this gives the mol. w. of c. 260. As the at. w. of Te is not known with certainty, it is not possible at present to correct the value deduced for the mol. w. from the determinations of V.D. The

gaseous molecule of Te is diatomic.

Reactions and Combinations.—1. Heated in air, or oxygen, Te is burnt to TeO2.—2. Heated in excess of chlorine TeCl, is produced; with bromine TeBr, is produced; with iodine the product is Tel.—3. Te and sulphur can be melted together in all proportions; it is doubtful whether definite compounds are formed or not (v. Tellurium sulphides, p. 655). Te and selenion can also be melted together in all pro-Te and portions; no definite compound has been isolated .- 4. Nitric acid forms H2TeO3; according to Klein a. Morel (Bl. [2] 43, 198), cold HNO₃Aq first forms 4TeO₂.N₂O₃.xaq (v. Tellurium DIOXIDE, p. 655). Aqua regia produces a mixture of H₂TeO₃ and H₂TeO₄.—5. Te dissolves in hot conc. potash solution, forming a red solution that contains K2Te and K2TeO3; on addition of water Te separates $(2K_2TeAq + K_2TeO_3Aq)$ = 3K₂OAq + 3Te).—6. Fusion with potassium nitrate produces K₂TeO₄.—7. Fusion with potas sium carbonate produces a mixture of K, Te and K₂TeO₂.—8. Te dissolves in sulphur trioxide, forming TeSOs (v. thio-oxide under Tellurium oxides, p. 655).

Detection and Estimation. — Free Te is readily detected by fusing with charred cream of tartar and dissolving in water, whereby a red solution is obtained that deposits Te in darkbrown flakes on standing in air. Tellurous acid and tellurites dissolve in dilute HClAq, and addition of water ppts. white H₂TeO₃; SO₂ ppts. Te from solutions of tellurites in dilute HClAq or HNO₃Aq. Solutions of tellurates in acids give no pps. with water; solutions in HClAq give off Cl on boiling, being reduced to tellurites, and are then ppd. by water. For a test for Te based on the formation of red TeSO₂ by passing TeH₂ into H₂SO₄Aq v. Divers a. Shimosé, C. J. 43, 329. Te is generally estimated by ppg. the element, from solution of H₂TeO₂ or a tellurite, by alkali sulphite, SO₂, or grape sugar (Kastner, abstract in C. J. 29, 440; cf. Donath, abstract in C. J. 60, 242). In C. J. 59, 238, Brauner describes a method for the volumetric estimation of Te by KMnO₂Aq.

Tellurium, acids of; tellurhydric acid H.Te (vol. ii. p. 727); tellurous acid H.TeO. (this vol., p. 656); telluric acid H.TeO. (this vol., p. 649). The existence of the potassium salt of HCNTe is probable (v. Tellurocyanides,

vol. ii. p. 353).

Tellurium, alloys of. The compounds of Te with metals are sometimes classed with the more definite alloys (v. Tellurides, p. 649).

Tellurium, antimonides of, v. Antimony; Combinations, No. 7, vol. i. p. 283.

Tellurium, arsenides of, v. Arsenic; Combinations, No. 6, vol. i. p. 303.

Tellurium, bromides of. Two bromides have been isolated, TeBr₂ and TeBr₄; both are formed by the direct combination of Te and Br. The bromides can be sublimed, under reduced pressure, without change. The V.D. of neither has been determined; but, from the analogy of TeCl₂ and TeCl₄, the formulæ are probably molecular.

Tellurium dibromide TeBr₂. Formula probably molecular, from analogy of TeCl₂. Prepared by fusing TeBr₄ with excess of powdered Te, and subliming either in vacuo or in a current of dry CO₂. A blackish-green crystalline mass; obtained in steel-grey needles by slow sublimation (Brauner, O. J. 55, 410). Melts c. 305°, and boils c. 343° (Carnelley, Melting- and Boiling-point Tables, 1, 21). TeBr₂ is very hygroscopic; it is decomposed by water to H₂TeO₂Aq and HBrAq. Dissolves in tartaric acid solution, forming TeBr₄ and Te (Brauner, Lc.).

I.c.).
TELLURIUM TETRABROMIDE TeBr₄. (Telluric bromide.) Formula probably molecular, from analogy of TeCl. Prepared by shaking together powdered Te and Br, in the proportion Te:4Br, warming in a current of dry CO₂, and then sub-liming in vacuo (for details and description of apparatus, v. Brauner, C. J. 55, 896). Forms a crystalline crust, which is fiery red when hot and orange coloured when cold. S.G. $\frac{150}{40} = 4.31$ (Brauner, I.c., p. 407). Melts at c. 380°, and boils at 414°-427°, according to Carnelley a. Williams (C. J. 35, 563; 37, 125). Dissolves in a little water without change; on dilution a colourless liquid is obtained, containing HBr and H₂TeO₃; crystals are obtained by evaporating over H₂SO₄, and are said by Berzelius to be TeBr₄...æH₂O. By adding AgNO₃Aq to a solution of TeBr₄ in tartaric acid, Brauner (C. J. 55, 398) obtained AgBr and also crystalline scales probably (C,H,O₅)2.Ag2O.TeO2.2aq; this compound is called by B. tellurium-silver tartar emetic (cf. Klein a. Morel, C. R. 100, 1140). TeBr, forms double salts with alkali bro-The salt TeBr. 2KBr (= K2TeBr.) is prepared by dissolving the proper quantity of KBr in water, adding Te, running in Br, shaking till the Te is dissolved, filtering, and evaporating at 100° (Wills, C. J. 35, 711; cf. von Hauer, W. A. B. 25, 135). Dark-red, opaque, lustrous erystals; orthorhombic, a:b:c=1:1.4901:1.3658(W., l.c.); cf. Wheeler (Am. S. [3] 45, 267), where some other salts of the form M2TeBr are described.

Tellurium, chlorides of. Two compounds are known, TeCl₂ and TeCl₄; both have been gasified, and the formulæ of both are molecular. The chlorides are formed by the direct combination of their elements.

TELLURIUM DICHLORIDE TeCl₂. Mol. w. c. 196. Powdered Te is heated in a distillation-flask in a stream of Cl until the Te is completely converted into TeCl₄; Te is added equal

in weight to the original quantity used, the whole is heated for a little time, an inverted condenser being attached to the flask, and the TeCl₂ is then distilled off at 320°-330° (Michaelis B. 20, 2488). TeCl₂ is a black, amorphous solid melting at 175° (M., l.c.) to a black liquid Carnelley a. Williams give m.p. as 209° (C. J. 37, 125). Boils at 327° (C. a. W., l.c.). The vapour of TeCl₂ is reddish, and gives a characteristic absorption-spectrum (M., l.c.). V.D. at 440°, in vapour of N, = 98·2 (M., l.c.). The vapour becomes yellowish in presence of air or O; TeO₂ and TeCl₄ are formed, and after continued heating these react to produce TeOCl₄ (M., l.c.). TeCl₂ absorbs moisture when exposed to the air, but does not fume; on addition of much water H₂TeO₃ is formed; HClAq forms Te, and TeO₂ which remains dissolved in the acid (M., l.c.).

(M., l.c.).

TELLURIUM TETRACHLORIDE TeCl., (Telluric chloride.) Mol. w.c. 269. Powdered Te is heated, in a distillation flask, in a stream of dry Cl until the solid becomes pale yellow, when it is distilled in a stream of dry CO, (Michaelis, B. 20, 1780). TeCl, is a colourless, crystalline solid, melting at 214°, and boiling without decomposition at 380° (M., l.c.; Carnelley a, Williams give the m.p. as 224° [C. J. 37, 125], and the b.p. as 414° [C. J. 35, 563]). V.D. 131 at 440°, 125 at 530° (M., l.c.). The vapour of TeCl, shows no absorption-spectrum (M., B. 20, 2488). TeCl, dissolves without change in dilute HClAq. It deliquesces in the air, cold water ppts. an oxychloride, and H₂TeO, is also formed; boiling water dissolves TeCl, and H₂TeO, separates on cooling.

TeČl, combines with alkali chlorides to form salts TeCl, 2MCl = M, TeCl,; these salts are best obtained by adding MClAq to a slight excess of TeCl, dissolved in dilute HClAq and crystallising (v. Wheeler, Am. S. [3] 45, 267). The salts crystallise in regular yellow octahedra, without water. Weber (P. 104, 422) described a compound TeCl, 3AlCl, obtained by melting together TeCl, and AlCl,. TeCl, absorbs NH,, forming a greenish-yellow solid that is unchanged in air and has the composition TeCl, 2NH, (Espenschied, A. 113, 101).

Tellurium, fluorides of. Only one fluoride of Te has been isolated. The formula TeF, is probably molecular, from the analogy of TeCl.

probably molecular, from the analogy of TeCl₄.

Tellurium tetrafluoride TeF₄. (Telluric fluoride.) Prepared by dissolving H₂TeO₂ in HFAq, evaporating at 100° to a syrup, allowing to cool, separating the white nodules that are formed, and heating in a Pt basin (Högborn, Bl. [2] 35, 60). If carbonate or hydroxide of an alkali metal, or of Ba, is added to the solution of H₂TeO₂ in HFAq before evaporation, double salts of the forms TeF₄.M⁴F and 2TeF₄.M⁴⁴F are obtained; these salts are decomposed by water (H. l.c.).

water (H., l.c.).

Tellurium, haloid compounds of. These compounds belong to the forms TeX2 and TeX4, where X = Cl, Br or I; the only fluoride that has been isolated is TeF4. The chlorides TeCl2 and TeCl4 have been gasified, and the formulæ are molecular; the formulæ of the other haloid compounds are also probably molecular. An iodide containing more I than TeI4—perhaps TeI4, corresponding with SI4—may exist in the

solution of H₂TeO₂ in HIAq. No compound corresponding with S₂Cl₂, Se₂Cl₂, Se₂Br₂, and Se₂I₄ has been isolated. The haloid compounds of Te are more stable towards heat than the corresponding compounds of Se or S. The haloid compounds of Te are decomposed by water, giving H₂TeO₃ and HXAq; cold water is said to ppt. an oxychloride from TeCl₄. The compounds TeX₄ combine with the haloid compounds of the alkali metals to form salts M₂TeX_a; when X = F the salts are said to be of the form MTeF₂.

Tellurium, hydride of, TeH₂ v. Hydrogen telluride, vol. ii. p. 727.

Tellurium, iodides of. Two iodides have been isolated, TeI₂ and TeI₄; a third—perhaps TeI₈—may exist in the solution of H₂TeO₂ in HIAq. TeI₂ is formed by directly combining Te and I; TeI₄ is obtained by digesting H₂TeO₂ with HIAq.

TELLURIUM DI-IODIDE Tel. Prepared by gently warming a mixture of powdered Te and excess of I; may be sublimed as a black, crystalline crust; when strongly heated I is given off. Not changed by water, hot or cold (Berzelius, Lehrbuch, 5th edit., 3, 1139).

TELLURIUM TETRA-IODIDE Tel. (Telluric iodide.) Obtained by digesting powdered H₂TeO₃ with HIAq in a closed flask, whereby hard black granules are formed. Melts when heated, and gives off I. Boiling water causes decomposition, probably forming an oxyiodide. If a solution of H₂TeO₃ in HIAq is evaporated over H₂SO₄ and CaO₂H₂ in vacuo, lustrous prisms are obtained that are perhaps a compound of Tel₄ and HI (B., l.c.). By dissolving alkali iodides in Tel₁ dissolved in dilute HIAq, and crystallising, black salts of the form M₂Tel₄ are formed in regular octahedra; the potassium salt contains 2H₂O and crystallises in monoclinic forms (Wheeler, Am. S. [3] 45, 267).

Tellurium, oxides of. Three oxides have been isolated, TeO, TeO,, and TeO,. TeO is probably slightly basic; TeO, is the anhydride of the weak acid H,TeO,, but it also shows feebly basic properties; TeO, is a very weak acidic oxide. None of the oxides has been gasified; all are known in the solid state only. An oxide of Te and S, TeSO, is also known.

Tellurium monoxide TeO. This oxide was prepared in 1883 by Divers a. Shimosé (C. J. 43, 819). It is obtained by long-continued heating TeSO₃ to 180°-230° in vacuo until SO₂ ceases to be given off, powdering the residual solid, digesting it with water containing a little Na₂CO₃, washing with hot water, then with alcohol, and drying in a steam oven. TeO is a black, amorphous solid, with a slight brown shade, showing a graphitic lustre when pressed with a hard body. It is unchanged in air, but when heated it burns to TeO₂; when strongly heated in vacuo TeO₂ and Te are formed. TeO is scarcely affected by cold KOHAq; boiling KOHAq decomposes it, giving Te. SO₃, even if boiling, scarcely acts on TeO. The oxide dissolves in hot H₂SO₄Aq; D. a. S. say that the solution deposits Te(SO₄)₂ on cooling, but no analyses of the crystals that separate are given; Magnus (P. 10, 491) gave this formula to the product of the interaction of Te with hot conc.

H₂SO₄. Heated in HCl gas TeOl₂ is formed. TeO is slowly reduced to Te by SO₂.

TELLUBIUM DIOXIDE TeO₂. (Tellurous oxide.

Tellurous anhydride.)

Occurrence.—As tellurite or tellurium ochre, in the Seven Mountains.

Formation.—1. Te is heated in the air.—2. By decomposing a boiling solution of TeCl, in HClAq by boiling water.—3. By heating H₂TeO₃.
4. By oxidising Te by HNO₃Aq, and heating the solution.—5. According to Hilger (A. 171, 211) by heating Te with H₂SO₄, SO₂ is evolved and TeO₂ separates on cooling.

Preparation.—Powdered Te is dissolved in warm HNO₃Aq, S.G. 1·25, and the solution is heated somewhat above 20°; below 8° H.TeO₃ separates, between 8° and 20° both TeO₂ and H.TeO₃ are formed, and the solid that forms from the solution above 20° is TeO₂ only (Klein a. Morel, Bl. [2] 43, 198). By allowing the solution of Te in HNO₂Aq to stand for some hours, and then adding some alcohol, TeO₂ is obtained in crystals (Oppenheim, J. pr. 71, 267;

cf. Berzelius, P. 28, 392).

Properties. - A white, crystalline solid; melts at a red heat to a transparent, yellow liquid, which on cooling forms a white crystalline mass, giving off so much heat that the solid glows feebly. TeO₂ may be sublimed in a slow stream of air. Clarke (Am. S. [3] 14, 285) gives S.G. as 5.7559 at 12.5°. As obtained from a solution of Te in HNO₃Aq, TeO₂ forms quad ratic octahedra; S.G. 5.65 to 5.68 at 0° (K.a. M., l.c.). As obtained by heating H₂TeO₃ till all water is removed and the residue melts, TeO₂ forms orthorhombic needles; S.G. 5.88 to 5.91 at 0° (K. a. M., l.c.). Freshly-prepared TeO2 has no taste, but after a time it acquires a disagreeable metallic taste (Berzelius, P. 28, 392). It does not redden litmus paper. It is almost insoluble in water; 1 pt. dissolves in 150,000 pts. of water (K. a. M., *l.c.*). TeO₂ is very slightly soluble in most acids; it is more soluble in HClAq (v. infra, Reactions, No. 3). Dissolves readily in solutions of caustic alkalis, but in alkali carbonate solutions and in ammonia only after long-continued boiling. TeO_2 does not form H_2TeO_2 by reacting with water, but as it is obtained by heating this acid it may be called tellurous anhydride.

Reactions and Combinations.-1. Reduced to Te by heating in hydrogen to the temperature whereat Te vaporises.—2. Easily reduced to Te by heating with carbon.—3. TeO2 absorbs hydrogen chloride, giving off much heat and forming compounds TeO_{xx}HCl. At -10° , after saturation with HCl, the compound TeO_xSHCl is formed; on slightly warming HCl is given off, and TeO₂.2HCl remains, and does not change when heated to 90°; on heating more strongly, TeOCl₂ is formed (Ditte, A. Ch. [5] 10, 82). Hydrogen bromide is also absorbed by TeO2; by saturating TeO_2 with HBr at -15° , a black solid, resembling I, and having the composition TeO2.3HBr, is formed; this compound begins to decompose at -40°, above this temperature TeO_{2.}2HBr is produced, which remains unchanged to c. 800°, at which temperature it decomposes to TeOBr₂ and H₂O (D., *l.c.*). Hydrogen iodide and hydrogen fluoride are also absorbed by TeO, but the products have not been ex-

amined.—4. Tellurites (q. v., p. 650) are formed by dissolving TeO2 in caustic alkali solutions, or by fusing TeO₂ with alkali carbonates.—5. A compound of TeO₂ with sulphuric anhydride 2TeO2.SO3-sometimes called basic tellurium sulphate—is formed by dissolving TeO, in hot H2SO4 diluted with 3 to 4 times its weight of water, and evaporating; it crystallises in rhombic tablets, somewhat soluble in cold dilute H₂SO₄Aq (Klein, C. R. 99, 326).—6. A compound with nitric anhydride 2(4TeO2.N2O3).3H,O is described by Klein a. Morel (Bl. [2] 43, 198), as obtained, in rhombic needles, by dissolving TeO₂ in moderately conc. HNO₃Aq, and also by dissolving Te in excess of HNO₃Aq, S.G. 1.15 to 1.35, and evaporating at a gentle heat. This compound—which is sometimes described as basic tellurium nitrate—is decomposed at c. 330°, giving off N oxides and leaving TeO₂; with much hot water it gives a pp. of TeO₂.—7. A compound, probably TeO₂.Ag₂O₁(C₄H₄O₄)₂, 2aq, is formed by dissolving TeO, in HBrAq, adding tartaric acid, and ppg. by AgNO, Aq (v. Brauner, C. J. 55, 398; cf. Klein a. Morel, C. R. 100, 1140).

TELLURIUM TRIOXIDE TeO_s. (Telluric oxide. Telluric anhydride.) Obtained, as an orangeyellow, crystalline mass, by heating H₂TeO₄ (v. Telluric acid, p. 649) to somewhat above 160°. If the temperature becomes too high some of the TeO, is decomposed to TeO, and O; it is advisable, therefore, to digest the residue with cold HClAq (to remove any TeO2), to wash out HCl by water, and to dry at 100°. TeO, is not changed by cold, nor by boiling, water; nor is it acted on by cold HClAq, by hot or cold HNO, Aq, nor by dilute KOHAq. Very conc. boiling nor by dilute KOHAq. Very conc. boiling KOHAq slowly dissolves TeO_s, forming K₂TeO_s. TeO_s dissolves in boiling HClAq, Cl is slowly given off, and H₂TeO₃ is formed in the solution (v. Berzelius, P. 28, 392). Clarke (Am. S. [3]

14, 286) gives S.G. of TeO, as 5 0704 at 14 5°.
TELLURIUM THIO-OXIDE TeSO, (Tellurium sulphoxide. Tellurium-sulphur trioxide.) This compound is obtained by dissolving pure Te in pure SO₃, and warming after a time to c. 30°, pouring off SO₂, and drying in vacuo (Weber, J. pr. [2] 25, 218; for description of apparatus, and details, v. Divers a. Shimosé, C. J. 43, 323). TeSO, is a red amorphous solid, melting at 30°. When pure it remains unchanged, in a sealed tube, for months (D. a. S., l.c.). The colour gradually changes to brown by heating to 35°; the change is instantaneous at 90°. duct, which is a brown solid, has the composition TeSO_s (D. a. S., l.c.). At c. 180° SO₂ is given off and black TeO remains (D. a. S., Lc.). TeSOs is not acted on by SOs; it dissolves in H_2SO_4Aq , giving an amethyst-red solution; water decomposes it to H₂TeO₃, TeO, Te, H₂SO₄Aq, and H₂SO₄Aq (D. a. S., l.c.).

TeSO, may perhaps be regarded as tellurium

Tellurium, exyacids of, H2TeO2, v. Tellu-ROUS ACID (p. 656); H2TeO4, v. TELLURIC ACID (p.

649) Tellurium, oxybromide of, TeOBr₂. yellow solid; obtained by heating TeO₂.2HBr (v. TELLURIUM DIOXIDE; Reactions, No. 3, p. 654) above 300°. Melts at red heat, forming a very dark-coloured liquid, which gives off almost black vapour with partial decomposition to TeO, and TeBr. (Ditte, A. Ch. [5] 10, 82).

Tellurium, oxychloride of, TeOCl. Prepared by heating TeO.2HCl (v. Tellurium pr OXIDE; Reactions, No. 3) to above 90°. Very similar to TeOCl₂ (Ditte, A. Ch. [5] 10, 82). Decomposed at fairly high temperature to TeO, and TeCl..

Tellurium, oxyiodide of. According to Berzelius (Lehrbuch [5th edit.] 3, 1141), a greyishbrown, heavy powder is formed by digesting TeI, with boiling water, and this powder is probably an oxyiodide of Te; no analyses are given.

Tellurium, salts of. No compounds have been prepared by directly replacing the H of oxyacids by Te. According to Divers a. Shimose (C. J. 43, 319), Te(SO₁), is formed by dissolving TeO in hot H2SO, Aq and cooling, but no analyses are given; Magnus (P. 10, 491) gave the formula Te(SO₄)₂ to the product of the interaction of Te and hot cone. H₂SO₄. The compounds 2TeO₂.SO₃ and 2(4TeO₂.N₂O₃). Saq are sometimes called basic tellurium sulphate and basic tellurium rium nitrate respectively; the compound TeSO, (v. supra, Tellurium thio-oxide) may perhaps be looked on as tellurium sulphite; and a compound, probably TeO2.Ag2O.(C4H4O3)2. 2aq, is called tellurium-silver tartar emetic (v. Tellurium di-OXIDE; Reactions, Nos. 5, 6, and 7, supra).

Tellurium silver tartar emetic v. Tellurium DIOXIDE; Reactions, No. 7, supra.

Tellurium, sulphides of. Berzelius (P. 8, 411) described two sulphides, TeS₂ and TeS₃, corresponding with the oxides TeO₂ and TeO₄. According to Becker (A. 180, 257), CS₂ removes almost all the S from these supposed compounds. B. concludes that the substances described by Berzelius are not definite compounds; he thinks that compounds of Te and S are probably formed by passing H₂S into TeO₂Aq and TeO₃Aq. As the substances obtained by Berzelius seem to have very definite properties, they are shortly described here.

TELLURIUM DISULPHIDE TeS₂. A dark-brown pp. formed by passing H₂S into H₂TeO₄Aq, or into an acidified solution of an alkali tellurite; softens when heated and cools to a grey, somewhat lustrous mass; gives off S when strongly heated. TeS, reacts as an acidic sulphide, forming compounds xMS.TeS2, where M = $(NH_4)_2$, Cd, Li₂, Fe, Mg, Pb, K₂, Na₃, Zn, &c. The thiotellurites of the alkali metals are best obtained by saturating aqueous solutions of the tellurites with H2S, and crystallising in vacuo; the thiotellurites of alkaline earth metals are formed by boiling the sulphides of these metals with TeS2 and water; the thiotellurites of the heavy metals are obtained by ppn. from solutions of the alkali salts (Berzelius, P. 8, 411).

TELLURIUM TRISULPHIDE TeS,. A blackish grey, lustrous solid; formed by saturating H.TeO, Aq with H,S, and allowing to stand for some time in a closed vessel in a warm place. Thiotellurates of the alkali metals, xM2S.TeS. are formed by saturating M2TeO4Aq with H2S (v. Oppenheim, J. pr. 71, 267).

Tellurium, sulphoxide of, v. Tellurium Thio-

OXIDE, supra.

Tellurium, thio-oxide of, v. supra.

Tellurium, thio-salts of, v. Tellurium Disul-PHIDE, p. 655; and Tellurium Trisulphide, p. 655.

M. M. P. M.

TELLUROCYANIDES v. vol. ii. p. 853. TELLUROUS ACID H2TeO.

Formation.—1. By dissolving Te in HNO.Aq, S.G. 1.25, and at once ppg. by water; if the solution is left for some time before adding water the pp. is TeO₂.—2. By decomposing TeCl₄ by cold water.

Preparation.—TeO2 is fused with an equal weight of K2CO3 or Na2CO3 as long as CO2 is given off; the tellurite thus formed is dissolved in water, and HNO₃Aq is added until the liquid has a very distinctly acid reaction; the flocculent pp. thus produced is allowed to remain in contact with the liquid for some hours, and is then thoroughly shaken up with the liquid, a little HNO, Aq being added if the acid reaction disappears during this process; the pp. is then washed with ice-cold water, and dried at the ordinary temperature (Berzelius, P. 28, 392; cf. Oppenheim, J. pr. 71, 267). The solution of the alkali tellurite in HNO, Aq may be decomposed by adding water, but this must be done at once, as after standing water ppts. TeO2.

Properties and Reactions.—A white, light, amorphous solid, with a bitter, metallic taste. Reddens litmus paper. Dissolves slightly in cold water; when the aqueous solution is heated to c. 40° TeO₂ separates. When slightly heated gives TeO_2 and H_2O . H_2TeO_3 dissolves in many acids; from the solution in HClAq, water or alkalis ppts. H,TeO,; the solution in HNO,Aq gives a pp. of TeO₂ on addition of water after standing for some time. From the solution in H₂SO₄Aq and HNO₃Aq the compounds $2\text{TeO}_2.\text{SO}_3$ and $2(4\text{TeO}_2.\text{N}_2\text{O}_5).3$ aq have been obtained (v. Tellurium Dioxide; Reactions, No. 5 and 6, p. 655). H2TeO, forms tellurites (q. v., p. 650). Thomsen (Th. 2, 278) gives $[Te,O^2,H^2O] = 77,180$. M. M. P. M.

TELLURIUM, Organic Compounds of. Methyl telluride Me, Te. (82°). Formed by distilling K.Te with Ba(SO,Me), (Wöhler a. Dean, A. 93, 233; Heeren, Dissertation, Göttingen, 1861). Pale-yellow, mobile, heavy oil, with persistent alliaceous smell. Oxidised by HNO, to Me, TeOHNO, crystallising in prisms .-Me, TeHOAc. Transparent cubes, v. sol. water. Its solution gives with HCl a pp. of Me₂TeCl₂ which crystallises in long prisms [97.5°].—
Me,TeBr, [89°]. Prisms.—(Me,Te),H,CO,. Difficult to crystallise.—Formate Me,TeCH,O,.
Deliquescent needles.—Me,TeI,. Formed by heating Te with MeI at 80° (Demarcay, Bl. [2] 40, 100).—(Me,Te),H,C,O,.—Me,TeO. Got from Me,TeCl, and Ag,O. Deliquescent crystalline mass alkeling to litmus.—Me TeOCl. Got from mass, alkaline to litmus.—Me TeOCl. Got from the chloride and NH, Aq. Short prisms (from alcohol). — (Me,Te),OBr₂. — (M),Te),H₂SO₄. Cubes, v. sol. water, insol. alcohol.

Crystalline, sl. Methyl-iodide Me, TeI. sol. water. Converted by moist Ag₂O into an alkaline base which yields (Me₂TeCl)₂PtCl₄.

Ethyl telluride v. vol. ii. p. 519.

TEMPLIN OIL. An essential oil obtained from fir cones (Flückiger, J. 1855, 642; Berthelot, J. Ph. [3] 29, 38). Colourless, becoming greenish-yellow in air. Boils at 155°-177°. S.G. 12 Yields on rectification a laworotatory terpene (172°) S.G. 2.856. HNO, (1 vol.) and alcohol (\$ vol.) yield terpin hydrate C₁₀H₂₀O₂ aq [118°] (250°) S. 9 at 100°. Alcohol and HCl yield crystalline C10H15HCl and C10H152HCl

TERACONIC ACID C,H₁₀O₄ i.e. CMe₂:C(CO₂H).CH₂:CO₂H. Propylidene-succinic acid. [168°]. Formed in small quantity by distilling the isomeric terebic acid (Geisler, A. 208, 50). Na and NaOEt convert terebic ether into sodium ethyl teraconate (W. Roser, B. 15, 293). Formed also by treatment of the ether CO2Et.CBrPr.CH2.CO2Et with alcoholic potash (Schleicher, A. 267, 130). Crystals, sol. alcohol and ether, v. sol. cold water. Above 163° it yields H2O and an anhydride. Conc. HBrAq at 0° forms terebic acid.

Salts.—BaA".—CaA". Pp., v. sl. sol. water. -Ag₂A". Needles, m. sol. water.

Mono-ethyl ether EtHA". Oil. Decomposes Na₂CO₃, forming crystalline NaEtA", a solution of which gives with AgNO_s a pp. of the unstable AgEtA"

Di ethyl ether Et₂A". (255 i.V.). Oil. **TERACRYLIC ACID** $C_7H_{12}O_2$. (218° i.V.). A product of the distillation of terpenylic acid (Fittig a. Krafft, B. 10, 521, 1659, 1740; A. 208, 79; Amthor, J. pr. [2] 42, 389). Liquid, smelling like valeric acid, sl. sol. and lighter than water. Yields acetic acid on fusion with potash. Fuming HBr forms C₂H₁₈BrO₂, which gradually splits up into HBr and heptolactone C,H12O2. CaA' 5aq. Prisms or needles.—AgA': needles. Ethylether EtA'. (191°). Fruity oil.

TERBIUM. Tr. At. w. not determined with

certainty; probably c. 162 (v. infra).

The examination of gadolinite, a rare Swedish mineral, by Mosander, Cleve, and others made probable the existence therein of at least seven earths, to one of which the name terbia was given (v. Erbrum, vol. ii. p. 456, where the history of these earths is stated more fully, with references to original memoirs). It is still very doubtful whether the substance called terbia is a homogeneous body or a mixture of more than one compound (cf. METALS, RARE, vol. iii. p. 242). Delafontaine (A. Ch. [5] 14, 238) prepared an orange-yellow, earthy compound, which he regarded as pure terbia, from samarskite, by a long process of fractional ppn., first by K₂SO₄Aq, then by oxalic acid, and finally by formic acid (cf. Marignac, A. Ch. [5] 14, 247; Cleve, Bl. [2] 81, 197).

Terbia Tr.O, is described as an orange-yellow amorphous solid; when heated in H for some time it becomes white. Tr₂O₃ dissolves in acids, forming salts of the type Tr₂X₂, where X = SO4, CO3, 2NO3, &c. The emission spectrum of terbia has been mapped by Roscoe a. Schuster (C. J. 41, 283). The at. w. of Tr-the supposed element has not been isolated -- was determined by Delafontaine to be 113.5; Marignac found the value 148.5; Lecoq de Boisbaudran found 163.1, 161.4, and by later work 159.5 (C. R. 102, 395; M. M. P. M. 111, 474).

TERÉBENE v. Terpenes.

TEREBENTHENE v. TERPENES.

DITEREBENTHYL $C_{20}H_{30}$. V.D. 4.6. (845°). S.G. 12 '9688. $[\alpha]_D = 59^\circ$. A product of distillation of colophony (Renard, C.~R.~105, 865; 106, 856). Oil, resinified by air. KMnO, oxidises is

to propionic, acetic, and formic acids and CO2. Cold fuming HNO₃ forms C₂₀H₂₇(NO₂)₃, a yellow powder, sol. alcohol and ether. Br in CS, at -10° forms C₂₀H₂₆Br₂, which on heating yields diterebenthylene C₂₀H₂₈ (347°) S.G. ¹² ·9821, whence Br forms C₂₀H₂₄Br₄. HNO₂ produces C₂₀H₂₅(NO₂)₃, and H₂SO₄ forms C₂₀H₂₇SO-H, all amorphous solids. Bromine-water gives rise to C₂₀H₂₄Br₆, a brown, amorphous solid. Gaseous HCl passed into the ethereal solution forms C₂₀H₂₀HOl. H₂SO₄ yields C₂₀H₂₀SO₃H, which is sol. water, alcohol, and ether, forming fluorescent solutions. Diterebenthyl passed through a red-hot tube forms H, C₂H₄, C₃H₆, pentane, hexane, hexylene, hexinene, heptinene, cymene, and other hydrocarbons.

TEREBENTILIC ACID C₈H₁₀O₂. [90°]. (250°) Got by passing the vapour of the hydrate of oil of turpentine C₁₆H₁₆2H₂O over soda-lime at 400° (Personne, A. 100, 253; cf. Hempel, A. 180, 86). Crystalline powder, sol. hot water, v. sol. alcohol and ether. May be sublimed .- AgA'. Crystalline.

TEREBENTIC ACID CoH14O5. Got by digesting oil of turpentine with oxide of lead (Weppen, A. 41, 294). Crystals (from alcohol). TÉREBIC ACID C,H,O, i.e.

 $CMe_2 < CH(CO_2H) > CH_2$. Mol. w. 158. [176°]. H.C.v. 766, 642 (Ossipoff, C. R. 108, 812). S. (ether) 2.856 at 85° (Amthor, J. pr. [2] 42, 385). Formed by the action of nitric acid on oil of turpentine (Bromeis, A. 37, 297; Rabourdin, J. Ph. [3] 6, 185; Cailliot, A. Ch. [8] 21, 27; Svanberg a. Eckmann, J. pr. 66, 220; Mielck, A. 180, 47; Bredt, A. 208, 37; Erdmann, A. 228, 179). Formed also by oxidising pinol (Wallach, A. 253, 256; 259, 317). Terebic acid is perhaps identical with oxyhexic acid (Gorboff, J. R. 1887, 605). Monoclinic crystals (from alcohol), sl. sol. cold, v. sol. hot water. V. sol. alcohol and ether. Not attacked by fuming HNO₃. Split up by water at 150° into CO, and pyroterebic acid. On boiling with H₂SO₄ (2 pts.) and water (1 pt.) it is converted into the lactone of oxyisohexoic acid $CMe_2 < CH_2 > CH_2$, CO_2 being On distillation it yields pyroevolved. terebic (hexenoic) acid, oxy-isohexoic lactone, and a little teraconic acid. On heating for 15 hours at 160° with excess of a saturated solution of baryta it is converted into acetone and succinic acid (Frost, A. 226, 863). Fuming HIAq yields CH₂Pr.CH₂.CO₂H. Alkalis form salts of diaterebic acid, of which terebic acid is the lactone.

Salts.—NH₄A': very soluble prisms.— KA'\frac{1}{2}aq.—NaA'\frac{1}{2}aq.—BaA'\frac{2}{2}aq: amorphous.— AgA'. Prisms, v. sol. water.

Ethyl ether EtA'. (274° i.V.). S.G. 19 1-111. Formed from terebic acid, alcohol, and HCl. Dilute NaOH dissolves it, forming monoethyl diaterebate; on cautious acidification the ether separates again, but only after warming (Ekmann). Sodium dissolves in its ethereal solution, giving off hydrogen and forming the sodium salt of acid ether of teraconic acid (q.v.). Alcoholic NaOEt acts in the same way, excess forming a white pp. of disodic teraconate.

Diaterebic acid $OMe_2(OH).CH(OO_2H).CH_2.CO_2H.$ The salt BaA" Saq is formed by boiling terebic acid with excess of baryta-water. It crystallises from Vol. IV.

alcohol, but on acidification at once yields the lactone, terebic acid. AgNO, added to a solution of the Ba salt ppts. Ag, A".—CaA" Saq.—CaA".—PbA" Saq.—Pb, A", (OH), aq.

Ethyl ether Et, A". Formed from Ag, A"

and EtI. Oil. Converted by acetyl chloride into CMe2(OAc).CH(CO2Et).CH2.CO2Et, an unstable

B-Bromo-terebic acid C, H, BrO, i.e.

 $\mathrm{CMe_2} < \mathrm{OH_2CO_2H) \atop CO} > \mathrm{CH_2}$. [151°]. Formed by adding Br (1 mol.) to powdered teraconic acid (1 mol.) covered with water (Frost, A. 226, 863). Large crystals, m. sol. CS₂, v. sl. sol. chloroform and benzene. Decomposed by boiling with water into HBr and terebilic acid C,H,O,. Reduced by sodium-amalgam to terebic acid.

OXY-TEREBIO References.—Chloroand

ACID.

TEREBILENIC ACID C, H,O, i.e.

 $CMe_2 < CO_H > CH$ (?) [169°]. Formed by heating a-chloro-terebic acid at 200° (Roser, B. 15, 296; A. 220, 261) and by evaporating βchloro-terebic acid with water (Frost, A. 226, 370). Small prisms or needles (from water), v. sol. alcohol and ether. Crystallises from alcohol or cone. HBrAq in trimetric forms; a:b:c= ·809:1: ·858. May be sublimed. Excess of KOH forms the diaterebilenate K2C,H3O,, but this splits up on warming with water into KOH and potassium terebilenate. Does not combine with Br or HBr. Sodium-amalgam reduces it to terebic

acid. - CaA'2. - AgA'. Prisms.

Reference. - Chloro-terebilenic acid.

TERECHRYSIC ACID C.H.O. A product of the action of HNO, (S.G. 1.2) on turpentine (Cailliot, A. 64, 376). Orange-red crystals, v. e. sol. water, alcohol, and ether.—PbA": crystals.

TERELACTONE C_sH_sO₂ i.e. C_bH_s $<_{\dot{\text{CO}}}^{\text{O}}$. [12°]. (210° i.V). Formed by the action of boiling water on di-bromo-isohexoic acid derived from pyroterebic acid and Br (Geisler, A. 208, 47). Mobile liquid, sol. water. Boiling baryta-water converts it into amorphous Ba(C,H,O,)2, which is sol. water and deposits BaCO, on boiling.

C"H"O" TEREPHTHALIC ACID C₂H₄(CO₂H)₂. p-Phthalic acid. Mol. w. 166. H.C.p. 770,900. H.C.v. 771,200. H.F. 188,100

(Stohmann, J. pr. [2] 40, 139).

Formation.—1. By the action of HNO, on oil of turpentine (Cailliot, A. Ch. [3] 21, 28; De la Rue a. Hugo Müller, A. 121, 86; Schreder, B. 7, 704; cf. W. C. Williams, B. 6, 1094).—2. By the oxidising action of chromic acid mixture on p-xylene (Beilstein, A. 133, 32; 137, 301), cuminic acid, cuminic aldehyde, cymene (Hofrmann, A. 97, 197), p-toluic acid (Beilstein a. Yssel, A. 187, 308), diethyl-beazene, and amyltoluene (Fixig, A. 141, 167).—3. By the action of boiling dilute HNO, on cymene and on terpenes (Schwanert, A. 182, 257; Homeyer, Ar. Ph. [3] 5, 326).—4. By exidation of o-ethyl-cyluene by severely Ph. (Clears of Pierzel). Ph. toluene by aqueous KMnO, (Claus a. Pieszcek, B. 19, 3083).—5. By fusing potassium p-sulphobenzoate with sodium formate [Remsen, B. 5, 879).-6. From its nitrile, which is obtained by distilling K.FeCy, with potassium benzene p-disulphonate (Garrick, Z. 1869, 551), p-chlorobenzene sulphonate (Nölting, B. 8, 1113), or pbromo-benzene sulphonate (Irelan, Z. 1869, 164; Barth a. Senhofer, A. 174, 242; Limpricht, A. 180, 88) .- 7. The semi-nitrile is also formed by the action of a hot solution of cuprous potassium cyanide upon p-diazobenzoic acid (Sandmeyer, B. 18, 1497). -- 8. By heating p-di-bromo-benzene with chloroformic ether and 1 p.c. sodiumamalgam at 110° and saponifying the product (Bonz, B. 18, 2305).—9. By the action of 5 p.c. KMnO, on an oil (258°-263°) which remains as a residue in the manufacture of aniline and toluidine (Hell a. Rockenbach, B. 22, 505).

Preparation.—1. Br (2 mols.) is added to boiling p-xylene (100 g.), and the resulting C_oH₄(CH₂Br)₂ decomposed by alcoholic KOAc on the water-bath. After evaporating the alcohol the C₂H₄(OAc)₂ is extracted with ether, and oxidised by adding 4.5 litres of 10 p.c. KMnO₄ to its solution in water (1 litre) and NaOH (500 g. of S.G. 1.22). Finally more KMnO, is added till the solution remains violet, the solution kept at 100° for 3 hours, filtered, and ppd. by HCl (Baeyer, A. 245, 139).—2. By oxidising p-xylene with chromic acid mixture (Beilstein, A. 133, 41). 3. From p-toluidine by Sandmeyer's reaction, the resulting p-toluic acid being oxidised by

KMnO, (Baeyer a. Herb, A. 258, 1). Properties. - White crystalline powder, nearly insol. water, alcohol, ether, chloroform, and HOAc. Sublimes without previous fusion. Does not yield an anilide on boiling with aniline (Michael a. Palmer, B. 19, 1376). It is ppd. from its salts by phthalic acid. Yields benzene when distilled with slaked lime. Reduced by sodium-amalgam in a current of CO_2 to the $\Delta^{2.t}$ or (1,4)- dihydride. If CO_2 be not used the product is the $\Delta^{1.4}$ or (3,6)- dihydride, which is also get by boiling the (1,4)-dihydride with NaOHAq (Baeyer, A. 269, 153; cf. Mohs, Z. [2] 3, 68). The (1,4)-dihydride gives benzoic acid on oxidation by MnO2 and dilute H2SO4, while the (3,6)-dihydride forms terephthalic acid (Baeyer, A. 269, 182). By heating its alkaline solution with sodium-amalgam terephthalic acid is reduced to a tetrahydride, and this is further reduced by HIAq at 240° to a hexahydride (Baeyer, B. 19, 1805).

Salts. - (NH₄),A". Small crystals. -CaA" 3aq. S. 08 at 6°.—BaA" 4aq. S. 28 at 5°.—SrA". S. 19 at 17°.—Ag₂A". Curdy pp.

Mono-methyl ether MeHA". [c. 230°]. Formed by the action of conc. H,SO, or alcoholic potash on the di-methyl ether (Baeyer, A. 245, 141). Needles, sol. Na₂CO₂Aq and hot water.

Di-methyl ether Me₂A". [140°]. S. 3

at 100°. H.F. 172,300 (Stohmann, J. pr. [2] 43, 2). Formed by heating the acid with PCl, on the water-bath and pouring the product into methyl alcohol. Trimetric plates; $a:b:c=843:1:8\cdot083$. Insol. cold water, sl. sol. hot MeOH, m. sol. EtOH.

Di-ethyl ether Et₂A". [44°]. Prisms. Propyl ether Pr₂A". [81°]. Needles. Isopropyl ether Pr₂A". [56°]. Plates. n-Butyl ether (C₄H₂)₂A". Liquid. Isobutyl ether (CH₂Pr)₂A". [52.5°

Needles, v. sol. ether (Berger, B. 10, 1742).

Isoamyl ether (C₂H₁₁)₂A". Pearly scales. Phenyl ether Ph₂A". [194°] (Baeyer, A. 258, 44). Formed from the chloride and phenol. Leaflets.

Ohloride C.H. (COC!)2. [78°]. (259°). Amic acid C.H. (CONH2).CO2H. [214°]. Formed from p-diazo-benzoic acid by Sandmeyer's reaction (Sandmeyer, B. 18, 1498). Minute plates (from water), m. sol. cold water.

Amide C,H,(CONH2)2. Formed from the

chloride and NH, Aq. Amorphous. Nitrile C_eH₊(CN)₂. [220°] (Luckenbach, B. 17, 1428). Formed from the amide and P2O3. Got also by distilling calcium bromo benzene p-sulphonate with K,FeCy, Conc. HIAq forms C_oH₁(CI₂NH₂)₂ (Biltz, B. 25, 2543). Γ cistrans Δ^{2,5} or (1,4)- Dihydride

CO,H.CH<CH:CH>CH.CO,H. S. 3 in the

cold. H.F. 182,600 (Stohmann, J. pr. [2] 43, 538). Formed by reducing terephthalic acid in a current of CO₂ by sodium-amalgam (Baeyer, A. 251, 257; 269, 153). Monoclinic prisms (from EtOAc); $a:b:c=982:1:1\cdot019$; $\beta=78^{\circ}$ 2'. M. solether. Not attacked by sodium-amalgam in the cold. Unites with bromine (4 atoms). Transformed into the \$\Delta^{1,5}\$ isomeride by boiling with water, and into the $\Delta^{1,4}$ acid by boiling with NaOHAq. Alkaline KMnO, re-oxidises the acid to terephthalic acid. A warm solution of the acid readily reduces AgNOs. On warming with aqueous cupric acetate it gives off CO, and forms a white pp. which, on adding HOAc and warming, yields Cu₂O, the liquid then containing benzoic acid. The (3,6), (3,4), and (2,3) isomerides do not reduce AgNO₃ and cupric acetate. The Ba salt crystallises in plates, and is v. sol. water.

Methylether Me2A". [77°]. Yields a diand a tetra-bromide. Br in CHCl, forms a dibromide [170°] and a tetrabromide [98°].

Di-phenylether Ph.A". [146°]. Formed from the chloride and phenol at 100°. Small crystals, sl. sol. alcohol, ether, and ligroïn.

 $\Gamma cis \Delta^{2,5}$ or (1,4)- Dihydride

CO₂H.CH CH:CH C(CO₂H)H. Extracted by

ether from the mother-liquor from which the preceding isomeride has separated. Closely resembles its cistrans isomeride, but the cis acid and its salts are the more soluble.

Δ1,5 or (3,4)- Dihydride

CO₂H.CH CH:CH C.CO₂H. S. 042 in the

cold. H.F. 185,300. Formed by boiling the (1,4)- dihydride with water, and obtained, therefore, by reducing terephthalic acid with sodiumamalgam (Baeyer, A. 245, 142; 251, 257; 269, 148). Converted into the (3,6)- acid by NaOHAq. Sodium-amalgam in the cold reduces it to the Δ² tetrahydride.—BaA" 4aq. Trimetric crystals; a:b:c = .319:1: .352 (Baeyer a. Herb, A. 258, 22).

Methyl ether Me,A". [40°]. $\Delta^{1,4}$ or (3,6). Dihydride

CO₂H.O CH₂·CH CO₂H. S. *0059 in the

cold. H.F. 191,900. Formed by reduction of terephthalic acid in alkaline solution by sodiumamalgam (Baeyer, A. 245, 142). Got also by boiling the (1,4)-dihydride with NaOHAq. Slender needles (from water), almost insol. ether. Much resembles terephthalic acid. limation it is partially converted into terephthalis

acid. Immediately oxidised by KMnO. Forms CH(CO,H) CH, CHBr CH, CH.CO,H by uniting with HBr. Sodium-amalgam in the cold reduces it to a slight extent, forming the two isomeric Δ^2 tetrahydrides.—BaA"4aq. Crystals, resembling its $\Delta^{1,5}$ and $\Delta^{1,8}$ isomerides.

Mono-methyl ether MeHA". 「225°7. Got from Me, A" and alcoholic potash. Needles

(from hot water).

Di-methy i ether Me₂A". [130°]. H.F. 172,700. Formed from the dihydride by successive treatment with PCl, (2 mols.) and MeOH. Monoclinic plates (from EtOAc); $a:b:c=1.52:1:2\cdot79$; $\beta=c.74^\circ$. Sl. sol. water, m. sol. hot alcohol, v. e. sol. ether. CHCl, yields a dibromide C,H,Br,(CO,Me)2 [90°] and a tetrabromide [150°]. Excess of Br yields CO₂Me.C₆H₆Br<0 [188°]. HBr forms C₈H₈Br₂(CO₂H)₂ which yields C₈H₈Br₂(CO₂Me)₂

Di-phenyl ether Ph2A". [191°]. Formed by treating the chloride with phenol (Baeyer a. Herb, A. 258, 31). Small scales (from hot

MeOH), sl. sol. ether.

Δ 1.8 or (5,6). Dihydride

CO,H.C & CH, CH, > C.CO,H. S. .0053 in the cold. Formed by the action of alcoholic potash on the dibromide of the Δ^2 tetrahydride and on the tetrahydride of di-bromo-terephthalic acid got by bromination of the hexahydride. Unites with hydrogen bromide (2 mols.) forming CO₂H.CH

CH₂ .CH₂ >CH.CO₂H. Easily reduced by Zn and HOAc and by sodium-amalgam to the two A2 tetraliydrides. Boiling water conit into the (3,4)- isomeride. Boiling NaOHAq forms the (3,6)-isomeride.—BaA" 4aq.

Needles (from hot water).

Methyl ether Me.A". [85°]. Monoclinic plates; $a:b:c=2\cdot241:1:3\cdot591$; $\beta=87^\circ$ 13'. Slowly converted by HBr into (2,3)-di-bromo-terephthalic acid hexahydride, which is reduced by zinc-dust and acetic acid to the Δ^2 tetrahydride of terephthalic acid. Yields a dibromide [64°].

Di-phenyl ether Ph.A". [175°]. needles (from MeOH), sl. sol. cold alcohol.

Δ' or (3,4,5,6)- Tetrahydride.

CH(CO,H) CH, CH, CO,H. [above 800°]. S. 102 at 16°; 83 at 100°. H.F. 214,200 (Stohmann, J. pr. [2] 43, 5). Formed by boiling a solution of terephthalic acid (1 pt.) in NaOHAq for twenty hours with gradual addition of 4 p.c. sodium-amalgam (100 pts.) (Baeyer, B. 19, 1805; A. 245, 160; 258, 82). Prisms, more sol. water than terephthalic acid or its dihydrides. Combines with HBr and with Br (1 mol.). Immediately reduces alkaline KMnO, yielding oxalic acid.—BaA"3 aq. M. sol. cold water. Reduced by HIAq at 240° to the hexahydride.—Ag.A". Amorphous.

Methylether Me,A". [39°]. H.F. 196,200. Plates. Its ethereal solution shows blue fluor-NaOEt. HBr gives C_sH_sBr(CO₂H)_s, whence Me_sA'' [95°]. Bromine forms C_sH_sBr₂(CO₂Me)_s [81°]. escence and gives a fugitive rose-red pp. with

Di-phenyl-ether Ph.A". [145°]. Formed

from the acid by successive treatment with PCl, and phenol. Monoclinic crystals; a:b:c= 2.824:1:2.470, m. sol. cold alcohol and ether.

 Γ cistrans Δ^2 or (1,4,5,6). Tetrahydride

CH(CO,H) CH; CH; CH, CH,CO,H. [c. 220°]. S. 17. Formed by reducing \$\Delta^{1,6}\$ dihydride and also the dibromide of the $\Delta^{1,8}$ dihydride. dised by KMnO, in the cold to succinic acid and a soluble acid [150°]. Alkaline K_{*}FeCy_{*} gives terephthalic acid. Yields three dibromides CH(CO₂H) CH₂ .CH₂ .CH₂ CH.CO₂H. [171°], [51°], and [94°]. The Ba and Cd salts are more sol. water than those of the A' isomeride.

Methyl ether Me2A". [c. 3°].

Amide. Dimetric needles; a:c=1:2·151. Diphenyl ether PhA". Formed, in two modifications [107°] and [c. 190°] by the successive action of PCl, and phenol (Baeyer a. Herb, A. 258, 39). The modification [c. 190°] probably is a mixture containing the Δ^1 isomeride.

Dibenzyl ether (C,H,)2A". [48°]. From

Ag2A" and benzyl chloride.

 Γ cis Δ^2 or (1,4,5,6)- Tetrahydride. [150°-155°]. S. 2.7 in the cold. Formed, together with the cistrans isomeride, by reducing the Δ1.4 dihydride by sodium-amalgam in the cold. The Ba, Cd, and Ag salts are amorphous. This acid and the cistrans isomeride are converted into the Δ^1 isomeride by boiling with NaOHAq.

I cistrans Hexahydride

CO.H.CH CH, CH, CH, CO. H. Fumaroid or stable modification. [300°]. S. 086 at 16°; 1.3 at 100°. H.F. 236,500 (Stohmann, J. pr. [2] 43, Formed by heating the tetrahydride with HIAq for 6 hours at about 240° (Baeyer, B. 19, 1806; 245, 170; 251, 257). Formed also by heating ethyl butane tetracarboxylate with NaOEt and ethylene bromide at 100°; the product being hydrolysed and heated at 220° till evolution of CO, ceases (Mackenzie a. Perkin, jun., C. J. 61, 174). Prepared by reducing the hexahydride of bromo-terephthalic acid with zinc-dust and HOAc. Small prisms, sol. hot water. May be sublimed. Not oxidised by cold alkaline KMnO₄. Br (2 mols. at 100°) forms C₄H₂Br(CO₂H)₂ [71°], and a maleic isomeride [205°]. Treatment with PCl₄ followed by Br at 150° forms C,H,Br,(CO,H), in a fumaroid modification which yields Me,A" [150°] and a maleic modification which yields Me,A" [68°] and an anilide [200°]. The K and Ba salts are v. sol. water; the Ca salt is sl. sol. water.

Methyl ether Me,A". [71°]. S. .5 at 100°. H.F. 218,100. Exhibits no fluorescence. Volatile with steam.

Diphenyl ether Ph.A". [151°]. Needles. I cis Hexahydride

 $CO_2H.CH \stackrel{CH_2.CH_2}{\underset{CH_2.CH_2}{\leftarrow}} C(CO_2H)H.$ Maleic or labile modification. [162°]. H.F. 287,400. Formed by reducing the very soluble hexahydride of bromo-terephthalic acid with sincdust and HOAc. Plates (from water), v. sol. alcohol and ether. Changes on heating with HClAq at 180° into its isomeride. Its Ba sals and methyl ether do not crystallise.

References. - BROMO-, CHLORO-, NITEO-, and

OXY- TEREPHTHALIC ACID.

TEREPRIHALIC ALDEHYDE

C.H.(CHO),[1:4]. [116°]. (247°). S. 1.7 at 100°. Formed by boiling C.H.(CH,Cl), or C.H. (OH.Br)2 (1 pt.) with lead nitrate (1 pt.) and water (20 pts.) (Grimaux, C. R. 83, 825; Löw, A. 231, 863). Formed also by the action of fuming HNO₃ on di-w-bromo-p-xylene (Lōw, B. 18, 2072), and by boiling $G_0H_4(CHCl_2)_2$ with water (Colson a. Gautier, Bl. [2] 45, 6, 508). Obtained from $CH_2(OH)_1G_2H_1.CH_2OEt$ by successive treatment with PCl, and water (Colson, Bl. [2] 42, 152).

Preparation.—p-Xylene (1 pt.) is heated with Br (6 pts.) at 140° to 200° with inverted condenser. The crystalline tetra-w-bromo-xylene [169°] thus obtained is heated with three times its weight of H2SO4 (S.G. 1.825 at 120°-180°), the product poured into water, and the needles that separate recrystallised from water (Hönig, M. 9,

1153).

Properties.-Long needles, v. sl. sol. hot water and cold ether, v. sol. alcohol. Slightly volatile with steam. Dissolves in 25 pts. of

saturated aqueous NaHSO, at 45°.

Reactions .- 1. Oxidised by chromic acid mixture to p-aldehydo-benzoic acid and then to terephthalic acid.-2. Conc. NaOHAq forms terephthalic acid, ω-oxy-toluic acid, and di-ωoxy-xylene.-8. KNO, and excess of H2SO, at 110° form nitro-terephthalic aldehyde.—4. Cold conc. NH₃Aq forms tri-p-aldehydo-hydrobenz-amide (C₆H₄(CHO).CH)₃N₂, a crystalline powder, insol. water, alcohol, and ether, yielding on oxidation by KMnO₄ crystalline N₂(CH.C₆H₄.CO₂H)₈ (Oppenheimer, B. 19, 574).—5. Dry or alcoholic NH₂ forms crystalline C₆H₄(CH:NH)₂.—6. NaOAc and Ac₂O give p-aldehydo-cinnamic acid.-7. Acetone and dilute NaOH give a white pp. of C_eH₄(CH:CH.CO.CH₃)₂, which crystallises from ether-acetone in matted needles [156°], insol. water and alcohol, and gives a deep-red solution in conc. H₂SO₄. An intermediate body is CaH, (CH(OH).CH, CO.CH,)2.-8. NPhMe2 and ZnCl, form C,H,(CHO).CH(C,H,NMe,), the leuco derivative of the aldehyde of malachite green .-9. Cold conc. aqueous KCy added to a cold saturated alcoholic solution of the aldehyde ppts. amorphous C₁₆H₁₂O₄, which probably has the formula C,H,(CHO).CH(OH).CO.C,H,(CHO) [170°-174°]. This body reduces cold ammoniacal AgNO_s, forming a mirror. It also reduces warm Fehling's solution and combines with phenyl-hydrazine. KMnO, oxidises it to benzoin di-p-carboxylic acid. NaOHAq dissolves it, forming benzoin di-p-carboxylic acid, di-z-oxy-p-xylene, and other bodies (Oppenheimer, B. 19, 1814).

Phenyl-hydraside [230°].

Oxim C_eH₄(CH:NOH)₂. [200°]. Formed from the aldehyde and an alkaline solution of hydroxylamine (Westenberger, B. 16, 2994). Crystalline, v. sol. alcohol and etter, sl. sol. water. Yields C_sH_{*}(CH:NOEt)_s [55°] and

C.H. (CH:NOAc), [155°].

Reference.— Nitro-terephthalic aldehyde. TEREPHTHALIC AMIDINE C,H,ON, i.e. C₅H₄(C(NH).NH₂)₂. The salt B"2HCl, formed by digesting terephthalic imido-ethyl ether with alcoholic NH₂, is crystalline, and yields B"H₂PtCl₄ (Luckenbach, B. 17, 1436).

TEREPHTHALOPHERONE C. PETHALO-

PHENORE.

TERPENES C₁₀H₁₆; also Sesquiterpenes C₁₃H₂₄, and Polyterpenes (C10H16)n. The greater number of these hydrocarbons exist ready formed as constituents of essential oils secreted by plants. Others are produced from the natural terpenes by the action of heat or of chemical agents. A dihydrocymene isomeric with the natural terpenes has been obtained synthetically from methylisopropyl succino-succinic ester and the homologous dihydroparaxylene and dihydrobenzene by corresponding processes (Baeyer, B. 25, 1840, 2122; 26, 232). The natural terpenes are generally optically active liquids, with right- or lefthanded rotatory power. The only exception is the racemic compound dipentene (q. v.). The following isomerides are known:-

1. Pinene. This includes two varieties, australene or dextropinene and terebenthene or lævopinene. American spirit of turpentine conists chiefly of dextropinene. (156°). [a]₁ = +21·5 (Berthelot, A. 83, 105; 88, 345, 110, 367, Suppl. ii. 226). S.G. ${}_{0}^{0}$ = *8765; ${}_{25}^{25}$ = *8586; ${}_{100}^{100}$ = *8278 (Tilden, unpubl. expis.). French turpentine oil consists almost wholly of lævopinene. (156·5°). [a]₁ = -40·32. S.G. ${}_{0}^{0}$ = *8767; ${}_{1780}^{1780}$ = *8619 (Riban). C. R. 78, 788; 79, 314). It is also present in oil of rosemary (Bruylants, J. 1879, 944), oil of lemon (Tilden, Ph. [3] 9, 654), sage and juniper (C. J. 31, 554), thyme and anise (Brühl, B. 21, 156), and other essential oils.

For production and properties of turpentine oils v. Oils, Essential, Thorpe's Dictionary of

APPLIED CHEMISTRY.

Different specimens of the pinenes, both dextro- and lævo-, obtained by fractional distillation from turpentine exhibit considerable variations in their rotatory power. An optically inactive liquid has been obtained by Wallach (A. 258, 343) from pinene nitrosochloride by treatment with aniline, whereby a diazo- compound is formed, which with the neighbouring chlorine atom gives rise to diazobenzenechloride and regenerates the hydrocarbon

 $-CCl - C.NO - + H_2NC_8H_5$ $= -CCl - C(N:NC_8H_5) - + H_2O \text{ and}$ $-\operatorname{OOl} - \operatorname{O(N:NO_6H_5)} - = -\operatorname{C} = \operatorname{C} - + \operatorname{OlN_2O_6H_5}.$ The hydrocarbon thus obtained boils at 155°-156°, has a density 858 at 20°, a refractive index for D 1.46553 at 21°, and is supposed to be identical in constitution with the pinenes. It is apparently not resolvable into a mixture of dextro- and lævo- pinenes, but with nitrosyl chloride and other reagents it behaves in the same manner as pinene.

Dry pinene unites with one molecule of dry hydrogen chloride, forming a crystalline com pound formerly called artificial camphor [125°]. which appears quite saturated, as it is unacted upon either by excess of hydrogen chloride or bromine. The hydrochloride prepared from dextropinene is dextrorotatory, while that from lævo-pinene is lævorotatory. The compound is very stable; it may be distilled almost without change, mere traces of hydrogen chloride being evolved; and it is unacted upon by aqueous solutions of the alkalis, except at high temperatures under pressure. Heated with sodium stearate, benzoate, or acetate, or with alcoholic potash, it loses the elements of hydrogen chloride and yields solid camphene (v. Camphene, infra). In the presence

of water, alcohol, ether, or acetic acid, pinene takes up two mols. of hydrogen chloride, producing a compound of different character [50°], which on the application of heat is readily split up into hydrogen chloride and a mixture of liquid hydrocarbons (v. DIPENTENE, infra). Nitrosyl chloride passed into a solution of pinene in chloroform at -10° yields a compound C10H16NOCl [103°], which is ppd. as a white crystalline powder on the addition of alcohol (Tilden, C. J., June 1875). The same compound is formed by adding hydrochloric acid to a cooled mixture of the terpene with amyl nitrite and glacial acetic acid (Wallach, A. 245, 245). The nitrosochloride gently heated with alcoholic potash yields a nitroso-compound, $C_{10}H_{13}NO$ [129°] (Tilden). For crystallography v. Story-Maskelyne (C. J., June 1875, and Phil. Mag.). Nitrosopinene unites with two atoms of bromine, forming a crystalline dibromide, which decomposes on melting [132°]. Mixed with alcohol and nitric acid both pinenes form crystallised terpin hydrate C₁₀H_{1.}(OH)₂·OH₂ (Wiggers, A. 33, 358; 57, 247; Tilden, C. J. 83, 247; Hempel, A. 180, 71). Pinene dissolved in carbon tetrachloride and mixed with bromine yields a mixture of products, from which a well-defined crystalline dibromide $C_{10}H_{16}Br_2$ [169°-170°] may be isolated. On heating this with aniline it gives up hydrogen bromide, and ordinary cymene is produced. The yield, however, amounts only to about 10 p.c. of the bromide employed (Wallach, A. 264, 1). Exposed to the action of air or oxygen, in sunlight, the pinenes produce a crystalline compound C10H18O2 (Sobrero, C. R. 33, 66) which when distilled with dilute acid yields a compound called by Armstrong 'Sobrerone' (Armstrong a. Pope, C. J. 59, 311), which is identical with one of the products of the action of nitrous acid on pinene, isolated by Wallach a. Otto, and called by them 'Pinol' (A. 253, 249). Pinene distilled with bleaching powder and water yields a large quantity of chloroform. Pieric acid has no action upon pinenes in the cold, but at the b.p. of the latter a brisk reaction sets in, and if the liquid is afterwards allowed to cool a compound is deposited in colourless scales, which on boiling with alkali yield borneol (Lextreit, C. R. 102, 555; Tilden a. Forster, C. J. 63, 1388).

2. Limonene. This compound, like pinene, occurs in two optically active varieties. Dextrolimonene (175°–176°), S.G. $\frac{20}{20}$ ·846, $[a]_D = +106$ ·8 (Wallach, A. 252, 145) occurs in oils of the fruit of oranges and lemons, also in caraway and dill, &c. The most convenient source is the essential oil of sweet orange, Citrus aurantium. Lævolimonene is obtained from the oil distilled from the leaves of Pinus sylvestris and P. picea. This oil occurs in the drug houses as 'Fir-Wool oil,' but is now much adulterated with common turpentine, and the commercial oil seldom yields more than a small percentage of limonene. (175°-176°), S.G. $\frac{20}{30}$ = 846, [a]_p = -105° (Wallach, A. 227, 287, 246, 221). The limonenes, treated with perfectly dry hydrogen chloride, yield a liquid monochloride. In the presence of alcohol they give, with excess of hydrogen chloride, a quantitative yield of a dihydrochloride [50°], identical with the compound obtained from the pinenes. By the action of nitrosyl chloride (Tilden a. Shenstone, C. J. May, 1877), or by the use of amyl or ethyl nitrite and hydrogen chloride (Wallach), they yield a crystalline nitrosochloride, which, whether formed from dextro- or lavo-limonene, is always a mixture of two isomerides separable from each other by cold chloroform or ether. There are therefore four isomeric limonene nitroso-chlorides, as follows:

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From d. Limonene: 

\alpha compound [103^{\circ}-104^{\circ}] \alpha_{\rm D}=+318^{\circ}

\beta , [105^{\circ}-106^{\circ}] \alpha_{\rm D}=+240^{\circ}

From l. Limonene: 

\alpha compound [103^{\circ}-104^{\circ}] \alpha_{\rm D}=-314^{\circ}

\beta , [100^{\circ}] \alpha_{\rm D}=-242^{\circ}
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From the nitroso-chlorides by the action of heat (Tilden a. Shenstone), or by boiling with alcohol (Goldschmidt a. Zürrer, B. 18, 2220), is produced an isonitroso-limonene, which when made from lævo-limonene is identical in every respect with carvoxim C₁₀H₁₄N.OH, from carvol, extracted from oil of caraway (Goldschmidt a. Zürrer, B. 18, 1732). For a comparison of the rotatory powers of the two limonenes and their derivatives, see Wallach (A. 252, 141). The limonenes are further characterised by the formation of a crystalline tetrabromide [104°-105°] by direct addition of bromine (Wallach, A. 227, 277).

By mixing together equal quantities of dextroand lovo-limonenes an optically inactive mixture is obtained, which behaves in many respects as a distinct hydrocarbon, and appears to be identical with dipentene (q. v.). Strong sulphuric acid acting upon citrene (d-limonene), causes polymerisation, the chief product being a colophene (310°-320°) (Bouchardat a. Lafont, C. R. 115, 1083).

The oil of Licari kanali contains a compound, licareol $C_{10}H_{18}O$, from which a hydrocarbon, licarene $C_{10}H_{18}$, is obtainable by the action of acetic anhydride at 150° (Barbier, C. R. 114, 674, and C. R. 116, 993, and 1062). This terpene appears to consist of impure d-limonene, as it boils at 176°-178°, gives a tetrabromide [103°-104°], forms a nitrosochloride, from which carvoxim [72°] is formed by alcoholic potash; but its specific rotatory power is low, $[\alpha]_D = 7.51^\circ$.

3. Sylvestrene occurs in Swedish turpentine (Atterberg, B. 10, 1206) and in Russian turpentine (Wallach, A. 230, 245), in company with australene and other hydrocarbons of higher boiling-point. (176°-177°). S.G. 18 - 2510; 30 - 3470. [a] - + 66°32 (Wallach, A. 245, 197). The odour of sylvestrene differs from that of pinene and limonene, and resembles the odour of fresh firwood. It forms a liquid monohydrochloride, and a crystalline dihydrochloride [72°], from which the hydrocarbon may be recovered unchanged by heating with aniline. Sylvestrene also gives a nitrosochloride [106°-107°], which, by treatment with alcoholic potash, yields only oily products. The tetrabromide crystallises in monoclinic tables [135°] (Wallach, A. 289, 29).

4. Phellandrene. The seeds of Phellandrium aquaticum were found by Pesci (G. 16, 225) to yield about 2½ p.o. of a volatile oil, consisting chiefly of a dextrorotatory terpene (171°-172°). The same hydrocarbon is contained in the oil of bitter fennel, Faniculum vulgars (Wallach, 4.

662 TERPENES.

289, 40), and in the oil of elemi. Lævo-phellandrene is found, according to Wallach (A. 248 282), in the oil of Eucalyptus amygdalina. Phellandrene is characterised by the formation of a nitroso-nitrite $C_{10}H_{16}N_2O_3$. Pesci obtains it by the action of a nitrite and dilute sulphuric acid upon the hydrocarbon. It crystallises in needles [94°], and although obtained from the dextroctatory hydrocarbon it rotates the plane of polarisation to the left. $[a]_D = -183^{\circ}$ (Pesci). By reducing agents this compound is converted into a base $C_{10}H_{10}(NH_2)_2$ (209°-214°). Ammonia converts it into nitrophellandrene

C₁₀H₁₅NO₂ and an acid.

5. Dipentene, formerly called terpilene or terpinylene. As already stated, this compound is produced when equal quantities of dextroand lavo- limonenes are mixed together (see Pentylenes). It is formed by heating isoprene to a temperature of about 300°. It is also the chief constituent of 'isoterebenthene,' formed by the action of heat on levo-pinene. It occurs among the products of the destructive distillation of caoutchouc. Dipentene is also produced, together with terpinene, terpinolene, and terpineol, by the action of sulphuric acid or phosphoric acid on terpin, or by the action of sulphuric acid on the pinenes under certain conditions (Armstrong a. Tilden, C. J. November, 1879). It is also formed from the dihydrochloride [50°], whether made from pinene or limonene, either by the action of heat upon the chloride alone (Tilden, B. 12, 1133) or by boiling it with a mixture of sodium acetate and acetic acid, or with alcohol and aniline (Wallach). The products thus obtained were formerly supposed to consist of a single hydrocarbon, to which the name 'terpilene' was given. The hydrocarbons known as cinene and cajeputene formed from cineol and cajeputol respectively also consist of dipentene. Dipentene occurs ready formed in the volatile oil of the camphor tree, also in oil of elemi and in Russian and Swedish turpentines (Wallach, A. 227, 296, &c.).

Dipentene boils at about 176°, and has a pleasant smell of lemons. It combines with bromine forming a tetrabromide [124°] unites with 2HCl forming a chloride [50°], from which it may be regenerated by methods given above. It is to be noted, however, that in all cases more or less isomeric change occurs, and the hydrocarbon, whether reproduced by heat alone or by the action of aniline, always contains terpinene and terpinolene, beside cymene and a small quantity of a paraffinoid hydrocarbon (Tilden a. Williamson, C. J. 63, 292). The nitrosochloride C₁₀H₁₄NOCl yields an inactive carvoxim C₁₀H₁₄NOH [93°] (A. 245, 267). A mixture of equal volumes of dextro- and lawolimonene behaves in many respects as though it were a distinct hydrocarbon, as it was supposed to be previously to Wallach's researches. Thus the tetrabromide melts at 124°, while the limonene compound melts at 104°. The inactive isonitrosodipentene or carvoxim, melting at 93°, is formed by mixing together equal quantities of lævo-carvoxim [72°] from dextro-limonene and dextro-carvoxim [72°] from lævo-limonene. The resulting compound gives, by Raoult's freezingpoint method, a molecular weight corresponding to the simple formula C₁₀H₁₈NO (Wallach, A.

246, 230).

Certain derivatives of dipentene exist in two optically inactive forms. When d-limonenenitrosochloride, prepared from either dextro- on levo-limonene, is warmed with an alcoholic solution of piperidine, a mixture of two crystallisable nitrolamines is formed in each case, as follows:

NITROLPIPERIDINES $C_{10}H_{16}NONO_{0}H_{10}$, FROM LEVOLIMONENE:

(a) [93°-94°] Rhombie, from alcohol.
 [a]_D = -67·60°. Hydrochloride dextrorotatory.

(β) [110°-111°]. Monoclinic. Ealmost inactive.

FROM DEXTROLIMONENE:

(a) [93°-94°]. Rhombic. [α]_D = +67.95°.
 Hydrochloride lævorotatory.

(β) [110⁵–111°]. Monoclinic. $[\alpha]_D = -60.48^\circ$. Hydrochloride (?). When the two a-piperidine bases [93°-94°] are dissolved in petroleum-spirit, and the solutions mixed, an inactive nitrolamine [154°] is at once precipitated. A second inactive compound [152] is formed by mixing the β -piperidine bases (Wallach, A. 252, 123). Similar results have been obtained by the employment of aniline and benzylamine. The salts of the bases thus produced possess a rotatory power opposite in direction to that of the free base. No method is at present known of producing from limonene active addition-compounds with two molecules of hydracid. With excess of hydrogen chloride the same inactive dipentene dihydrochloride [50°] is always obtained. The corresponding dihydriodide seems to crystallise in two forms differing

slightly in melting-point (Wallach, A. 239, 13). 6. Terpinene. (180° about). This compound is formed, together with dipentene, by the action of acids upon pinene and terpin, &c. It is said to occur in oil of cardamoms (Ev. Weber, A. 238, 98). Terpinene has not been obtained in an absolutely pure state, but is characterised by forming a nitrosonitrite $C_{10}H_{10}N_2O_8$ [155°] (Wallach, A. 239, 33). The tetrabromide is fluid, and it yields no crystallisable hydrochloride. The nitrosonitrite by the action of bases yields nitrolamines $C_{10}H_{15} < NOH_{15}$

[116°-118°], C₁₀H₁₃<NOH [130°-131°], &c. (A. 241, 315).

7. Terpinolene. (185° about). This hydrocarbon is formed along with dipentene and terpinene by the action of acids upon pinene, &c.

With hydrogen chloride and hydrogen bromide terpinolene unites to form the dipentene dihydrochloride [50°] and dihydrobromide [64°]. It forms an optically inactive tetrabromide which crystallises in monoclinic tables [116°] but is gradually converted at ordinary temperatures into a porcelein-like mass

into a porcelain-like mass.
8. Fenchene. A liquid isomeride of camphor (190°-193°), obtained from oil of fennel, is treated with alcohol and sodium, by which it is reduced to the alcohol C₁₀H_{1,}OH, a colcurless crystalline compound. By treating this with phosphoric chloride the chloride C₁₀H_{1,}Cl is formed, and from this, by heating with aniline, fenchene is optically inactive. It differs from other terpenes in resisting the action of nitrie acid unless heated (Wallach, A. 263, 149).

Synopsis	OF TERPENE	S AND THE	R CHIEF	DERIVATIVES	J.
Boiling-points a	approximately	stated. I	Rotatory	power +	, or 0.

Pinene (156°)	Limonene (176°)	Dipentene (176°)	Sylvestrene (175°)	Phellandrene (170°)	Terpinene (180°)	Terpinolene (185°)	Fenchene (160°)
+ and -	+ and -	±	+	+ and -	0	0	0
O ₁₀ H ₁₀ HCl saturated [125°]	O ₁₀ H ₁₄ HCl unsaturated liquid	C,.H,.HCl unsaturated liquid	OHHCl liquid				O.,H,,HOL
C10H102HCI	C,.H,.2HOl [60°]	C,.H,.2HCl [50°]	OH2HO1 [72°]				
O _{to} H _{ie} Br _e [170°]	O. H. Br. [104°]	O ₁₀ H ₁₄ Br ₄ [124°]	O ₁₀ H ₁₆ Br ₄ [135°]		O ₁₀ H ₁₄ Br ₄ liquid	O.H.Br. [116°]	O, H, Br. liquid
O,.H,.NOOI [103°]	C ₁₀ H ₁₄ NOCl four isomerides [100° to 106°]	OHNOCI [101°-102°]	O _{to} H _{to} NOCl [106°]	O, H, N, O, [102°]	O ₁₆ H ₁₆ N ₉ O ₉ [155°]		
O, H, NO, H, [122°]	C, H, NO NHC, H,	O, H, NO NHO, H, [109°]a	C, H, (NO NHO, H, (7)°)		O ₁₀ H, _e N,O, [137°]		
O, H, NO [132°]	C _{1.} H _{1.} N.OH [72°]	[98°] O'•H'*NOH				7 -	

Isoterpenes.

Camphenes $C_{10}H_{1s}$. Pinene monohydrochloride [125°] was formerly described under the name 'artificial camphor,' from its resemblance to camphor in appearance and to a certain extent in odour. This compound is remarkably stable, but may be decomposed by heating with sodium stearate or benzoate, with potassium acetate, with alcoholic potash, or, better, with a mixture of sodium acetate and alcoholic soda (Brühl, B. 25, 147). The resulting hydrocarbon is a camphene [51°-52°] (160°), dextro- or lavorotatory or inactive according to the nature of the hydrochloride used and the reagent employed, though the exact conditions which determine the production of one or other are scarcely known (Riban, A. Ch. [5, 6, 353).

Camphene is also formed from bornyl chloride by the action of alcoholic potash (Riban), by the action of water and magnesia (Kachler, A. 197, 86), or by heating with aniline to the boiling-point of the latter (Wallach, A. 230, 234); it is also formed from camphor chloride and from pinene hydrochloride by the action of sodium (Montgolfier, C. R. 89, 102). In the last case it is accompanied by a liquid hydrocarbon (170°), C₁₀H₂₀, which behaves like a paraffin, and is probably identical with a liquid obtained by the action of hydrogen iodide on turpentine (Berthelot), also by the action of iodine (Armstrong a. Gaskell, B. 12, 1756) and by the action of sulphuric acid on turpentine (Armstrong, B. 12, 1759).

Camphene is also formed by the action of strong sulphuric acid on turpentine, and constitutes the characteristic ingredient in the liquid formerly known as 'terebene' (Armstrong a. Tilden, C. J. 1879, 733).

Camphene closely resembles camphor in appearance and even somewhat in odour. It is soluble in alcohol, ether, and benzene, and crystallises in leaflets from a concentrated solution. It is incapable of combining with bromine, but it forms a compound [157°] with hydrogen chloride, which is distinguished from pinene hydrochloride by its instability, being rapidly

decomposed by water with reproduction of camphene (Riban, C. R. 80, 1830). It is also dissociated into camphene and hydrogen chloride when volatilised (Ehrhardt, C. N. 54, 239). Camphene hydrochloride is said to be identical with bornyl chloride (Kachler a. Spitzer, A. 200, 340; v. also Brühl, B. 25, 160). Camphene does not combine with nitrosyl chloride. In contact with bromine it is slowly attacked, forming an oily monobromo-derivative (230°-240°).

Oxidised by chromic liquor, camphene yields camphor, dextro-, levorotatory or inactive according to the character of the camphene.

By the action of phosphorus pentachloride on camphene, and subsequent treatment of the mass with an alkaline solution, salts of two phosphonic acids are formed (Marsh a. Gardner, C. J. 65, 35).

When camphene is heated to about 300° it is converted into liquid products which seem to include dipentene, but have not been sufficiently investigated.

Homologues of camphene.

Ethyl-camphene obtained by the conjoint action of sodium and ethyl iodide upon camphor monochloride $C_{10}H_{10}Cl$, is a colourless mobile liquid having a smell like turpentine (197-9°-199-9°, bar. 742·1 mm.). Isobutyl-camphene is also a liquid (228°-229°, bar. 750·4 mm.) (Spitzer, B. 11, 1817).

Sesquiterpenes C18H24.

Oils of clove, calamus, cascarilla, patchouli, and cubebs contain a hydrocarbon of this composition (Gladstone, C. J. 1872), as also do the oils of gabbanum, and savin and 'huile de cade,' a kind of tar made by distillation of the wood of Juniperus oxycedrus (Wallach, A. 238, 81). The hydrocarbon obtained from cade or cubebs (274°-275°) (S.G. 18 - 921) forms the following crystalline compounds: C_{1.}H_{.2}HCl [117°-118°], C_{1.}H_{.2}2HBr [124°-125°], C_{1.}H_{.2}2HI [105°-106°]. The hydrocarbon, especially when partially resinified by exposure to the air, gives the following characteristic colour reaction: dissolved in chloroform or glacial scetic acid and then

shaken up with a few drops of strong sulphuric acid, the liquid assumes an intense green and then blue colour, which, when heated, passes into red (Wallach). An attempt to prepare from isoprene a polymeride of the composition $O_{1s}H_{24}$ proved unsuccessful (Wallach).

Conimene C₁₈H₂₄, a liquid (264°) having a pleasant odour, is obtained from the incense resin (*Icica heptaphylla*) of British Guiana (Stenhouse a. Groves, C. J. 1871, i. 175).

Polyterpenes C20H32, &c.

This group includes (1) the constituents of certain natural essential oils; (2) products of artificial polymerisation of terpenes; and

(3) caoutchouc and guttapercha.

1. Oleo-resin or so-called 'balsam' of copaiba (Copaifera Langsdorffii and other species) is a mixture of a resin with an oil (250°-260°). The latter unites with water and absorbs hydrogen chloride, with production of a deep violet colour but no crystalline hydrochloride. A similar hydrocarbon is obtained from Gurjun balsam or 'wood oil,' the product of various species of Dipterocarpus growing in the East, and from other essential oils.

2. When turpentine oil and other terpenes are heated for some time to about 300°, or treated with concentrated sulphuric or phosphoric acid or other agents, a large part of the hydrocarbon undergoes polymerisation. The name 'colophene' was given by Deville to the less volatile portions of the product obtained in this manner from French turpentine, apparently under the impression that it was closely related to the oil obtained by distillation of resin. Resin-oil, however, contains oxygen, and exhibits quite different characters.

Colophene is a yellowish viscid fluid, usually

fluorescent, which begins to boil at about 800°, but the distillation even under reduced pressure is attended by decomposition. The boiling-point continually rises, and even at a temperature approaching dull redness a viscid residue is left which on cooling becomes nearly solid (Armstrong a. Tilden, C. J. Nov. 1879). The portions which distil at 300° and upwards appear to consist of saturated compounds, for they absorb mere traces of hydrogen chloride. By the action of antimony trichloride upon turpentine a solid, C₄₀H_{e4}, is formed. This combines with hydrogen chloride to form two compounds C₄₀H_{e4}HCl, and C₄₀H_{e4}2HCl (Riban, C. R. 1874, 389).

3. CADUTCHOUG v. vol. i. p. 677. For further information concerning the products of its decomposition by heat v. Pentinenes (vol. iii. p. 807) and DIPENTENE (supra).

GUTTAPERCHA, v. vol. ii. p. 658.

Caoutchouc is formed from isoprene by polymerisation, which sometimes occurs spontaneously under circumstances not fully understood (Tilden, C. N., May 1892).

Oxidised compounds connected with terpenes.

A considerable number of oxidised compounds are known which are evidently closely connected with the terpenes, and in many cases directly derivable from them. Some of these, as, for example, borneol and camphor, occur as natural products in essential oils; others, such as terpin, are the products of the addition of water to a terpene, or, like camphor and some of its isomerides, may be formed from terpenes by oxidation. These compounds are enumerated in the following table, together with the hydrocarbons with which they are presumably immediately connected.

-			
Hydrocarbons	Alcohols	Glycols	Ketones (?)
Cymene C ₁₀ H ₁₄ (ii. 361)	Carveol C ₁₀ H ₁₈ OH (i. 711)	_	Carvol C ₁₀ H ₁₄ O (i.
Pinene C ₁₀ H ₁₆ (v. supra)	Myristicol and absinthol (v. infra)	Sobrerol $C_{10}H_{16}(OH)_2$ (v. infra)	Sobrerone (pinol) C ₁₀ H ₁₆ O (v. infra)
Dipentene $C_{10}H_{16}$ (v. supra)	Terpineol $C_{10}H_{17}OH$ (v. infra)	$\begin{array}{c c} \text{Terpin} & C_{10}H_{18}(\text{OH})_2 \\ (v. infra) \end{array}$	
	Cineol C ₁₀ H ₁₇ OH (ii. 187). Syn. Euca- lyptol, cajeputol	· · <u>-</u>	
Fenchene $C_{10}H_{10}$ (v. $supra$)	Fencheol $C_{10}H_{17}OH$ (v. infra)	_	Fenchone $C_{10}H_{18}O$ $(v. infra)$
Camphene $C_{10}H_{10}$ (v. $supra$)	Borneol C ₁₀ H ₁₇ OH (i. 522)	Camphene glycol C ₁₀ H ₁₆ (OH) ₂ (v.infra)	Camphor C ₁₀ H ₁₆ O
_	Geraniol $C_{10}H_{17}OH$ (ii. 609)		Pulegone C ₁₀ H ₁₆ O (v. infra)
_	Linalool C ₁₀ H ₁₇ OH (iii. 146)	_	Puleoné $C_{10}H_{16}O$ (v. infra)
_	•	_	Tanacetone or Thu- jone C ₁₀ H ₁₆ O (v. infra)
Menthene C ₁₀ H ₁₀ (iii. 202)	Menthol C ₁₀ H ₁₉ OH	-	Menthone C ₁₆ H ₁₈ O (iii. 204)
Dihydrocamphene C ₁₀ H ₁₈ (v. DECINENE, ii. 867)	'-	_	<u>, , , , , , , , , , , , , , , , , , , </u>
Tetrahydrocamphene C ₁₀ H ₂₀ (v. DECYLENE, ii. 869, et infra)	- 1	-	_

Cymene $C_{10}H_{14}$ (176°); ii. 361. Dipentene $C_{10}H_{16}$ (176°); v. supra. Fenchene $C_{10}H_{16}$ (160°); v. supra. Camphone $C_{10}H_{16}$ (160°); v. supra. Menthene C₁₀H₁₈ (167°); iii. 202.

Dihydrocamphene C₁₀H₁₈ v. Decinene, ii. 367. The liquid produced, together with the solid monohydrochloride, by passing hydrogen chloride into pinene, yields when treated with sodium a mixture from which a liquid hydrocarbon C10H18 (148°-149°), having a smell of oranges, may be separated by fractionation, &c. (Bouveault, C. R.

116, 1067).

Tetrahydrocamphene C10H20 (DECYLENE, ii. A liquid obtained by the action of hydrogen iodide and phosphorus upon oil of turpentine (Berthelot) and another (160°) by action of phosphonium iodide on turpentine (Baeyer) have this composition. The same formula is ascribed by Armstrong (B. 12, 1758) to a liquid (160°-170°) obtained by the action of iodine upon oil of turpentine and upon camphor, also by the action of sulphuric acid upon turpentine. these products are insoluble in strong sulphuric acid, and resemble the paraffins in chemical characters.

Carveol C₁₀H₁₅.OH (219°); i. 711.

Terpineol C₁₀⁵H_{1,1}OH. [35°]. (215°-218°). Wiggers, A. 33, 358; 57, 247; List, A. 67, 362; Tilden, C. J. 33, 247, and 35, 286; Wallach, B. 18, 618, Ref.; Kannonikoff a. Flawitzky, J. pr. [2] 32, 497; Bouchardat a. Voiry, C. R. 104, 996. Dextro- and lavo-terpineol are obtained by the action of alcoholic sulphuric acid upon d- and l-pinene respectively, or inactive by distilling terpin with very dilute sulphuric or hydro-chloric acid. Terpineol as thus obtained is a viscous liquid, having an odour of white lilac. By fractional distillation under reduced pressure a portion is obtained (130°-135° at 40 m.m.), which may be crystallised by cooling to -50° , or by the introduction of a crystal of the same. These crystals melt at 30°-32°, and boil undecomposed at 218° (Bouchardat a. Voiry, C. R. 104, 996). Terpineol treated with sodium evolves hydrogen, but in consequence of its viscosity the action soon ceases. Contact with dilute acids converts terpineol into terpin hydrate. Hydrogen chloride and iodide react to form C₁₀H₁₈Cl₂ [50°] and C10H18I2 [77°] respectively. Potassium hydrogen sulphate at 200° produces dipentene. Terpineol unites with two atoms of bromine, but excess of bromine gives rise to dipentene tetrabromide. With phenyl cyanate it yields phenylterpinyl-urethane CoHs.NH.CO.OC10H17 [110°]. These and other reactions have been repeated by Wallach, using crystallised terpineol (A. 275, 103). Terpineol oxidised by permanganate yields a substance C₁₀H₂₀O₃ [121°-122°], which probably has the constitution of an oxyterpin C₁₀H₁₇(OH)₃. Further oxidised by means of chromic acid this yields a crystalline compound

C₁₀H₁₀O₂ [62°-63°] (Wallach, A. 275, 145).
Cineol C₁₀H₁₆O (176°); ii. 187. Cineol appears to contain no hydroxyl, since it is not acted upon by metallic sodium nor by benzoyl chloride at 120°. It is also unaffected by hydroxylamine and by phenylhydrazine. Hence it appears to be neither an alcohol nor a ketone. Its relation to terpineol (infra) is shown by the fact that when heated with

alcohol and sulphuric acid cineol is converted into terpinene and terpinolene: also that terpineol, and therefore also terpin, are partly converted into cineol by prolonged heating with phosphoric acid. Cineol oxidised by permanganate of potassium yields the potassium salts of carbonic, oxalic, and cineolic acids, with a small quantity of acetic acid. Cincol yields 45 p.c. of its weight of cineolic acid. This compound forms well-defined anhydrous crystals, which dissolve in 70 pts. of water at 15°, and in 15 pts. at 100°. They melt with decomposition at 196°-197°.

Calcium cineolate C₁₆H₁₄CaO₅.4H₂O is soluble in cold water, but is completely precipitated by boiling the aqueous solution. Silver cineolate C₁₀H₁₄Ag₂O₅.H₂O is soluble in both water and alcohol, but cannot be crystallised. Ethyl cineolate $C_{10}H_{14}(C_2H_3)_2O_3$, obtained by passing hydrogen chloride gas into an alcoholic solution of the acid, is a colourless liquid (155° under 11-12 mm.). By dry distillation cineolic acid yields cineolic anhydride, water, carbon dioxide, and a liquid which appears to consist of a monobasic acid (A. 246, 265).

Fencheol C₁₀H₁₇.(OH). Fenchyl alcohol [40°-41°]. (201°). (Wallach, A. 263, 143; 272, 99-125). S.G. 50 933 (W.). $[\alpha]_D = \pm 10.86$ (W.). Dextro- and lævo-fenchyl alcohols are obtained by the reduction with sodium and alcohol of lævo- and dextro-fenchones respectively, centrated HNO, oxidises l-fenchyl alcohol to dfenchone. PCl₅ gives rise to fenchyl chloride C₁₀H₁₁Cl (84°-86° at 14 mm.).

Borneol C₁₀H₁₁OH, i. 522.

Geraniol C₁₀H₁₁OH, ii. 609.

Linalcol C10H17.OH, iii. 146. Linalcol is said to be convertible into the isomeric geraniol by treatment with acetic anhydride, and saponifying the resulting ester with alcoholic potash (Bouchardat, C. R. 116, 1253).

Menthol C₁₀H₁₀(OH), iii. 203. Terpin C₁₀H₁₀(OH)₂ [104°-105°]. (258°) (Wiggers, A. 33, 358; 57, 247; Tilden, C. J. 33, 247; 85, 286; Wallach, A. 230, 225-272). Terpin is best known in the form of its hydrate $C_{10}H_{20}O_2.H_2O$ [116-117°], a beautifully crystalline compound which on heating to 100° loses water and leaves terpin as a vitreous mass. Terpin hydrate is readily obtained by shaking turpentine oil with alcohol acidified with sulphuric or nitric acid (Flawitzky, B. 12, 1022; Tilden, C. J. 33, 247; 35, 286), and leaving the liquid to evaporate. It is an optically inactive, saturated compound, slightly soluble in boiling water, soluble in alcohol and crystallising in rhombic prisms (Rammelsberg, P. 63, 570; Maskelyne, P. M. 1879). It possesses the characters of a glycol and with hydrogen chloride yields dipentene dihydrochloride [50°]. Boiled with water containing a mere trace of mineral acid, terpin dissolves with production of terpineol and water.

Camphene glycol C10H15(OH)2.

Wagner, B. 23, 2311).

Formation .- By the oxidation of camphene in benzene solution with a 1 p.c. solution of KMnO, (l.c.).

Properties.—Colourless prismatic needles (from bensene) very readily soluble in ether, alcohol, CS, and chloroform. Exhibits the 666 TERPENES.

same phenomenon as camphor when thrown on water. Sublimes when heated above 100°. Melts when warmed with water, but only dissolves in it with difficulty. Heated with acetic anhydride in a closed tube at 120°, the greater part loses water, only a small portion being converted into the acetic ester. Treatment with dilute HCl results in the loss of 1 mol. of H,O, giving rise to a solid body, C₁₀H₁₀O, isomeric with camphor, which reduces ammoniacal AgNO, and reacts with hydroxylamine.

Carvol $C_{10}H_{14}O$ (228°), v. i. 711. Fenchone $C_{10}H_{14}O$. [5°-6°]. (192°-198°) (Wallach, A. 263, 130; 272, 102). S.G. 12 9465 (W.). $R_A 44.23$ (W.). $[a]_D + 71.70$ and -66.94.

Occurrence.-Dextro-fenchone is present in fennel oil, the fraction boiling between 190° and 195° consisting almost exclusively of the ketone, together with traces of anethol; derived from Thuja oil, fenchone possesses lævorotatory optical properties. D- and l-fenchone form a series of derivatives which bear to one another a relation similar to that subsisting between the two tartaric acids, and the analogy extends to the formation of racemic modifications when equal proportions of the two classes of derivatives are mixed together.

Preparation.—The fraction of fennel oil boiling between 190° and 195° is treated with concentrated HNO, the ketone remaining unattacked. After being freed from acid and distilled in a current of steam, the dry oil is cooled,

and the fenchone crystallises out.

Reactions.—With bromine in the cold, an addition compound is formed, substitution taking place if the mixture becomes heated. P2Os, P2Ss, and PCls have the same action upon fenchone as upon camphor. Fenchone is dissolved by cold concentrated HCl, being thrown out of solution on warming. HNO, has but little action in the cold, but in a sealed tube at KMnO. 120° HCN is formed (A. 263, 134). oxidises fenchone to a mixture of oxalic, acetic, and dimethyl-malonic acids. D- and l-fenchone yield d- and l-fenchyl alcohols respectively on reduction with Na and alcohol (v. Fencheol). On heating d- and l-fenchone with ammonium formate, l- and d-fenchylamines respectively are formed. Fenchone does not react with phenyl-hydrazine or with alkaline sulphites, but both varieties form oxims [161°] with hydroxylamine.

d-Fenchon-oxim C₁₀H₁₅NOH [161°] (240°) loses water on treatment with dilute H.SO., an unsaturated dextrorotatory nitrile (218°) being

formed.

l-Fenchon-oxim C₁₀H₁₆NOH [161°] behaves in a similar manner on treatment with dilute H2SO. The resulting nitrile yields with potash a lavorotatory a isoxim C₂H₁₅.CONH₂ [114°-115°] which is readily converted by dilute H.SO. into the \$\beta\$-isoxim [136°-137°].

Camphor $C_{10}H_{18}O$ (i. 669). Menthone $C_{10}H_{18}O$ (2066) (iii. 204).

Pulegone C. H. O. (130°-131° under 60 mm.). (Beckmann a. Pleissner, A. 262, 1; Wallach, A. 272, 122; Semmler, B. 25, 3515). S.G. $\frac{20}{9323}$ (B. a. P.); $R_{\rm A}$ 45.55 (B. a. P.); $[a]_{\rm D}$ 22.89 (B. a. P.). Pulegone is the chief constituent of Spanish oil of pennyroyal (Mentha pulegium). It is a dextrorotatory colourless liquid, having

an odour of peppermint, and rapidly becoming yellow on exposure to the air.

Reactions.-With HBr it combines forming a lsworotatory ([α]_p = -33.8) addition compound $C_{1o}H_{1o}O.HBr$ [40.5°]. When heated with ammonium formate, Wallach (A. 272, 123) failed to obtain a base isomeric with fenchylamine and thujonamine, a mixture of bases being formed. Slow oxidation with KMnO, gives rise to acetone and β-methyl-adipic acid [84.5°]. Rapid oxidation with the same reagent yields a γ -valero-lacton- γ -acetic acid (Semmler, B. 25, 3516). Pulegone combines with hydroxylamine to form pulegonoxim C₁₀H₁₀NO, [157°]. The oxim contains one molecule of H₂O more than camphoroxim. It is laworotatory ([a]_b -83.4) and forms a hydrochloride [117°-118°], a benzoyl ester [137°-188°], and an acetyl ester [149°]. On heating its hydriodide pulegonamine $C_{10}H_{18}O.NH$ is formed.

Pulegone hydrobromide C₁₀H₁₆O.HBr [40.5°] is laworotatory; it forms an oxim [38°] which becomes converted on standing into normal pulegonoxim [157°]. Pulegone hydrobromide, on treatment with moist silver or lead oxide, loses HBr, pulegone being regenerated. Distillation with zinc-dust gives rise to an oil which resembles menthone in every respect but the melting-point of its oxim (85° instead of 59°).

Constitution .- From the results of oxidation experiments carried on under various conditions with $KMnO_4$ Semmler (B. 25, 3519) attributes to pulegone the constitution:

Pulsone is the name given to a compound isomeric with camphor, obtained by Barbier (B. 25, 110c), from pennyroyal oil, described as boiling at 222°-223°, and possessing the following physical properties: S.G. 9482; 28 9293; [a], +29·15. Its oxim C₁₀H₁₈NOH is an oil (170° under 48 mm.) yielding a liquid anhydride $C_{10}H_{18}N$ on treatment with dilute H_2SO_4 . Chromic acid mixture oxidises it to carbonic, acetic,

and propyl-succinic [89°-91°] acids.

Tanacetone C₁₀H₁₆O (195°-196°; 84·5 at 13 mm.), identical with the thujone of Wallach (A. 272, 109; Bruylants, B. 11, 450; Semmler, B. 25, 3343, 3519; Wallach, l.c.). S.G. $\frac{90}{20}$ 9126

(S.). R. 44.54 (S.). Tanacetone occurs in tansy, sage, wormwood and thuja oils. It has ketonic properties, and possesses an optical rotatory power amounting to $+38^{\circ}$ 30' in a 2-dcm. column.

Reactions.—Sodium in alcoholic solution reduces it to tanacetyl alcohol (92.5° at 13 mm.). With alkaline hypobromite it yields bromoform and tanacetogen-dicarboxylic acid (113.5° at 15 mm.). Oxidised with KMnO, it yields tanacetocarboxylic acid, which exists in two modifications, the α and β thujaketonic acids of Wallach (l.c.), [75°-76°] and [78°-79°] respectively. Heated with ammonium formate, it gives tanacetylamine (198°-199°), the same body being formed when tanacetoxim (see below) is reduced with sodium and alcohol. The hydrochloride of this base on dry distillation yields tanacetene (172°-175°), a hydrocarbon of the formula $C_{10}H_{10}$, identical with Wallach's thujene. Tanacetone reacts with hydroxylamine to form tanacetoxim $C_{10}H_{10}$. NOH (51·5°) (135°-186° at 22 mm.), which yields tanacetylamine on reduction, while treatment with dilute alcoholic H_sSO_4 converts it into a cymidine (Pr:NH_:Me:1:3:4), which gives carvacrol with nitrous acid. Tanacetophorone C_0H_{10} (89°-90° at 13 mm.). R_a 37·6. When tanacetogen-dicarboxylic acid (produced by the action of NaBrO upon tanaceto-carboxylic acid) is heated with soda-lime, tanacetophorone is formed. It has ketonic properties, and combines with hydroxylamine.

Constitution.—Semmler (B. 25, 8519) suggests for tanacetone and tanacetogen-dicarboxylic

acid the formulæ

HC
$$CH_2$$
- $C(Me)_2$ C.Ac and CH_2 — C.COOH CH_2 —HC.COOH. Tanacetone. Tanacetogen-dicarboxylic acid.

Sobrerone $C_{10}H_{10}O$. Pinol. (183°-184°) (Wallach, A. 253, 254; Armstrong, C. J. Proc. 1890, 100). S.G. 20 ·953 (W.); μ_D 1·469 (W.). Sobrerone is obtained from the mother-liquors accumulated in the preparation of pinene nitrosochloride; also by boiling sobrerol with dilute

H₂SO₄ (Armstrong, l.c.).

Reactions.—It combines readily with Br to form a dibromide [94°], from which, on treatment with alcoholic potash, sobrerone is regenerated. Sobrerone and its dibromide yield terebic acid [175°-176°] on oxidation with KMnO₄. HNO₂ produces the same result. It forms a nitroso-chloride [103°] which, by the action of bases, is readily converted into nitrolamines, e.g. sobrerone-nitrol-piperidine [154°] and sobrerone-nitrol-benzyl-amine [183°-134°]. The dibromide O₁₀H₁₀O.Br₂ [94°] (143°-144° at 11 mm.) yields on treatment with alcoholic potash, in addition to sobrerone, its glycol-ether C₁₀H₁₀O(O₂H₃)₂ [52°-53°]. Sobrerone-glycol-diacetate is easily-obtained by heating the dibromide to 150° with lead acetate in glacial acetic acid solution. When heated at 100° for three hours with formic acid, sobrerone dibromide is converted into pinene.

Constitution.—Its behaviour towards bromine and nitrosyl chloride and its molecular refraction indicate the existence of one ethylene linking, while its indifference towards acid chlorides, hydroxylamine, phenylhydrazine and H₂S render it probable that the O- atom is united to two different C- atoms. From these facts, and from its behaviour on oxidation, Wallach (A.

253, 259) assumes the constitution

Sobrerel C₁₀H₁₈O₂. [150°] (Sobrero, C. R. 33, 66; Armstrong, C. J. Proc. 1890, 100; C. J. 59, 815). Sobrerol is obtained by oxidising French and American turpentines in

sunlight. It occurs in two optically active forms, both melting at 150° , and on allowing a mixed solution to crystallise inactive sobrerol [131°] separates out. When boiled with dilute H_2SO_4 it loses 1 mol. of H_2O , sobrerone—identical with Wallach's pinol—being formed.

Eucalyptol C₁₀H₁₀O (176°) (Jahns, B. 17, 2941; Bouchardat a. Voiry, C. R. 106, 663). Identical with cincol, spicol, cajeputol, and ter-

pan (B. a. V.).

Myristicol C₁₀H₁₀O (224°) (Wright, B. 6, 1320; Gladstone, C. J. 23, 147; 25, 1; Brühl, B. 21, 471). S.G. 22 9446 (G.). R_A 46·42 (G.). Myristicol is a dextrorotatory alcohol, obtained from the ethereal oil of nutmeg, Myristica aromatica. It yields a chloride with PCl₂, and is converted into cymene by the action of ZnCl₂.

Absinthol C₁₀H₁₆O (217°) (Beilstein a. Kupffer, B. 6, 1183; Wright, l.c. 1320; Gladstone, C. J. 45, 241; Brühl, B. 21, 471). S.G. ²²/₂₂ 9128 (G.). R_a 44·62 (G.). Absinthol is obtained from oil of wormwood, and is dextrorotatory. It is converted by ZnCl₂ and P₂S₄ into cymene. In what form the oxygen exists is at present unknown.

Constitution of the terpenes.

I. Pinene. The following facts must be taken into consideration:—

1. Dry pinene combines with one molecule of hydrogen chloride to form a saturated compound, from which hydrogen chloride is withdrawn with difficulty, the resulting hydrocarbon being a solid camphene. Moist pinene combines with 2HCl, yielding dipentene-dihydrochloride. This also is saturated. The formation of both these compounds is undoubtedly attended by isomeric change, inasmuch as pinene cannot be recovered from either of them; but on removal of the elements of hydrogen chloride a new hydrocarbon results.

2. Pinene seems to combine with two atoms or with four atoms of bromine, according to the method of operating. By adding bromine to a cooled solution of pinene in carbon tetrachloride, Wallach has obtained a crystalline dibromide C₁₀H₁₆Br₂ [170°]. The yield is, however, very small, about 7 p.c. of the hydrocarbon employed (A. 264, 1). By shaking up a solution of pinene in chloroform, with an excess of sodium hypobromite, acidified with hydrochloric acid, so as to liberate the bromine, and immediately afterwards determining the unabsorbed excess of bromine, Tilden obtained results which pointed to the union of the terpene with four atoms of bromine. The combination is, however, very unstable, and hydrogen bromide is soon evolved C. J. 53, 882). Schtschukaroff finds that a chloroform solution of pinene takes up, in the dark, four atoms of bromine; but the bromide formed is unstable, and quickly gives off hydrogen bromide, thus C₁₀H₁₀Br₄ = HBr + C₁₀H₁₅Br₅ (B. 23, 432, Ref.; J. pr. 47, 191-6).

3. Pinene combines with NOCl to form a saturated compound which is not an oxim but a nitroso-compound, and yields up the NO.Cl by the action of aniline, with liberation of an

optically inactive pinene.

4. By addition of two atoms of bromine to pinene, and subsequently heating the product

(Otto a.

alone, or, better, with aniline (Oppenheim, B. 5,

629), cymene is formed.

5. In contact with slightly diluted mineral acids, or when heated above its boiling-point, pinene is gradually converted into an optically inactive mixture of dipentene with terpinene, terpinolene, and polymerised terpene ('colophene').

6. When heated to a temperature short of

6. When heated to a temperature short of visible redness, pinene yields a considerable quantity of isoprene C₅H_s, together with a

notable amount of meta-xylene.

7. Pinene submitted to oxidation yields about 1 p.c. of its weight of p-toluic or terephthalic acid. This is perhaps due to the presence of a little cymene. It must, however, be remembered that the oxidising agent usually employed is either nitric acid or acidified chromic liquor, and hence that the pinene is first changed into dipentene or one of the other hydrocarbons referred to above (5). When nitric acid is used oxalic acid is the most abundant product, accom
(CH₃)_xC.OH(COOH).CH₂

and dimethylfumaric acid || CH.C.COOH

Beckurts, B. 18, 826). By treatment with chromate and sulphuric acid pinene yields much acetic acid, together with some terebic acid and

terpenylic acid
$$C_2H$$
,.CH.CH(COOH).CH₂ C_2H ,.CH.CH(COOH).CH₂ C_2H

8. Natural pinene rotates the plane of polarisation to the right or to the left. It therefore must be supposed to contain an asymmetric atom of carbon.

In order to epitomise these facts many formulæ have been proposed. The most important fall under two classes—namely, those in which pinene is represented as a dihydrocymene, and those in which a cross- or para-linkage is assumed.

Of the former class, the first example is the formula given many years ago by Oppenheim (B. 5, 98)—

This kind of formula is now attributed with greater probability to limonene, and it must be remarked that of the possible formulæ of this type there are only four which contain an asymmetric carbon atom and iso-propyl.

It has been shown that cymene, with which pinene is closely connected, contains iso- and not normal propyl (Widman, B. 24, 439). Hence pinene and its isomerides are believed also to be isopropyl compounds.

Formulæ containing cross- or para- linkages

have been proposed as follows:-

II. Limonene.—1. Dry limonene combines with one molecule of hydrogen chloride, but unlike the hydrochloride derived from pinene the resulting compound is optically active and is not saturated, as it combines with halogens, with nitrosyl chloride, and, in the presence of moisture, with a second molecule of hydrogen chloride (Wallach, A. 270, 188). Limonene in the presence of water or alcohol readily yields dipentene dihydrochloride [50°], identical with the product formed from pinene. The dihydrochloride is optically inactive.

2. Limonene unites with four atoms of bromine, forming a crystalline tetrabromide.

 Limonene unites with one molecule of nitrosyl chloride. The product, of which two (stereo?) isomerides occur, gives the reactions of an oxim.

4. Limonene readily yields abundance of cymene identical with that which is obtained from pinene (Oppenheim, B. 5, 628).

5. Limonene is polymerised by heat much less readily than pinene, and is not convertible into camphene.

6. Limonene vapour at a low red heat yields

isoprene in the same way as pinene.

7. Limonene oxidised by nitric acid or by chromate yields oxalic, acetic, terebic or terpenylic acids according to circumstances, usually without any trace of toluic or terephthalic acid.

8. Limonene is dextro- or lavo-rotatory.

mixture of equal volumes of the active hydrocarbons constitutes dipentene, which, on account of its peculiarities, quite different from those of its components, was long regarded as a distinct hydrocarbon.

The formula which agrees best with the characters of limonene is either I. or II. given

above (see Pinene).

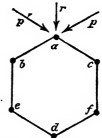
In both of these there is an asymmetric carbon atom, and as the mono-hydrochloride is optically active while the dihydrochloride is inactive, these compounds will probably be represented by the following formulæ:—

Limonene nitrosochloride deprived of the elements of hydrogen chloride yields a compound which has been identified with carvoxim. If the following formulæ be assumed for carvacrol and carvol, the interrelation of these compounds is obvious.

The corresponding formula for limonene should be No. II., but if that correctly represents the constitution of the hydrocarbon, the production of the nitrosochloride, or of the oxim from it, must be attended by isomeric change involving a re-distribution of the hydrogen and of the double linkages. Thus:—

The oxim is optically active.

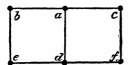
The very ready conversion of pinene into dipentene (= limonene) is a point of considerable importance, and it appears probable that the products formed by the action of heat and acid oxidising agents are in reality derived, not directly from pinene, but from the isomeric hydrocarbon into which it is first converted. The splitting up of pinene by heat into isoprene has been regarded (Wallach, A. 239, 48; Collie, B. 25, 1111) as supporting the formulæ already given VI. and VII., in which a para- or cross linkage is adopted. But remembering that isoprene is converted by polymerisation, not back again into pinene, but into dipentene, this evidence is of little importance. On the other hand, the difficulties involved in the assumption of a cross link in a cycloid of six carbon atoms are very considerable. In any closed ring of carbon atoms, the stability of the ring is easily accounted for upon simple mechanical principles. For if we consider six carbon atoms situated at the angles of a regular hexagon, and each attracted to its two neighbours on either side, whatever be the forces of attraction between them the resultant of those forces will have the effect of urging each carbon atom toward the centre of the figure. Thus, if the carbon atom a is attracted towards the carbon atom b with a force represented by p, and towards c with a force represented by p', in the figure, the resultant of these two pressures will be r, which acts towards the centre; and so with each of the five other atoms.



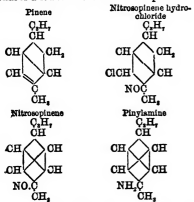
This conception is, of course, independent of any hypothesis concerning the fourth unit of valency, and the formula has no reference to the well-known 'centric' formula. The modern hypothesis according to which the carbon atom is supposed to be situated at the centre of a regular tetrahedron while the radicles with which it is united are placed at the angles of the same supplies no distinction between the force and the direction of the unit of affinity. The difference of specific volume observed between carbon compounds of the aromatic and fatty series seems to indicate that carbon atoms in the former are more closely united together, but it appears that no compounds are known in

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which the specific volume of carbon differs very greatly from its normal value. Hence it appears probable that the carbon atom, in uniting with other atoms, preserves a nearly constant distance from them, and in the figure above to suppose that a is combined with d as firmly as with b or c requires the assumption that a carbon atom is capable of acting through a distance twice as great in one case as in another. If this is possible, and we imagine the force of attraction between a and d to be as great as between a and b or c, then, unless c acts upon e and b acts upon f at the same time with the same force the figure will be changed from the hexagon to a double square.



A compound of this kind would be incapable of many of the changes which are characteristic of pinene, and a second cross linkage between c and e, or between b and f, would be impossible. Hence the following formulæ, which are attributed by Wallach to nitrosopinene and pinylamine cannot be accepted even if a single diagonal is allowed in the formula of pinene.



Formulæ III. and IV. now remain to be considered. So far as the writer is aware, formulæ of this type have not been proposed for any of the terpenes. Nevertheless there is much to be said in favour of an expression of this kind for a compound so unstable as pinene, which appears to be incapable of withstanding the action of heat or of contact with any reagent without isomeric change. For example, the conversion of pinene inte dipentene, terpinene, and terpinolene may be explained with the minimum amount of hypothesis by the following formulæ, in which the tendency towards the conversion of an unsymmetrical into a symmetrical molecule is clearly shown. The question as to the relative positions of the methyl and propyl groups is left open.

The conversion of pinene into dipentene through the dihydrochloride is readily shown.

Such a formula for pinene also explains the extreme instability of its tetrabromide.

III. CAMPHENE. Camphene behaves towards all reagents-such as bromine and nitrosyl chloride—as a saturated compound, but with hydrogen chloride it unites, forming a compound which in external appearance resembles pinene monochloride, but differs from that compound by its instability. When camphene is oxidised by chromic acid it yields camphor, but when treated with permanganate a glycol is formed. This compound crystallises in odourless small monoclinic prisms, which sublime at temperatures above 100° (192° ca.). Its most characteristic property is seen in its reaction with dilute acids, whereby water is separated and a compound isomeric with camphor formed. This reaction appears to indicate the existence of one ethylenic bond in camphene (G. Wagner, B. 23, 2307). Brühl (B. 25, 160) also finds, as the result of his latest determinations of the molecular refraction and dispersion of camphene, that this hydrocarbon contains one ethylenic linkage. Wallach (B. 24, 1555) upon other grounds appears to hold the same view. Camphene is, however, so closely connected with camphor that the constitution of the latter compound when settled may serve as a guide in determining the constitution of the former. The formula for camphor given by Kekule (B. 6, 931) was based upon the supposed intimacy of its relation to cymene and carvacrol.

There can be no serious doubt of its character as a ketone, inasmuch as it yields an oxim, though it gives no compound with NaHSO₂; but the action of dehydrating agents upon camphor is very complicated, and there can be no doubt that the production of cymene is attended by isomeric change. Armstrong a. Kipping (C. J., Jan. 1893) have shown that among the products of the action of strong sulphuric acid upon camphor

there is a considerable quantity of acetylorthoxylene (C₆H₂Me.Mc.Ac, Me:Mc:Ac = 1:2:4). From observation of its molecular refraction Brühl (B. 21, 467) suggests a formula containing a para- linkage, either

$$\begin{array}{c|cccc} C_{a}H, & & & C_{3}H, \\ C & & & CH_{a} & & CH_{a} \\ CH_{a} & & CH_{a} & & CH_{a} \\ \hline CH_{a} & & CH_{a} & & CH_{a} \\ \hline \\ CH_{a} & & CH_{a} & & CH_{a} \\ \hline \end{array}$$

The former of these two formulæ has also been used by Wallach (B. 24, 1555). These expressions are equivalent to representing camphor as a derivative of tetramethylene.

A formula of a different character has been suggested by Armstrong a. Miller (B. 16, 2260) (v. vol. i. p. 670), and another of somewhat similar type by Collie (B. 25, 1114):

Camphor by oxidation with nitric acid yields camphoric acid in large quantity, and as the characteristic and chief product. Camphoric acid is therefore probably derived from camphor by an operation of a simple kind without constitutional disturbance. The following formula proposed by Collie (B. 25, 1116) has been shown to possess a very high degree of probability (Walker, C. J., April 1893):

This formula sufficiently accounts for the chief facts concerning camphoric acid, viz. that it is a saturated dibasic acid, which readily yields the anhydride by heating; that it is optically active, and when treated with hydrogen iodide yields tetrahydro- and hexahydro- we when the contract of the co

From this formula are deducible the following formulæ for camphor and camphene:

This formula for camphene explains pretty well all its relations, except its derivation from pinene, if either of the formulæ containing a para-linkage be adopted. The chief objections to such formulæ are stated above.

J. Bredt (B. 26, 3047) has recently proposed formulæ for camphene, camphor, and camphoric acid, which are derived from a pentamethylene ring.

Synthesis of Terpenes.

Von Baeyer in a series of papers has shown that various dihydro-derivatives of benzene and its homologues may be synthetically produced. Starting with succino-succinic ester, this compound, by treatment with dilute sulphuric acid, yields diketo-hexamethylene, and this, by reduction with sodium-amalgam, gives the corresponding glycol. This glycol possesses the characters of a sugar, and has been named chinite or quinite (B. 25, 1037). It occurs in two isomeric forms, which by treatment with hydrogen bromide yield two dibromohexamethylenes. Both of these by heating with quinoline yield dihydrobenzene (B. 25, 1840), a liquid which has nearly the same boiling-point as benzene, but combines with HBr and with Br. It also reduces permanganate. By a corresponding series of processes dihydro-p-xylene was obtained from dimethylsuccino-succinic ester (B. 25, 2122). Dihydrop-xylene (134° circa) smells of turpentine, unites with hydrogen bromide, but forms no crystallisable nitrosite.

In order to produce a dihydrocymene it is necessary to start from the methyl-propyl com-This compound of succino-succinic ester. pound was obtained by heating the sodium compound of the monopropyl ether with methyl iodide (B. 26, 233). Methyl-isopropyl-dihydrobenzene boils at about 174°. It smells like turpentine and resinifies in the air. It decolourises permanganate and unites with bromine, forming an uncrystallisable bromide. Although closely resembling a terpene, however, it is obvious that this hydrocarbon is not identical with any natural terpene known, containing, as it does, no asymmetric carbon. It may be identical with terpinene. The relations of the paradibromohexamethylene, or of its alkyl derivatives, to the corresponding hydrocarbon is shown in the following formulæ (B. 25, 1840, 2122):

TERPENYLIC ACID C₂H₁₂O₄. [72°] [Fittig); [58°] (Wallach). Formed, together with terebic acid, by oxidising oil of turpentine and other terpenes with chromic acid mixture (Hempel, B. 8, 357; A. 180, 77; Fittig a. Krafit, A. 208, 71). Formed also by oxidising pinol hydrate [181°] with KMnO₄ (Wallach, A. 259, 318). Monoclinic prisms (containing aq).

Beactions .- 1. Yields acetic acid and CO, on

further oxidation by chromic acid.—2. On dry distillation it yields teracrylic acid (218°), a very little oxy-heptoic lactone (212°) and other bodies (Amthor, J. pr. [2] 42, 388).—8. Reduced by HI and P to \$\beta\$-isopropyl-glutaric acid

(Schryver, C. J. 63, 1344).

Salts.—BaA's. Amorphous powder, v. e. sol. water. Converted by boiling with baryta into barium diaterpenylate Ba(C,H18O,), aq, which separates in needles on heating the cold saturated solution and is v. sl. sol. water. On adding an acid the liberated diaterpenylic acid at once changes to its lactone, terpenylic acid.-CuA', xaq.-AgA'. Crystalline mass. Silver diaterpenylate AgC_sH₁₅O₅ is ppd. by adding AgNO₅ to a solution of barium diaterpenylate.

Ethyl ether EtA'. [38°]. (305° i.V.). Monoclinic crystals; a:b:c=1.051:1:.381; $\beta = 50^{\circ}$ 58'. V. sol. alcohol and ether, sl. sol. cold water. Sodium converts it into (a)- and

cold water. Southern Coll. H₂₂O₅. (β)-diterpodilactone C₁₅H₂₂O₅. [154°]. Formed as above (Fittig, A. 256, 113). Long Years, m. sol. alcohol. Yields needles (from water), m. sol. alcohol. Yields $CaC_{1a}H_{24}O$, 6aq, $Ag_2C_{1a}H_{24}O$, and $BaC_{1a}H_{24}O$, 6aqcrystallising in monoclinic prisms; a:b:c= $\cdot 935:1:\cdot 595$; $\beta = 65^{\circ} 54'$, which, when decomposed by HCl and immediately extracted with ether yields (a)-diterpolactonic acid $C_{15}H_{24}O_{8}$ [160°]. This acid decomposes above 160° into water and its lactone.

(β).Diterpodilactone C₁₅H₂₂O₅. ്135°ി. product of the action of Na on terpenylic ether (Fittig, A. 256, 122). Needles, m. sol. alcohol. Converted by baryta into (B)-diterpolactonic acid [187°] and (B)-diterpoxylic acid C15H25O7. (B)-diterpolactonic acid forms trimetric crystals; a:b:c = .782:1: 259 and is re-converted by heat

into the lactone. (3)-Diterpoxylic acid forms the salts BaA" 3½aq, CaA", and Ag,A".

(a)-Diterpylic acid C_{1a}H₂₂O₃. (a)-Diterpodilactone carboxylic acid. [216°]. An intermediate product of the action of Na on terpenylic ether. Needles, m. sol. alcohol. Converted by boiling HClAq or by heating above 216° into

(a)-diterpodilactone.

TERPILENE v. TERPENES.

TERPILENOL $C_{10}H_{18}O$. [c. 32°]. (c. 130° at 50 mm.). S.G.c. 95. H.C. (inactive) 113,318; (active) 102,923 (Louguinine, C. R. 107, 1165). Got by saponification of the terpilene acetates C₁₀H₁₆HOAc or terpilene formates C₁₀H₁₆CH₂O₂ by alcoholic potash (Lafont, A. Ch. [6] 15, 205). Crystals, insol. water, v. sol. oil of turpentine, alcohol, and ether. Boils with decomposition at 220°. Occurs in several forms varying in their action on polarised light.

Reactions.—1. HOl forms C10H162HCl [480], which is inactive even if the terpilenol were active.—2. Alcohol and HNO₃ give $C_{10}H_{22}O_3$ [104°].—3. Phthalic anhydride forms terpilene C10H10.-4. Ac2O at 100° forms the acetyl de-

rivative or terpilene acetate.

Acetylderivative C₁₀H_MOAc (220°). S.G. 2 .971. H₁₆HOAc. V.D. 6.6. Obtained by heating terpilene with HOAc at 150°. Liquid (Bouchardat a. Lafont, C. R. 102, 1555).

Formyl derivative $C_{1e}H_{17}O.CHO$. [a]_D = -69° 25'. S.G. 2 .999. Formed from oil of turpentine and cold formic acid (Lafont, C. R. 106, 140). By heating with water at 100° 65 p.c. of the ether is saponified.

TERPIN v. TERPENES. TERPINEOL v. TERPENES.

TERPYLONIC ACID $C_9H_{14}O_6$. [185°]. product of the oxidation of turpentine by chromic acid mixture (Schryver, C. J. 63, 1328). Needles, v. e. sol. water, m. sol. ether. Differs from camphoronic acid in electrical conductivity.

TETANINE v. PTOMAÏNES.

TETRADECANAPHTHENE C14H28. cor.). S.G. 2 ·839. Occurs in petroleum from Baku (Markownikoff a. Ogloblin, J. R. 15, 339).

n-TETRADECANE C_{1.}H_{so.} Diheptyl. [4·5⁶]. (253°). S.G. § ·775; § ·768. Formed by heatmyristic acid with HI and P (Krafft, B. 15. 1700). Got also by the action of Na on n-heptyl iodide (Krafft, B. 19, 2223; Sorabji, C. J. 47,

Reference. - DI-BROMO-TETRADECANE.

TETRADECENOIC ACID C14H26O2 C_6H_{13} .CH: $C(C_5H_{11})$.CO₂H. Amylhexylacrylic acid. Formed by the action of alcoholic potash on cenanthol (Perkin, B. 15, 2803; 16, 211). Liquid, boiling in vacuo at 275° to 280°.

TETRADECENOIC ALDEHYDE C_{1.}H_{2.0}O i.e. CH₃[CH₂]₃CH:C(CHO)[CH₂]₄CH₃. (278°). S.G. $\frac{15}{5}$ 849. Formed by the action of alcoholic potash or of ZnCl2 on cenanthol (Perkin, C. J. 43, 47). Oil, not solid at -20° . Forms crystalline C₁₄H₁₈ONaHSO₃. and heptoic acids on oxidation. Yields hexoic

TETRADECENYL ALCOHOL C,4H28O, (282°). S.G. $\frac{15}{15}$ ·852. Formed by reducing the preceding aldehyde with sodium-amalgam (Perkin, B. 15, 2808). It is also a product of the action of sodium-amalgam on a solution of cenanthol in HOAc (Perkin, C. J. 43, 68). Oil. May be reduced to $C_{14}H_{30}O$.

Acetyl derivative C14H27OAc. 290°). S.G. 15 868. Combines with Br (2

atoms).

TETRADECINENE C14H26 i.e. CMe:C.C11H29. [6.5°]. (134° at 15 mm.). S.G. $\frac{6.5}{4}$ ·8064; $\frac{15.2}{4}$ ·8000. Formed from tetradecylene bromide and alcoholic potash at 150° (Krafft, B. 17, 1872; 25, 2249).

TETRADECOIC ACID $C_{11}H_{28}O_{2}$ i.e. $C_{3}H_{11}.CH(C_{7}H_{13}).CO_{2}H.$ (300°-310°). Got by the action of moist Ag₂O on the corresponding aldehyde (Perkin, C. J. 43, 74). Liquid, not solid at -10° .

Tetradecoic acid v. MYRISTIC ACID. Reference.—OXYTETRADECOIC ACID.

TETRADECOIC ALDEHYDE C1, H20 i.e. C₃H₁₁.CH(C₇H₁₃).OHO. [80°]. (267°). S.G. 30 827. A product of the action of Na on an ethereal solution of cenanthol (Perkin, C. J. 43, Tables, v. sol. alcohol. Reduces ammoniacal AgNO. Yields heptoic, hexoic, and acetic acids on oxidation.

Isomeride v. MYRISTIC ALDEHYDE.

n-TEFRADECYL ALCOHOL C,4H_{s0}O. [38°]. (167° at 15 mm.). S.G. 30° 8153. Formed by reducing myristic aldehyde (Krafft, B. 16, 1720). [13°]. Acetyl derivative C1. H2. OAc. (176° at 15 mm.).

Tetradecyl alcohol C,H,1.CH(C,H,s).CH,OH. (270°-275°). S.G. 15 8368. Formed by reducing the corresponding aldehyde or the aldehyde C₁₄H₂₆O (Perkin, B. 15, 2811; C. J. 43, 76). Liquid, solidifying at -10° .

Acetyl derivative C14H29OAc. (c. 278°).

S.G. $\frac{15}{15}$ ·850. Oil, not solid at -10°. n-TETRADECYLENE C14H28 i.e.

CH₂:CH.C₁₂H₂₅. [-12°]. (127° at 15 mm.) S.G. $\frac{9}{4}$ 7852; $\frac{15}{4}$ 7745. Formed by distilling tetradecyl palmitate at 500 mm. (Krafft, B. 16, 3021). Liquid.

Bromide v. DI-BROMO-TETRADECANE.

TETRADECYLENE GLYCOL v. Di-iso-BUTYL PINACONE

TETRADECYL-MALONIC ACID C17H32O2 i.e. $C_{14}H_{29}CH(CO_2H)_2$. [118°]. Formed boiling its nitrile with alcoholic potash (Hell a. Jordanoff, B. 24, 988). White powder (from HOAc), insol. water, v. sl. sol. ether, sl. sol. cold alcohol. At 150°-170° it is split up into CO₂ and palmitic acid.—Ag₂A'': white pp.

Amide C₁₄H₂₉.CH(CO,H)(CONH₂). Pearly scales. At c. 140° it yields palmitic amide. Nitrile C₁₄H₂₉.CH(CO,H).CN. [76°].

Formed by heating p-bromo-palmitic ether with alcoholic KCy. Silky plates (from HOAc).

n-TEIRA-ICOSANE C₂₄H₅₀. [51°]. (243° at 15 mm.). S.G. 79 .763; 99 .748. Formed by the action of HI and P on C₁,H₃₀,CCl₂,C₆H₁, made from the ketone obtained by distilling barium stearate with barium heptoate (Krafft, B. 15, 1718).

TETRA-ICOSOIC ACID C23H47.CO2H. [72.5°]. Occurs in the soap got by heating carnaüba wax with aqueous NaOH (Starcke, A. 223, 307). Crystalline powder, v. sol. hot alcohol.—PbA'2. [111°]. Sol. toluene and hot HOAc.

TETRAZOLE $N \leqslant_{CH.\dot{N}H}^{N=N}$. [155°]. Formed by treating amido-phenyl-tetrazole carboxylic acid with alkaline KMnO, followed by HNO, (Bladin, B. 25, 1412). Plates (from toluene). Reddens blue litmus.

TETRAZYL-HYDRAZINENH, NH. $C \leqslant_{NH.\dot{N}}^{N-\dot{N}}$ [199°]. Formed from amido-tetrazotic acid by diazotisation and reduction by SnCl, (Thiele, A. 273, 155). Yellowish crystalline aggregates, v. sol. hot water.—B"2HCl. [176°]. Aceto-acetic ether forms $\stackrel{CH}{CMe.NH}$ N.C $\stackrel{N-N}{NH.N}$ [215°] crystallising in needles.

TETRIC ACID C10 H12O. i.e.

CH₂:C(OH).CMe < 0 .CO > CMe.C(OH):CH₂

(Nef, A. 266, 92) or CH₂CO.C(CO₂H):CH₂ Acetyl-acrylic acid. [189°]. (262°). S. 1·5 at 14°. Electrical conductivity: Walden, B. 24, 2027. Got by heating CH₃CO.CMeBr.CO₂Lt at 100°, but not by heating the isomeric acid CH₂Br.CO.CHMe.CO₂Et (Nef; cf. Pawloff, B. 16, 486). Formed by successive action of bromine and alcoholic potash on methyl-acetoacetic ether (Demarcay, C. R. 87, 351; 88, 126). Triclinic prisms (from water), v. sol. hot water, alcohol, and ether. Gives a violet-red colour with FeCl, PCl, gives rise to several compounds boiling between 169° and 185° which take up chlorine forming C.H.Cl.O (Pawloff, Bl. [2] 45, 181; C. R. 97, 99). Demarcay obtained oily C,H,Cl,O (?) which united with Cl and Br forming C,H,Cl,O [48°] and C,H,Cl,Br,O [67°]. Resinified by dilute HClAq at 150°. Potash-Vot. IV.

fusion gives formic and propionic acids. Distillation with lime forms methyl ethyl ketone Sodium-amalgam has no action.

Salts.—(NH₄)₂A". Crystals, v. sol. water.
—Na₂A" 6aq.—K₂A".—BaA" 3aq. Needles, v. e. sol. water.—BaA" 2aq.—Ba₂(OH)₂A". Sl. sol. water.—CaA" 2aq.—MgA" 10aq.—ZnA" aq.—CuA".—Ag₂A". Needles, sl. sol. water.

Ethyl ether CMe:C(OEt) O (?). 180°1.

(180° at 70 mm.) (Moscheles, B. 21, 2607).

Amide C,H,NO₂. [212°]. Got by heating

the acid with alcoholic NH,

TETROLE CYANURAMIDE v. Nitrile of PYRROLE CARBOXYLIC ACID.

TETROLIC ACID C.H.O. i.e. CH. C:C.CO.H. Butyrolic acid. Mol. w. 84. [77°]. (203°). Formed by boiling \$\beta\$-chloro-crotonic acid or \$\beta\$chloro-iso-crotonic acid with dilute potash (Geuther, Z. 1871, 245; Friedrich, A. 219, 343; Kahlbaum, B. 12, 2338). Formed also by the action of Na and CO2 on allylene chloride (Pinner, B. 14, 1081).

Preparation.—From acetoacetic ether and PCl_s, the product being freed from POCl_s by heating to 115, mixed with water, and freed from chloro-crotonic acid by distillation with steam. The residue is treated with alcoholic potash (Fittig a. Clutterbuck, A. 268, 96; cf. Friedrich, A. 219, 322).

Properties.—Plates (from ligroin). May be sublimed. V. e. sol. water, alcohol, and ether. Not reduced by sodium-amalgam to crotonic acid (F. a. C.; cf. Aronstein, B. 22, 1181).

Slightly volatile with steam.

Reactions. - 1. Br forms di-bromo-crotonic acid [120°], which is reduced by sodium-amalgam to tetrolic acid. Bromine also yields C4H4Br2O2. [97°] (Pinner).—2. HBr forms bromo-crotonic acid [91°] .- 8. Heated with conc. KOHAq it forms acetone.-4. Fuming HClAq at 15° in a few weeks converts it into β -chloro-crotonic acid [94.5°].—5. Boiling with Na and MeOH yields butyric acid (A.).—6. Chlorine forms C₄H₄Cl₄O₂. 7. On heating with iodine in CHCl, in sealed tubes it yields di-iodo-crotonic acid (B. 26, 843).-8. Split up at 211° into CO₂ and allylene.—9. KMnO, forms acetic acid.

Salts.—NaA'. Crystalline.—KA' (dried at 100°). Prisms (from alcohol). — NH,A'. LiA'aq. -MgA', 3aq. - BaA', 3aq. - CaA', 3aq. Small needles.— ZnA', 2aq (Lagermark, J. R. 12, 290).— ZnA', 2aq (Fittig). Groups of prisms.—CdA', 4aq.—PbA', aq. S. 1.5 in 92 p.c. alcohol at 20°.

Chloride C4H3OCl. Fuming liquid, carbonised by heat. Quickly decomposed by water (Lagermark, Bl. [2] 35, 171).

DI-TETROLURE A v. CARBONYL-PYRROL. TETROLE-URETHANE v. Ethyl ethyl of PYRBOLE CARBOXYLIC ACID.

TEUCRIN C₂₁H₂O₁₁ or C₂₁H₂₆O₁₁. [230°]. A glucoside occurring in *Teucrium fructicans*, a plant used in Sicily as febrifuge. Extracted by alcohol (Oglialoro, G. 9, 440; 13, 498). Slender yellow prisms. Boiling dilute nitric acid forms anisic, tartaric, and oxalic acids. Boiling dilute H₂SO, yields glucose and a yellow insoluble residue.

TEWFIKOSE. $[a]_p = 53.1^\circ$. A sugar occurring in the milk of the Egyptian buffalo (Bos bubalus). Yields glucose on hydrolysis (Pappel a. Richmond, C. J. 57, 759), v. Sugars.

THALICTRINE. Macrocarpine. Occurs in Thalictrum macrocarpinm (Hanriot a. Doassans, Bl. [2] 34, 83). Yellow needles, insol. water, sol. alcohol and ether. Resembles aconitine, but is less poisonous.

THALLIN v. OXY-QUINOLINE.

THALLIUM. Tl. At. w. 203.64. Mol. w. 407.28 as gas at c. 1700° (v. infra). Melts at 293.9° (Crookes, C. J. 17, 123; cf. Lamy, A. Ch. [3] 67, 385). Boils at full red heat. S.G. pressed 11.88, cast 11.81, in wire 11.91 (Crookes, l.c.; cf. Lamy, l.c.; Werther, J. pr. 91, 385; de la Rive, C. R. 56, 588). V.D. 232-7 at 1636°, 203-7 at 1728° (Biltz a. V. Meyer, B. 22, 725). S.H. 0325 (Lamy, l.c.); 0335 (17°-100°) (Regnault, A. Ch. [3] 67, 437). C.E. (linear) 00003021 at 40°, .00003135 at 50° (Fizeau, C. R. 68, 1125). E.C. 5.225 at 0° (Hg at 0°=1) (Bénoit, P. M. [4] 45, 314; cf. Matthiessen, P. 118, 431). H.C. [Tl²,O] = 42,240 (Th. 3, 354). Latent heat of fusion c. 1040 for 203.6 g. (Heycock a. Neville, C. J. 65, 35). The characteristic green line of the emission spectrum has the wave-length 5347 according to Huggins (T. 1864. 139), 5349.6 according to Mascart (Ann. de l'écol. norm., 4, 7 [1867]), and 5349.5 according to Thalen (Determin. des long. d'onde des raies metalliques [Upsala, 1868]). Concerning the emission spectrum of Tl v. also Crookes, Pr. 12, 536; Nickles, C. R. 58, 132; Kirchoff a. Bunsen, A. 155, 230, 366; Lockyer a. Roberts, Pr. 23, 344; Liveing a. Dewar, Pr. 27, 132; Hartley a. Adeney, T. 1884. 104; Becquerel, C. R. 99, 376; Wilde, Pr. 53, 369.

Historical.—In 1861 Crookes was preparing Se from the flue-dust of a sulphuric acid manufactory at Tinkelrode in the Harz mountains, when on distilling the impure Se he obtained a residue which showed a marked single green line in the spectrum; this line was not known to belong to the spectrum of any element, hence Crookes supposed that the impure Se contained a new element. Crookes isolated the element towards the end of 1861, and called it thallium (from $\theta a \lambda \lambda \delta s = a$ green bud) (v. C. N. 3, 193, 303). Lamy (A. Ch. [3] 67, 385) noticed a new green line in the spectrum of a leaden chamber deposit about the same time as Crookes was investigating the cause of the line he had observed; in 1862 Lamy isolated a specimen of the metal which Crookes had obtained a few months before.

Occurrence. — Never uncombined. Compounds of Tl are widely distributed, but only in very small quantities. The sulphide occurs in minute quantities in many specimens of copper pyrites and iron pyrites (v. Crookes, C. J. 17, 112; Lamy, l.c.; Wöhler, A. 142, 263; Carstaljen, J. pr. 102, 65; Gunning, Ar. N. 3, 86); also in native sulphur from Lipari and Spain. Compounds of Tl are found in small quantities in many ores of As, Bi and Hg, and in preparations from these ores; also in commercial Se and Te, probably as selenide and telluride (v. Crookes, l.c.; Röppler, Am. S. [2] 35, 420; Herapath, Ph. 4, 302; Werther, J. pr. 89, 129). Some specimens of lepidolite contain small quantities of Tl compounds (v. Schrötter, W. A. B. 48 (ii.) 734; 50 (ii.) 268). A Swedish

mineral called crookesite was found by Nordenskjold to contain from 16·3 to 18·55 p.c. Tl as selenide, along with selenides of Cu and Ag (Bt. [2] 7, 409). Small quantities of TlCl are found in some mineral springs and in the saline deposits from such springs (v. Böttger, J. pr. 89, 378; 90, 22; 96, 294). Traces of Tl alum have been found accompanying the double sulphates of Cs, Li, and Rb in alumite from the island of Volcano (Cossa, Acad. dei Lincei [3] 2). Small quantities of Tl compounds have been noticed in carnallite, sylvine, and kainite (Schramm, A. 219, 374). Many specimens of commercial sulphuric and hydrochloric acids contain traces of Tl compounds (v. Crookes, l.c.).

Formation.—(Crookes, i.c.; or T. 1872.)—1. By reducing Tl₂CO₃ by fusion with KCN.—2. By electrolysis of Tl₂CO₄ covered with water.—3. By electrolysis of Tl₂SO₄.—4. By heating Tl₂C₂O₄ in a covered crucible.—5. By ppg. by zinc from solution of a thallous salt to which excess of NH₂Aq or NaOHAq has been added.

Preparation.-The most economical source is the flue-dust from sulphuric acid manufactories where pyrites or sulphur containing Tl compounds has been used. Many specimens of flue-dust contain no Tl compounds; in some specimens 8 p.c. Tl has been found, but only in a very few does the quantity amount to 25 p.c. Tl (Crookes). After having been heated to dull redness, to remove any HCl, the dust is well stirred in wooden tubs with its own weight of boiling water and allowed to settle for 12 hours; the liquid is siphoned off, and the residue is again stirred with boiling water and let settle; the mixed liquids are ppd. by a considerable excess of conc. HClAq; the pp., which is crude TICl, is well washed on a calico filter and squeezeddry. About 68 lbs. crude TlCl were thus obtained from 3 tons flue-dust (Crookes, T. 1872). The crude TlCl is heated with its own weight of pure conc. H₂SO₄ in a Pt dish till all HCl and most of the H₂SO₄ are removed; the TlHSO4 is dissolved in c. 20 times its weight of water, the solution is nearly neutralised by CaCO_s, filtered, ppd. by conc. HClAq, and the TICI so obtained is well washed, and is then boiled for a few minutes with (NH₃)₂SAq; the pp. of Tl₂S is filtered off, well washed with water containing H2S, and dissolved in dilute H,SO,Aq; the solution is boiled to remove H2S, ammonia is added in slight excess, and after boiling the solution is filtered and crystallised; (NH₄), SO₄, being much more soluble in water than Tl₂SO₄, remains in solution when Tl₂SO₄ crystallises out.

If small quantities of Tl are to be prepared from Tl₂SO₄, the salt is dissolved in c. 20 times its weight of water, the solution is acidulated with H SO₄, and electrolysed, Pt electrodes being used. If large quantities of Tl are to be prepared, Tl₂SO₄ is placed in a deep porcelain dish (holding about 4½ litres) and covered with water (c. 3 kilos Tl₂SO₄ are used at a time) and plates of pure zinc are arranged vertically around the sides of the vessel; heat is applied, and the whole of the Tl separates in a few hours as a spongy mass easily detached from the zinc by shaking. The spongy Tl is washed several times, pressed by the fingers, and then fused with KÖy in a porcelain crucible to get it into a

lump (for details v. Crookes, l.c.; v. also Gunning, J. pr. 105, 343).

Crookes (l.c.) gives several methods for preparing pure Tl. If the starting-point is commercial Tl₂SO₄, he recommends to dissolve this salt in water, to saturate the cold solution with pure H2S, filter, heat to boiling, and pour into boiling, dilute, pure HClAq; to filter while hot, and allow to cool; to wash the TlCl that crystallises out, by decantation, till the washings are free from H_2SO_4 , and to purify by re-crystallising twice, from water. The TlCl thus obtained is dried, mixed with pure Na₂CO₃, and projected, in small successive portions, into molten pure KCy in an unglazed porcelain crucible; reduction to metal is rapid. After cooling, the contents of the crucible are treated with water, the ingot of Tl is boiled for some time in water, dried, and fused over a spirit lamp (coal-gas should not be used on account of the S compounds in it) in an unglazed porcelain crucible, with free access of air and frequent stirring with a piece of unglazed porcelain, to facilitate oxidation, and the metal is poured into a porcelain mould.

Properties.—A white metal without any tinge of blue; when fused under coal-gas, liquid TI is exactly like Hg; may be highly polished by rubbing under water; very soft, may be cut with the nail, or soratched by lead. Less tenacious than lead; does not become brittle between —18° and its m.p.; very malleable, Lamy (A. Ch. [3] 67, 385) says that plates \(\frac{1}{16.000} \) mm. thick

can be obtained; can be squeezed, but not drawn, into wire; very inelastic. Tl crystallises very easily in octahedral forms; it crackles like tin when bent; a wire of Tl seems amorphous when prepared, but under water it soon becomes crystalline. Tl quickly tarnishes in air; the film of oxide formed protects the mass of metal. The metal may be kept unchanged for years under water that has been deprived of air (v. Böttger, D. P. J. 197, 374). Tī melts easily (at 294°) without first becoming pasty; it expands considerably before melting is complete, and con-tracts much on cooling. The boils below white heat; it may be distilled in a current of H. Pieces of the metal may be welded together by Tl is pressure at the ordinary temperature. nearly as diamagnetic as bismuth. Tl gives an intense green colour to a non-luminous flame (v. infra, Detection). Heated in air, Tl readily burns to Tl₂O; it combines with S, Se, P, Cl, Br, I, &c., when heated with these elements; it dissolves in solutions of most acids (v. Reactions). The compounds of Tl are very poisonous, producing symptoms like those of lead poisoning.

The at. w. of Tl has been determined (1) by ppg. Cl as AgCl from TlCl by AgNO, Aq (Lamy, A. Ch. [3] 67, 385 [1862]; Hebberling, A. 134, 11 [1865]); (2) by ppg. I from TlI by AgNO, (Werther, J. pr. 92, 136 [1864]); (3) by converting Tl₂SO, into BaSO, (Lamy, l.c.; Hebberling, l.c.); (4) by converting Tl into TlNO, (Crookes, T. 1873.277); (5) by determining S.H. of Tl (Lamy, l.c.; Regnault, A. Ch. [3] 67, 437 [1862]); (6) by determining V.D. of TlCl (Roscoe, Pr. 27, 426).

The mol. w. of Tl in the gaseous state is double the at. w.; in other words, the gaseous

molecule is diatomic (Biltz a. Meyer, B. 22, 725). Ramsay (C. J. 55, 521) determined the depression produced in the vapour pressure of Hg by dissolving Tl in that solvent; assuming that equal volumes of dilute solutions contain equal numbers of molecules, and that the mol. w. of liquid Hg is 200, the results obtained by R. indicated the molecule of Tl dissolved in Hg to be monatomic.

Tl is distinctly a metallic element; no compounds have been isolated wherein Tl certainly forms part of the negative radicle, although there are indications of the existence of such compounds (v. THALLIUM DIOXIDE, p. 680); the compounds which TIBr_s, TlCl_s, and TlI_s form with alkali halides may contain Tl in the negative radicles. Tl is the last element in the odd-series family of Group III.; this family comprises Al, Ga, In, and Tl. The element Tl also shows marked resemblances to the alkali metals; and in its physical properties it is more like lead than any other element. The position assigned to Tl by the periodic classification of the elements clearly indicates the relations between this element and other elements. The relations of Tl (III.-11) to Al (III.-3) should be similar to those of Hg (II.-11) to Mg (II.-3), and also similar to those between Pb (IV.-11) and Si (IV.-3). The resemblances between the compositions and properties of the oxides of Hg and Mg on the one hand, and of the oxides of Pb and Si on the other hand, are shown only in the highest oxides of these elements: there is Hg₂O but no Mg₂O, there is PbO but no SiO; HgO and MgO are basic, PbO, and SiO, are feebly acidic. Nevertheless Hg and Pb form lower oxides than HgO and PbO₂; these lower oxides, Hg₂O and PbO, are basic; Hg₂O feebly, and PbO strongly, basic. Hence it is likely that the resemblances between the oxides of Tl and Al will be shown in the highest oxides. This supposition is confirmed; Tl₂O₃ and Al₂O₃ are basic, forming salts M₂X₃, where X = the radicle of a dibasic acid; these oxides are more basic than PbO2 and SiO2, but less basic than HgO and MgO. As Hg and Pb form lower oxides than HgO and PbO2, and these lower oxides are more basic than HgO and PbO₂, so Tl will likely form an oxide lower than Tl2O2, and this oxide will be more basic than Tl2O3; Tl forms the strongly basic oxide Tl₂O. Then consider the position of Tl in series 11. Consider the oxides of the members of this series: the lower oxides Hg₂O, PbO, and Bi₂O₃ are basic; the higher oxides HgO, PbO₂, and Bi₂O₃ pass from basic (HgO) to feebly acidic (PbO2 and Bi2Ob); therefore it is probable that the higher oxide of Tl will be less basic than HgO, but that it will not be acidic; Tl2O3 is a feebly basic oxide.

If the chlorides of series 11 are considered, the position of Tl in III.—11 is seen to be justified: HgO gives HgCl₂, which can be gasified unchanged, PbO₂ gives PbCl₄, which is very easily decomposed by heat, no chloride corresponding with Bi₂O₅ has been isolated; therefore Tl₂O₅ will probably give a chloride more stable towards heat than PbCl₄, but less stable than HgCl₂. TlCl₂ is decomposed by heating to c. 80° into TlCl and Cl. Hg₂O gives HgCl, PbO gives PbCl₂, and Bi₂O₅ gives BiCl₂, and these chlorides can all be gasified unchanged; there-

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fore the chloride of Tl corresponding with Tl₂O will probably not be decomposed by heat; TlCl has been gasified unchanged. The relations between the elements Hg. Tl, Pb, Bi, and Mg, Al. Si may be summarised in the expressions Tl: Hg: Pb-Al: Mg: Si; and Tl: Hg=Pb: Tl=Bi: Pb. (For a further working out of the relations of Tl, v. Mendeléeff, C. N. 1880. 2; v. also Alkalis, metals of the, vol. i. p. 114; and Earths, metals of the, vol. ii. p. 424.) The atom of Tl is monovalent in the gaseous molecule TlCl; the V.D. of no other compound of Il has been determined.

Reactions and Combinations .- 1. Exposed to air or oxygen, a film of Tl2O is at once formed; when heated in air or O to somewhat below redness (c. 300°) Tl burns to Tl₂O₃, and at a higher temperature this gives Tl₂O and O (Lamy, A. Ch. [3] 67, 385). Moist ozone acts on Tl giving TlO₂H₃. -2. A compact piece of Tl does not decompose boiling water, decomposition begins at a red heat; but spongy Tl, as ppd. by zinc, dissolves in water forming TlOHAq.-3. Tl dissolves in solutions of most acids, forming thallous salts Tl_2X , where X = 2Cl, $2NO_3$, SO_4 , &c. -4. Tl combines with sulphur and selenion (v. sulphides and selenides).—5. Forms alloys with many metals (v. alloys).-6. Forms compounds with antimony and arsenic (v. antimonide and arsenide).-7. A phosphide is formed by heating Tl in vapour of phosphorus (v. phosphide).—8. Combines with chlorine, bromine, and iodine, when heated with these elements (v. chlorides, bromides, and iodides) .- 9. Tl ppts. the noble metals, also copper, lead, and mercury from solutions of salts of these metals (Reid, C. N. 12, 242).

Detection and Estimation.-Tl is best detected by the intense green colour which the metal and its compounds give to the non-luminous flame of the Bunsen lamp; according to Lamy (l.c.) $\frac{1}{50,000}$ mgm. can be detected by the use of the spectroscope; if induction sparks are used it is said that $\frac{1}{80,000,000}$ mgm. can be detected (Cappel, P. 139, 628). This generally estimated as TH or Th.SO.; it can also be determined volumetrically by KMnO, Aq (Willm, Z. 1863.

Several alloys of Tl Thallium, alloys of. were prepared by Carstanjen in 1867 (J. pr. 102, 82) by heating the metals together. Many of the alloys are soft, and most of them readily tarnish in the air; the alloy with \(\frac{1}{2} \) pt. Sb is fairly hard; and those with Bi and Pb, Bi and Cd, and Bi and Sn are hard and brittle. Alloys are described with Al, Sb, Cd, Cu, Pb, Mg (v. also Mellor, C. N. 19, 245), Hg (v. also Crookes, l.c., and Regnauld, C. R. 64, 60), K, Na, and Zn; also with Bi and Cd, Bi and Pb, and Bi and Sn.

Thallium, antimonide of. A hard, grey, crystalline solid was obtained by fusing together Tl and Sb in the ratio Tl : Sb; with cold dilute H,SO,Aq the substance gave off SbH2 (Carstan-

jen, l.c.).

Thallium, arsenide of. Carstanjen (l.c.) obtained a white, crystalline, soft substance by fusing Tl and As in the ratio Tl : As; with cold acids AsH, was formed.

Thallium, bromides of. The compounds TIBr and TiBr, have been isolated, and also two compounds the simplest formulæ of which are Tl₂Br, and TlBr, As none of the bromides has been gasified, the formulæ are not necessarily molecular; but from the analogy of TiCl, the formula TlBr probably represents the composition of the gaseous molecule of this compound.

THALLOUS BROMIDE TIBr. (Thallium monobromide.) Tl reacts very slowly with Br. Thallous bromide is obtained, as a white pp., by adding HBrAq or solution of a bromide to solution of a thallous salt; almost insoluble in TlBr melts at water (Willm, A. Ch. [4] 5, 28). 458° to a brownish-yellow liquid, which solidifies to a yellow solid (Carnelley, C. J. 33, 278). Thomsen gives H.F. [Tl,Br] = 41,290 (Th. 3, 354).

THALLIC BROMIDE TIBrs. (Thallium tribromide.) Prepared by adding Br to TlBr suspended in water until solution is effected, and evaporating in vacuo. TlBr, forms a yellow, crystalline solid; it is very deliquescent, becoming gradually brown and giving off Br. Very soluble in water, also in alcohol. Reduced to TIBr by SO₂Aq and other reducing agents (Willm, l.c.). Thomsen gives H.F. [Tl,Br',Aq] = 56,450 (Th. 3, 354). TlBr₃ combines with NH₃ to form TlBr_s.3NH_s; a white solid that gives off NH₃ and Br at 100° , and when more strongly heated leaves TlBr (W., l.c.). Also combines with NH₄Br and KBr, forming TlBr, NH, Brand TlBr, KBr. 2aq (W., l.c.; Nicklès, C. Ř. 58, 537). Rammelsberg (P. 146, 597) described 2TlBr, 3KBr. 3aq.

THALLIUM DIBROMIDE TIBr₂ (or TIBr.TIBr₃ = Tl₂Br₄). Obtained in long, yellow, needles by adding TIBr to boiling TIBr₃Aq, in the ratio TIBr:TIBr₃ and cooling. Not december ratio TlBr:TlBr, and cooling. Not decomposed at 100°; water forms TlBr₃Aq and Tl₂Br₃ (v. infra) (W., l.c.).

THALLIUM SESQUIBROMIDE Tl2Br3, or TlBr. 3TlBr = Tl, Br. (Thallo-thallic bromide.) Orange-red hexagonal leaflets; obtained by adding HBrAq to a mixture of a thallous and a thallic salt in solution; also by adding TiBr to TiBr,Aq, or by adding a little water to TiBr, (3TiBr, + Aq = Tl,Br, + TiBr,Aq) (W., l.o.).

Thallium, chlorides of. Ti and Cl combine

to form at least four compounds: TlCl, Tl,Cl,

TlCl, and TlCl,

THALLOUS CHLORIDE TICI. Mol. w. 239.01. Melts at 427° (Carnelley, C. J. 33, 277). Boils at 708°-719° (C. a. Williams, C. J. 33, 284). S.G. 7.02, after fusion (Lamy, J. 15, 184). at 830° to 1025°=117.8 (Roscoe, Pr. 27, 426). S. 19 at 0°, 27 at 16°, 28 at 16.5°, 1.6 at 63° (Hebberling, A. 134, 15). H.F. [Tl,Cl]=48,580; [Tl,Cl,Aq] = 38,480 (Th. 3, 354).

Formation.-1. By heating Tl in Cl, keeping the Tl in excess.—2. By boiling powdered Tl in HClAq, and dissolving the film of TlCl that forms in much boiling water.—3. By adding

SO2Aq to TlCl3Aq.

Preparation. - HClAq, or solution of a chloride, is added to the solution of a thallous salt; the white curdy pp. that forms is washed with cold water, dissolved in a large quantity of boiling water, and the salt that separates as the solution cools is purified by two or three recrystallisations from boiling water (Crookes, C. J. 17, 112).

Properties and Reactions .- A white, crystalline solid; resembling PbCl, but separating from solution in hot water in smaller and less shining crystals than PbCl₂. Becomes slowly

violet coloured when exposed to light (Hebberling, A. 134, 15). Melts at 427° to a clear yellow liquid, which solidifies to a white crystalline Very slightly soluble in water (v. supra); less soluble in water containing a little HCl; insoluble in alcohol; insoluble in ammonia; soluble in hot Na₂CO₃Aq, KOHAq, or Na₂SO₄Aq. Hot conc. sulphuric acid forms Tl₂SO₄ and HCl; oxidisers such as ClAq, aqua regia, KMnO,Aq and HClAq, form TlCl,, Tl₂Cl₃, or TlCl₂ (v. these chlorides). TICl is reduced to Tl by zinc and sulphuric acid, also by adding aluminium to TICIAq at 90° (Cossa, Acad. dei Lincei [3] 2). Fusion with sodium hydrogen sulphate produces Tl₂SO₄ (Krause, D. P. J. 217, 823; Nietzki, D. P. J. 219, 262).

Combinations.—1. With platinic chloride to form 2TlCl.PtCl₄ (= Tl₂PtCl₈); obtained, as a pale-yellow, crystalline powder, by mixing solutions of TlCl and PtCl₄; this salt is extremely insoluble in water, one part dissolving in 15,585 parts water at 15°, and in 1948 parts boiling water (Crookes, l.c.). - 2. With mercuric chloride, to form TlCl.HgCl₂; white lustrous needles, separating from a hot solution of TlCl in HgCl₂Aq; all HgCl₂ is removed at 200° (Carstanjen, J. pr. 102, 141; Jörgensen, J. pr. [2] 6, 82).—3. With ferric chloride to form 3TlCl.FeCl₃; small red prisms; obtained by adding TlCl to a hot solution of FeCl₃ in HClAq, and crystallising the red pp. from hot conc. HClAq; decomposed to its constituent salts by water (Wöhler, A. 144, 350).—4. With chromic chloride, to form 3TlCl.CrCl3 (v. Neumann, A. 244, 328).-5. According to Lepierre a. Lachaud (C. R. 113, 196), a conc. solution of chromium trioxide reacts with TlCl to form small red prisms of CrO₃.TlCl [CrO₂(OTl)Cl], which are decomposed by water to TlCl and CrO,Aq.

THALLIC CHLORIDE TICIs. Mol. w. not certainly known; from analogy of AlCl, GaCl, and InCl₃, the formula TlCl₃ is probably molecular.

Formation .- 1. By heating TICl with hot aqua regia and evaporating the solution in Cl (Willm, A. Ch. [4] 5, 28).—2. By dissolving freshly ppd. TlO, H, in HClAq, adding HNO, Aq, and evaporating in Cl.—3. By heating TlCl at its melting-point in Cl; it is doubtful whether the product is pure TlCl₂, or contains also TlCl₂ or Tl.Cl. (Lamy, l.c.).

Preparation.-Cl is passed into water containing TICl in suspension until a little of the solution ceases to give a turbidity with PtCl₄Aq; excess of Cl is removed by a stream of CO₂, and the solution is evaporated in vacuo. Crystals of TlCl_s.H₂O separate, which lose water at c. 40°

(Werther, J. pr. 91, 385).

Properties and Reactions. — The hydrate forms large colourless prisms; the dehydrated salt is a white crystalline solid. Werther (l.c.) also obtained a hydrate 2TlCl, 15H, O from a solution of TlCl, prepared as described. Cl begins to be given off at c. 50° to 60°; at 100° a mixture of TlCl and chlorides intermediate between TlCl and TlCl, remains (Werther, l.c.; Lamy, l.c.). TlCl, is very deliquescent; it dissolves very easily in water. Thomsen gives [Tl,Cl*,Aq] =89,250 (Th. 3, 354). On diluting a conc. aqueous solution considerable quantities of TiO_sH_s and HClAq are formed. Reducing agents, such as SO_2 , form TlCl.

Combinations.—1. With ammonia to form TlCl, 3NH, (Willm, A. Ch. [4] 5, 28). A white, crystalline powder. Obtained by passing NH, over TlCl_s; by adding an alcoholic solution of NH, to TlCl, suspended in absolute alcohol; or by boiling TlO,H, with conc. NH, ClAq, ppg. by NH,Aq, washing with alcoholic ammonia, then with absolute alcohol, and drying in vacuo. The compound is unchanged in air; forms NH₄ClAq and Tl₂O₃; when heated NH₂, NH₄Cl, and N are given off and TlCl remains. Dissolves in HClAq, forming TlCl₃.8NH₄Cl.— 2. With ammonium chloride to form TlCl3.3NH4Cl; obtained by adding NH4Cl to a solution of TlOsHs in HClAq, and evaporating in vacuo; large, white, quadratic crystals, probably with 2H₂O (Rammelsberg, P. 146, 597; v. also Nicklès, J. Ph. [4] 1, 25).—3. Compounds with potassium chloride, casuum chloride, and rubidium chloride, TlCl₃.3MCl. xaq, have been obtained (v. Rammelsberg, l.c., also W. 16, 709; Godeffroy, Zeit. des allgemein. österr. Apothekerver. 1880. No.9; Neumann, A. 244, 348; Willm, Neumann (l.c.) also describes a compound of TlCl, with beryllium chloride. -4. Willm (l.c.) describes a compound with cupric chloride, 2TlCl_s.CuCl_s.—5. According to Nickles (C. R. 58, 537) TlCl, forms a compound with ether and hydrochloric acid, to which he gives the composition TlCl₃.Et₂O.HCl.H₂O; it is described as a yellow fuming liquid.

THALLIUM DICHLORIDE TICL, (or TICI.TICL, $= Tl_2Cl_4$). Lamy (A. Ch. [3] 67, 402) obtained a yellow, slightly hydroscopic, easily fusible solid, to which he assigned the composition TlCl, by heating Tl and TlCl in a stream of Cl so that the product always remained liquid, and allow ing to cool when saturated; the substance gave off Cl, and Tl₂Cl₃ remained.

Tl₂Cl₃; THALLIUM SESQUICHLORIDE TlCl₃.3TlCl = Tl₁Cl₆. (Thallothallic chloride.)
Formed by heating dry TlCl₃ somewhat above 100° as long as Cl is given off; the brown liquid cools to a clear yellow, crystalline mass (Lamy, l.c., p. 403). Also formed by the direct combination of Cl with molten Tl (L., l.c.); also by ppg. a solution containing a thallous and a thallic salt by HClAq (Willm, l.c.). Also obtained by dissolving Tl or TlCl in aqua regia, evaporating, heating till Cl ceases to be given off, and crystallising from water slightly aciditied with aqua regia (Lamy, l.c.; Crookes a. Church, C. N. 8, 1). According to Werther (J. pr. 91, 390) the product of this reaction, dried at 100°, is a mixture of compounds of TiCl and TiCl, besides TiCl. TiCl. crystallises in yellow, hexagonal leaflets; unchanged in air; melting between 400° and 500° to a dark-brown liquid that solidifies with much contraction, forming a yellowish mass with S.G. 5-9 (Lamy, l.c.). Very slightly soluble in water; S. 26 at 15°, 29 at 17°, 1-9 at 100° (Crookes; Hebberling, A. 134, 21). The solution is said to show the reactions both of thallous and thallic compounds. Heated above 500° Cl is given off; heating in H2S produces Tl2S, S, and HCl (Hebberling, *l.c.*). Heating in NH, produces TlCl, NH, Cl, and N (Crookes a. Church, C. N. 8, Tl₂Cl₂ reacts with KOHAq to form TlO₂H₃, TiCl, and KCl.

OTHER THALLO-THALLIC CHLORIDES. ing to Werther (J. pr. 91, 890), two compounds

besides Tl₂Cl₂ are obtained by evaporating TlCl | in aqua regia, heating till Cl ceases to be given off, crystallising from water slightly acidified with aqua regia, and drying at 100°. To these compounds W. gives the formulæ 3TlCl.4TlCl.

Thellium, converged of; v. vol. ii. p. 347.

Thallium, ferrocyanide of; v. vol. ii. p. 347.

Thallium, fluorides of. Only one fluoride of Tl has been isolated with certainty; there are indications of the existence of another.

THALLOUS FLUORIDE TIF. Formula probably molecular, from analogy of TlCl. Prepared by dissolving Tl₂CO₃ in HFAq, evaporating to dryness, heating gently to remove HF, and crystallising from water; forms white, lustrous, regular octahedra (Büchner, J. pr. 96, 404). TIF is also said to be formed as a white sublimate by heating Tho in HF gas (Kuhlmann, C. R. 58, 1037). Sol. in water; S = 80 at 15° , and increases with increase of temperature; solution in water has alkaline reaction; somewhat sol. in alcohol. A hydrate TlF.H₂O forms in monoclinic crystals by evaporating a solution of Tl₂O or Tl₂CO₃ in HFAq; it is deliquescent, and acts on glass (Kuhlmann, l.c.; Willm, A. Ch. [4] 5, 5). A compound with fluorhydric acid, TIF.HF = TIHF₂, is obtained by evaporating a solution of TIF in HFAq over H₂SO₄. Decomposed above 100° to TIF and HF. V. sol. water; S. = 100 at ordinary temperature; the solution has an acid reaction (Büchner, l.c.).

By digesting TlO3H3 THALLIC FLUORIDE. with HFAq, and also by adding HFAq to Tl(NO₂)₂Aq, Willm (A. Ch. [4] 5, 5) obtained a dark olive-green solid, becoming brown when heated, and melting to an orange-yellow liquid that became white on cooling, insoluble in water and in cold HClAq. This substance was probably thallic fluoride, or perhaps a thallothallic fluoride; no analyses are given.

Thallium, hydride of. Herapath (Ph. 302) supposed that a compound of Tl and H existed; but Crookes (C. J. 17, 132) failed to obtain any compound either by the interaction of an acid with Tl-Zn alloy, or by heating Tl in a stream of H; the escaping H burnt with a green flame, but this was found to be due to particles of Tl carried by the stream of H.

Thallium, hydroxides of. Tl forms three compounds with O and H, namely, TlOH, TlO,H

(or TlO.OH), and TlO.H.

THALLOUS HYDROXIDE TIOH. (Hydrated thallous oxide Tl₂O.H₂O.) Mol. w. not known with certainty. Thomsen (Th. 3, 354) gives $[Tl,O,H] = 56,915; [Tl^2,O,H^2O] = 45,470;$ H.F. $[Tl^{2}O,H^{2}O] = 3,230.$

Formation.—1. By dissolving Tl in water containing air, and evaporating.—2. By decomposing Tl.SO.Aq by BaOAq, filtering, and evaporating.—8. By the interaction of CaOAq and Tl₂C₂O₄Aq.—4. By dissolving Tl₂O in water and

Preparation.—1. Tl ppd. by zinc is allowed to oxidise in warm, moist air, and the oxide is dissolved in boiling water; these operations are repeated till a saturated hot solution is obtained, and this solution is evaporated a little, filtered from any Tl₂CO₃ that may have separated, and allowed to crystalkise (Crookes, C. J. 17, 132).—2. Tl₂SO₄Aq is mixed with the proper quantity of BaOAq, and the solution, filtered from BaSO4, is crystallised in vacuo; this method is tedious, and there is much difficulty in getting rid of all BaO.—3. Thin plates of Tl are suspended over a shallow dish containing absolute alcohol, within a bell-jar from which air has been exhausted, and pure O is passed into the jar, the temperature being kept at 20°-25°; after several hours or days, C₂H₅TlO is formed as an oil in the dish; excess of alcohol is removed by heating to 100°; a little water is added, whereby TIOH is ppd.; the pp. is drained, and exposed to warm dry air till all water and alcohol are

removed (Lamy, A. Ch. [3] 67, 385).

Properties and Reactions. — Pale-yellow, prismatic crystals. At 100° in air, or in vacuo over H2SO4 at the ordinary temperature, the crystals become black from formation of Tl2O, without altering their crystalline form. Easily soluble in water; Thomsen (Th. 3, 354) gives [TlOH,Aq] = -3,155. The solution in water resembles KOHAq and NaOHAq in its properties; it is strongly alkaline, absorbs CO, ppts. heavy metals as hydroxides or oxides, feels

soapy to the touch, stains the skin, neutralises acids, &c. The solution absorbs a little O on

heating; when evaporated to dryness and the residue re-dissolved, a little Tl2Os generally remains.

THALLIC HYDROXIDES. Two thallic hydroxides are known, TlO.OH and TlO,H3.

I. Monohydrated thallic oxide, TlO OH = Tl2O3.H2O. Mol. w. not known with certainty.

Formation.—1. By ppg. TlCl,Aq by alkali and drying at 100° (Lamy, A. Ch. [3] 67, 385).— 2. By passing ozonised air through TlOHAq (Schönbein, J. pr. 93, 35).—3. By the action of H₂O₂Aq on Tl; according to Weltzien (A. 138, 129) a mixture of TIOH and TIO.OH is thus obtained.—4. Various oxidisers—e.g. KMnO,Aq, hypochlorite solutions, &c.—convert TIOH into TIO.OH (Schönbein, i.c.).—5. By electrolysing a neutral or ammoniacal solution of Tl₂(SO₄), or Tl(NO₃)₃; the hydroxide separates on the positive (Pt) electrode (Flemming, J. Z. 1869, 292).

Preparation. -TlCl3Aq is ppd. by a slight excess of KOHAq or NH, Aq, the pp. is thoroughly washed, and dried at 100° (Lamy, A. Ch. [3] 67, 397; Crookes, C. J. 17, 132; Werther, J. pr. 91, 388). The air-dried pp. has the composition TlO.OH (Carnelley a. Walker, C. J. 53, 88).

Properties and Reactions.—A brown powder; insoluble in water and dilute alkali solutions. Heated to c. 60° begins to lose water; completely dehydrated at 115°, according to Werther (J. pr. 91, 385), but the residue contains some Th.O. as well as Tl₂O₃ (cf. Birnbaum, A. 138, 133). According to Carnelley a. Walker (C. J. 53, 88), dehydration is complete at 230°, and the residue is Tl₂O₂, which is stable to 360°. Dissolves in HClAq, forming TlCl₂; in conc. acid Cl is given off and some TlCl is produced. Dissolves in dilute H₂SO₄Aq; with conc. acid some O is given off. Dissolves in SO₂Aq, As₂O₈Aq, hot tartaric acid solution, and boiling oxalic acid solution, in each case forming a salt of Tl₂O. TlO.OH dissolves in warm conc. NH,ClAq, giving off NH, and forming TlCl, 3NH, on adding water Tl2O, is

ppd. as a black powder (Willm, A. Ch. [4] 5, 5).

II. Trihydrated thallic oxide, TlO,H. Tl.O. 3H.O. Formed, as light-brown, lustrous.

microscopic, hexagonal plates, by dissolving TIO.OH in molten KOH, fusing for some time, cooling, washing with hot water, then with alcohol, and drying at 100°. Insoluble in water. Not decomposed at 340°; soluble in dilute acids (Carnegie, C. N. 60, 113).

Tha lium, iodides of. Tl combines directly with I. At least two compounds are known: TII and Tl₂I₃; TII₃, and probably one, or perhaps more than one, other compound of TII and TII,

seem also to exist.

THALLOUS IODIDE TIL. (Thallium monoiodide.) Mol. w. not known with certainty, but formula TII is probably molecular from analogy

Formation. — 1. By heating together Tl and I in the ratio Tl:I.—2. By adding KIAq to

solution of a thallous salt.

Preparation.-KIAq is added to a dilute solution of a thallous salt, and the yellow pp. is thoroughly washed with water and dried at 100° (Lamy, A. Ch. [3] 67, 405; Crookes, C. J. 17,

Properties and Reactions.-A citron-yellow powder; melts at 439° (Carnelley, C. J. 33, 278), and boils at 800°-806° (C. a. Williams, C. J. 33, 284). S.G. 7.072 at 15.5°, after fusion 7.0975 at 14.7° (Twitchell, Clarke's Table of Spec. Grav. [2nd edit.] 35). Very slightly soluble in water. Determinations of solubility vary considerably; Werther (J. pr. 92, 130) says that 1 pt. requires 20,000 pts. water to dissolve it at 15° , and Crookes (*l.c.*) gives solubility as 1 in 4450 of water at 15° (v. also Hebberling, A. 134, 16). Crookes says TII is somewhat more soluble, Hebberling says it is decidedly less soluble, in KIAq than in water.

When heated to c. 190° TII becomes red, and when it melts a red liquid is formed which solidifies to a red crystalline mass having the composition TII (Hebberling, L.c., p. 18); the red variety is also obtained by crystallising from hot KOHAq (Willm, A. Ch. [4] 5, 28); also by crystallising from hot KC, H₃O₂Aq (Werther, l.c.). The red form of TII is produced by adding KIAq to a warm conc. solution of a thallous salt (Hebberling, *l.c.*, p. 16). Red TII crystallises in octahedral forms (W., *l.c.*). The red variety slowly changes on keeping to the yellow form.

When freshly ppd. yellow TlI is exposed to the sun's rays it becomes green and crystalline without any change in composition. The green variety of TII is somewhat more soluble in hot water than the yellow form; it crystallises from hot water in green microscopic crystals that are very refractive. The green variety changes slowly to the yellow on keeping; the change is hastened by heating in water containing I or KI, but not by heating with pure water. strongly heated, green TlI becomes red (v. Knösel, B. 7, 576, 893).

Til is decomposed by warm HNO, Aq, with separation of I. It is dissolved, without I being separated, by ClAq or aqua regia; Willm (l.c.) supposes that an iodochloride is formed. Fusion

with KCN produces Tl (Werther, l.c.).
THALLIG IODIDE TIL. This compound has not been isolated with certainty. By treating Tl with an ethereal solution of I, Nicklès (C. R. 58, 537) obtained a brown liquid from which brown needles gradually separated. These crystals may have been TlI; no analyses are

Double salts of TII, with alkali iodides have been prepared by digesting TII with solutions of I and alkali iodides in ether; these salts have the composition TII.MI.xH2O, where M = NH4 and K (v. Nicklès, J. Ph. [4] 1, 25; Willm, A. Ch. [4] 5, 5; Rammelsberg, P. 146, 597). A compound Tll, Cul. 8NH, is described by Jörgensen (J. pr. [2] 6, 82).

THALLIUM SESQUI-IODIDE Tl.I. or TlI.3TlI = Tl₄I_a. (Thallothallic iodide.) Black, lustrous needles; insoluble in water, slightly soluble cold alcohol, decomposed by boiling alcohol; decomposed slowly at ordinary temperature, quickly on heating; obtained by long-continued heating yellow TlI with excess of IAq (Knösel, B. 7, 576,

893).

OTHER THALLO-THALLIC IODIDES. By evaporating a solution of TlI in HIAq, in presence of I, at 70°, small, brown, rhombic, crystals were obtained by Jörgensen (l.c.). The composition of the crystals is represented by J. as 6TlI.TlI, or perhaps 5TII.TII3. The same compound is said to be obtained by digesting TII with an ethereal solution of I. Heating to c. 100° drives off I, and leaves yellow TII; boiling alcohol, and also KIAq, withdraws I, forming TII.

Thallium, oxides of. Two oxides have been isolated, Tl₂O and Tl₂O₃; there is evidence in favour of the existence of a third oxide, probably

Thallous oxide Tl₂O. (Thallium monoxide.)
Mol. w. not known with certainty. [Tl²,O] = 42,240 (Th. 3, 354). Prepared by heating TI in air to not above 100° (to prevent formation of Tl_2O_3); also by heating TlOH to 100° out of contact with air; also by heating Tl2Os to full redness. A black powder; melting at c. 300° to a dark-yellow liquid, which acts on glass, forming a silicate of Tl. Tl₂O absorbs moisture from the air, forming TlOH; it dissolves easily in water, forming a strongly alkaline solution of TlOH (v. THALLOUS HYDROXIDE, p. 678). Soluble in absolute alcohol, forming C₂H₈.OTl, a colourless, heavy oil (S.G. 3.55); this oil is decomposed by a little water with separation of TIOH (v. THALLOUS HYDROXIDE, Preparation, No. 3, p. 678). (Crookes, C. J. 17, 128; Lamy, A. Ch. [3] 67, 385). Tl2O is reduced to Tl by heating to redness in H, or in CO, or with Mg (Winkler. B. 23, 788).

THALLIC OXIDE Tl₂O₂. Mol. w. not known with certainty. Prepared by ppg. a thallic salt thoroughly, drying, and heating it to c. 260° (Crookes, *l.c.*, p. 182). Also formed by burning I'l in air to a little below redness; at a lower temperature the product is chiefe. temperature the product is chiefly Tl₂O, and at a full red heat Tl₂O₃ is decomposed to Tl₂O and O (Lamy, l.c.). Tl₂O₃ is also formed by passing an electric current (from a couple of Grove cells) through neutral Tl₂SO, Aq, using Pt electrodes: Tl deposits on the negative electrode, and Tl.O. forms, as a brown cake, on the positive electrode; the H₂SO₄Aq formed should be neutralised from time to time by NH,Aq (Crookes, l.c.). For other methods of formation v. Wöhler (A. 146, 263, 375), and Willm (A. Oh. [4] 5, 19). Tl₂O₈ is a brownish-black solid; insoluble in water Melts at c. 760° (?) (Carnelley a. O'Shea, C. J. 45, 409). Gives off O at a full red heat, forming Tl.O. Dissolves in hot H₂SO₄Aq, forming Tl₂SO₄Aq and giving off O. Soluble cold HClAq, forming TlCl₃Aq, and some TlCl with evolution of Cl. Takes fire in dry H₂S, forming Tl₂S₃ (Carstanjen, J. pr. 102, 65). A mixture of Tl₂O₃ and S takes fire when struck (Böttger, L. pr. 0.87). J. pr. 90, 27). Reduced by heating in H or CO to Tl₂O, and then to Tl. Tl₂O₃ absorbs NH₃ when heated in that gas; no definite compound has been isolated. For compounds with \bar{H}_2O v.

THALLIC HYDROXIDES, p. 678). THALLIUM DIOXIDE (?TlO₂). When a rapid stream of Cl is passed into conc. KOHAq, holding freshly ppd. TlO.OH in suspension, a violetcoloured solution is obtained; mineral acids added to this solution form thallous salts, and give off O, or Cl with HClAq. Carstanjen (J. pr. 101, 55) supposed the violet solution to contain a thallate of potassium. Piccini (G. 17, 450) obtained a violet pp. by adding Ba(NO₃)₂Aq to the violet solution; analyses of this pp. were not satisfactory, but the results indicated that an oxide of Tl of the composition TlO2 is formed by the method described. Piccini says that the same violet solution is obtained by electrolysing 30 p.c. KOHAq, using a plate of Tl as anode;

in water, and adding Tl₂SO₄.

Thallium, phosphide of. No compound of Tl and P has been isolated with certainty. According to Crookes (C. J. 17, 135), the black substance formed by passing P hydride into ammoniacal Tl₂SO₄Aq is a phosphide of Tl. Carstanjen (J. pr. 102, 80) failed to obtain a phosphide by reducing Tl_aPO₄ by C or H, or by heating the elements together. Flemming (Bl. [2] 10, 35) noticed that a thin, black coating formed on molten Tl when pieces of P were thrown on to it; by heating solution of a thallous salt with P in a sealed tube, he obtained a black solid, perhaps a phosphide, and also a phosphite and hydride of P (cf. Willm, A. Ch. [4] 5, 5).

also by digesting 1 pt. KOH with 4 pts. KClO,

Thallium, platino-cyanide of; v. vol. ii. p. 345

Thallium, salts of. Two series of salts are formed by replacing the H of acids by Tl; thallous salts, Tl2X, and thallic salts, Tl2X, The where $X = SO_3$, CO_3 , $2NO_3$, $2IO_3$, $\frac{2}{3}PO_4$, &c. chief thallous salts of oxyacids are arsenates, carbonates, chlorate and perchlorate, chromate, dichromate and trichromate, dithionate and trithionate, hypophosphite, iodate and periodate, molybdates, nitrate, phosphates, selenite and selenates, silicates; sulphates and sulphite, tellurate, thiosulphate. The principal thallic salts are arsenate, iodate and periodate, nitrate, phosphates, sulphates (v. Carbonates, Nitrates, Sulphates, &c.). Several double salts of both series are also known.

Thallium, salts of oxyacids of. No compounds of this kind have been isolated with certainty, but there is evidence in favour of the existence of salts of the hypothetical thallic acid, probably analogous to plumbates (v. Thallium DIOXIDE, supra).

Thallium, selenides of. Tl-Se has been isolated; another selenide, perhaps Tl₂Se_s, pro-

bably exists.
Thallous selenide Tl₂Se. Mol. w. not known. Obtained by fusing the elements toge-

ther in the ratio 2Tl:Se (Willm, A. Ch. [4] 5, 5; Carstanjen, J. pr. 102, 79); also by ppg. Tl₂CO₂Aq by H₂Se (Kuhlmann, Bl. [2] 1, 330). A grey, lustrous, crystalline solid; melting at c. 840° (Kuhlmann, *l.c.*). Dissolves in H.SO,Aq and HClAq, giving off H.Se.

THALLIC SELENIDE (? Tl₂Se₃). By fusing the elements together in the ratio 2Tl:3Se, Carstanjen (l.c.) obtained greyish-black prismatic crystals, resembling Sb; not acted on by dilute H₂SO₄Aq; decomposed by hot conc. H₂SO₄, with separation of Se and evolution of SO₂. composition of this substance has not been determined.

Thallium, silicofluoride of, Tl.SiF. tained by dissolving Tl2CO, in H2SiF6Aq, and evaporating. Colourless, regular, octahedra (Werther, J. pr. 91, 385). According to Kuhlmann (Bl. [2] 1,330), the crystals are Tl_SiF_s.H_.O. Easily soluble in water. Volatile without decomposition, according to Kuhlmann (l.c.).

Thallium, sulphides of. Tl2S and Tl2S2 have been isolated, and also probably more than one

compound of these two sulphides.

THALLOUS SULPHIDE TLS. Mol. w. not

Formation.—1. By fusing the elements toge ther in the ratio 2Tl:S (Carstanjen, J. pr. 102, 76).-2. By ppg. an alkaline solution of a thallous salt by H₂S.—3. By ppg. slightly acidified Tl₂SO₄Aq by H₂S (Hebberling, A. 134, 11).— 4. By heating an alkaline solution of a thallous salt with Na₂S₂O₃ (C., l.c.).

Preparation.—A cold, fairly conc., aqueous solution of Tl2SO, is acidulated with a trace of H2SO4Aq, and saturated with H2S, and the bluish-black pp., consisting of microscopic crystals, is washed with H2SAq and dried over H2SO4

(Hebberling, l.c.).

Properties and Reactions.—Prepared by ppn. from slightly acidulated Tl₂SO₄Aq, Tl₂S forms very minute, blue-black, lustrous tetrahedra; prepared by heating together Tl and S it is a black, crystalline, brittle solid; prepared by ppn. from alkaline solutions it is a black amorphous solid. When the amorphous solid is dried, and strongly heated, it melts, and solidifies on cooling to a black crystalline mass. After melting, S.G. is 8.0 (Lamy, J. 15, 185). H.F. [Tl',S] = 21,660 (Th. 3, 354). Moist ppd. TLS oxidises in air to Tl2SO4. On long continued heating Tl2S is completely decomposed (Lamy, l.c.). Tl2S is insoluble in solutions of alkalis, alkali sulphides, carbonates, and cyanides; it dissolves readily in H2SO4Aq and HNO3Aq, and less readily in HClAq. By ppg. from a very dilute solution of a thallous salt, and dialysing, Winssinger (Bl. [2] 49, 452) obtained a reddishbrown aqueous solution of colloidal Tl.S

Combination.—By passing H2S into Tl2SO4Aq containing As2O3Aq, and acidulated with dilute H_2SO_4Aq , Gunning (Ar. N. 3, 86) obtained a red pp. of Tl₂S.As₂S₃; the same compound was obtained by mixing a solution of As₂S₃ in NH₃Aq with an ammoniacal solution of a thallous salt. Heat decomposes the compound into its constituent sulphides; alkali solutions dissolve out

As, S, leaving Tl,S.

THALLIC SULPHIDE Tl2S. Obtained as a black, amorphous, easily fusible, solid, by melting Tl with excess of S, and removing uncom-

bined S by heating out of contact with air (Carstanjen, J. pr. 102, 65). Below 12° Tl₂S₂ is hard and brittle; above 12° it is so soft that it may be drawn out like pitch. It dissolves in warm dilute H2SO, Aq without separation of S. Strecker (J. pr. [2] 2, 162) obtained a lustrous, brownblack pp. by passing H2S into a solution of a thallic salt mixed with sodium tartrate and excess of NH, Aq, followed by boiling; this pp. was Tl,S, according to Strecker. When H,S is passed into an aqueous solution of a thallic salt, the pp. is a mixture of Tl₂S and S. Tl₂S₂ combines with potassium and sodium sulphides, forming Tl.S.K.S and Tl.S.Na.S. These compounds are obtained by fusing together 1 pt. Tl₂SO₄, 6 pts. S, and 9 pts. K₂CO₂ or Na₂CO₂, and washing the fused mass with water. The K compound is more stable than the compound of Na; when heated it melts, and then gives off S, leaving K₂S and Tl₂S; it is not changed by water or alkali solutions, but is decomposed by mineral acids (v. Schneider, A. 135, 215; cf. Krüss, B. 19, 2738). These compounds may be regarded as KTlS, and NaTlS, that is, as salts of a hypothetical thio-thallic acid.

THALLO-THALLIC SULPHIDES. By adding (NH₄)₂SAq to a solution of Tl₂Cl₃, a black pp. is obtained which is more fusible than Tl.S. and after melting solidifies to large greyish-black prisms. This pp. contains Tl and S. Similar substances are formed by heating together Tl₂S and Tl2S3, and by melting Tl with more S than 2Tl:S, and less than 2Tl:3S; Carstanjen (J. pr. 102, 65) gave the formulæ $5\text{Tl}_2\text{S.}3\text{Tl}_2\text{S}_3 = \text{Tl}_{16}\text{S}_{14}$ and $\text{Tl}_2S, \text{Tl}_2S_3 = \text{TlS}$ to compounds obtained in this way. By heating $\text{Tl}_2S_3, \text{Na}_2S$ (v. supra) with water, Schneider (A. 135, 215) obtained a compound to which he gave the formula

 $Tl_2S.2Tl_2S_3 = Tl_6S_7$.

Thallium, sulphocyanide of; v. vol. ii. p. 352. Thallium, thiosalts of. The compounds of Tl₂S₃ with K₂S and Na₂S may be regarded as K and Na thiothallates (v. THALLIC SULPHIDE, M. M. P. M. supra).

THALLIUM ETHIDE TIEt. Has not been

obtained in the free state.

Ethylo-chloride TlEt2Cl. S. 2.76 in the cold; 3.37 at 92°. S. (alcohol) .33 at 78°. Formed from TlCl, and ZnEt, in ether (Hartwig, A. 176, 257). Scales (from water), explodes feebly at 190°, yielding TlCl. Reacts with silver salts forming: TlEt₂NO₂, S. 5.67 at 70°, exploding at 236°.—(TlEt₂)₂SO₄, S. 65.4 in the cold; 87.3 at 90°; exploding at 205° and converted by BaI, into TlEt,I, S. 1, exploding at 195°.— (TlEt₂), PO₄, S. 23.7 in the cold; 20.7 at 75°, exploding at 189° .- TIEt2OAc crystallising in needles [212°] (245°), exploding if rapidly heated.

Ethylo-hydroxide TlEt,OH. S. 15:1 in the cold; 8 at 88°. Formed from (TlEt,).SO, and baryta. Silky needles (from water). More sol. cold than hot water, forming an alkaline solution. Does not unite with CO2, but forms

the above salts with other acids.

THAPSIC ACID C₁₀H₁₀O₄. [124°]. Obtained from the root-bark of *Thapsia garganica*. The bark is extracted with ether, which leaves on evaporation an amber-coloured resin. The resin dissolves in potash with evolution of heat, and the solution, on acidification, yields a neutral resin, n-octoic acid and thapsic acid. The

alkaline solution on standing deposits potassium thapsate (Canzoneri, G. 13, 514). White scales (from alcohol), nearly insol. water and benzene, sol. alcohol and ether.

Salts.-K2A" xaq. Prisms.-BaA" (dried at 120°). Crystalline pp.—Ag₂A": amorphous. Anhydride C16H28O3. [71°]. Crystalline powder (from benzene).

Anilide C_{1e}H_{2e}O₂(NHPh)₂. [163°]. THEBAINE C_{1e}H_{2i}NO₃. [193°]. S.G. 1·3. $[a]_j = -219^\circ$ in 97 p.c. alcohol at 15° ; = -230° in CHOl₃; S. (ether) ·7 at 10°; S. (isoamyl alcohol) 1·67 in the cold; S. (benzene) 5·27 in the cold (Kubly, J. 1866, 823). Occurs in opium to the extent of about ·6 p.c. (Pelletier, J. Ph. 21, 569; Couerbe, A. Ch. [2] 59, 155; Kane, A. 19, 9; Anderson, A. 86, 179; Hesse, B. 3, 367; A. 153, 47; 176, 196). Prepared by treating the aqueous extract of opium with Na2CO3 or lime, exhausting the filtrate with ether, agitating the ether with dilute acetic acid, and pouring the acid liquid into dilute KOHAq. The pp. is collected after 24 hours, warmed with dilute alcohol and dilute HOAc is added till just acid. The alcohol is evaporated off at 50°, papaverine and narcotine removed by filtration, and thebaine acid tartrate ppd. by adding powdered tartaric acid to the filtrate. The tartrate is re-crystallised from water (Hesse, A. Suppl. 8, 262). Thebaine may also be obtained by adding NaOAc to a solution of the hydrochlorides of the opium alkaloids, allowing to stand, filtering, evaporating, again filtering, and adding sodium salicylate. After 24 hours thebaine salicylate crystallises (Plugge, Ar. Ph. [3] 24, 993; 25, 343).

Properties.—Colourless leaflets (from dilute alcohol) or prisms (from conc. alcohol). Tasteless. Alkaline in reaction. Very poisonous. Begins to sublime at 135°; at 160° the sublimate contains V. sol. needles, cubes, and prisms (Blyth). alcohol and chloroform. Ppd. by alkalis and alkaline carbonates, the pp. being slightly soluble in NH₂Aq and lime-water. Conc. H₂SO₄ gives a deep-red solution. Boiling dilute H.SO, converts it into thebenine and thebaïcine. Hot conc. HClAq at 90° yields MeCl and morphothebaine (Howard, B. 17, 527). HI \q gives MeI (2 mols.) (Howard a. Roser, B. 19, 1597). Bromine-water forms C₁₉H₂₀BrNO₂, a flocculent pp. converted by excess of Br into C19H20Br5NO3, a yellow pp. Solutions of its salts give a pp. of the dichromate

on adding K2Cr2O7

Salts.-B'HClaq. Large trimetric prisms. S. 6.3 at 10°. $[a]_j = -168.32 + 2.33p$. — $B'_2H_2PtCl_5$ 2aq. Amorphous pp., quickly changing to orange prisms. Decomposed by boiling water (O. De Coninck, Bl. [2] 45, 131).—
B',H,C,O, 6aq. S. 1.03 at 10°.—B'H,C,O, aq. S.
2.3 at 20°.—B'C,H,O,aq. Prisms, very soluble in hot alcohol. S. .77 at 20°.—Meconate
B',C,H,O, 6aq. White prisms (from alcohol).— B',H,FeCy, (Plugge, Ar. Ph. [3] 25, 793).

Methylo-iodide B'MeI. Crystallises from

alcohol in prisms (containing EtOH), m. sol.

Ethylo-chloride B'EtCl. Needles. Ethylo-iodide B'EtI. Needles (from alcohol) (Howard, B. 17, 527).

Bensylo-chloride B'C, H, Cl. Crystals. Morpho-thebaine C1, H1, ON [1919]. Obtained by heating thebaine with strong aqueous HCl or

HBr, MeCl (or EtCl) being split off; the yield being 50 p.c. of the thebaine. Yellowish crystals. Soluble in alcohol, ether, and benzene, sparingly in water. It dissolves in H2SO, to a colourless solution.

Salts.—B'HCl. Small glistening crystals, sol. water, insol. alcohol.—B'HBr: fine silky needles.—B'HNO, 2aq: small white crystals, v. sol. water and alcohol.-B'2H2SO47aq: small crystals, v. sol. water, insol. alcohol. The oxalate is a sparingly soluble amorphous pp. The picrate is a yellow pp. which melts under water. Insoluble pps. are produced potassium ferrocyanide, ammonium molybdate, $HgCl_2K_2I_2$, $K_2Cr_2O_7$, and bismuth-potassium iodide.

Acetyl derivative C17H16AcO3N. [183°]: glistening plates (Howard, B. 17, 529).

Ethylo-iodide *B'EtI. Crystallises from

acetic acid.

Methylo-iodide B'MeI. Dimetric tables, insol. alcohol, sol. HOAc.

Benzylo-chloride *B'C,H,Cl. Needles, m. sol. water, sl. sol. alcohol (Howard a. Roser, B. 19,

Thebenine C₁₂H₂₁NO₃. Formed by boiling thebaine with HClAq (S.G. 1.04) (Hesse, A. 153, 69). Amorphous. Insol. benzene and ether, sl. sol. boiling alcohol. Conc. H2SO, forms a blue V. sol. KOHAq.—B'HCl 3aq. S. 1. '2H2HgCl, 2aq. Very long efflorescent Plates.-B'2H2HgCl, 2aq. prisms. - B'2H2SO4 2aq. Plates, insol. cold water.

-B'H₂C₂O₄ aq. Prisms, sl. sol. hot water. Thebaïcine. Formed by boiling thebaïne or thebenine with conc. HClAq (Hesse). Yellow amorphous pp., insol. ether, benzene, and water, sl. sol. hot alcohol, v. sol. KOHAq. Conc. H₂SO₄ forms a dark-blue solution. The salts are resin-

Theine v. Caffeine. THEOBROMINE C, H, N, O2 i.e.

CO < NMe.C:CH.NMe C.NH.COMol. w. 180. methyl-xanthine. S. 062 at 17°; 7 at 100°. S. (alcohol) 02 at 17°; 2 at 78°. S. (ether) -906 in the cold; 17 at 35°. S. (boiling CHCl,) 1. Occurs in the seeds of Theobroma cacao to the extent of 1.3 to 4.6 p.c. (Woskresensky, A. 41, 125; Glasson, A. 61, 335; Keller, A. 92, 71; Rochleder, A. 71, 9; 79, 124; Strecker, A. 118, 151). Formed by heating the lead salt of xanthine $C_5H_2PbN_4O_2$ (4 pts.) with MeI (5 pts.) for twelve hours at 130° (Fischer, B. 15, 454; A. 215, 311). Prepared by ppg. the aqueous extract of the seeds with lead acetate, filtering, removing excess of lead by H2SO4, adding MgCO3, evaporating to dryness, and extracting with alcohol (Dragendorff, Ar. Ph. [3] 13, 1). Got also by freeing cacao from oil by pressure, adding slaked lime, and extracting with 80 p.c. alcohol (Schmidt

a. Pressler, A. 217, 287). Properties.—Minute trimetric crystals, sl. sol. hot water, alcohol, and ether. Tastes bitter. At 134° it begins to sublime; at 170° (Blyth) or 290° (S. a. P.) it yields a crystalline sublimate. Very slowly attacked by pure HNO, (Franchimont, R. T. C. 6, 223). Not attacked by boiling KOHAq. Its salts are more or less easily de-composed by water. Ammoniacal AgNO, gives a gelatinous pp. which dissolves on warming, and deposits, on boiling, crystalline C,H,AgN,O,

Theobromine is ppd. by sodium (Strecker). phosphotungstate (Wolfram, C. J. 36, 406; Fr. 18, 346).

Reactions.-1. Yields methyl-parabanic acid on oxidation by CrO_s.—2. KČlO_s and HClAq at 50° yield methyl-alloxan and methyl-urea (Fischer, A. 215, 304). Chlorine-water yields amalic acid. 3. Will not combine with MeI to amalic acid.—3. Will not combine with Mer to form a methyloiodide. But in presence of dilute alcoholic KOH it reacts with MeI, forming caffeine, even in the cold; better at 100°. But the reaction is never complete, some theobromine being decomposed.—4. Conc. HCl has no action below 200°. At 250° it decomposes it, producing formic acid, carbonic acid, ammonia, methylamine and sarcosine, C,H,N,O2+6H2O $= 2CO_2 + NMeH_2 + 2NH_3 + C_3H_7NO_2 + CH_2O_2$; the NH, and NMeH, occur in the proportion indicated by the formula (Schmidt, A. 217, 298). Boiling aqueous Ba(OH), effects in 40 hours the same decomposition.—5. HNO_s forms CO₂, methyl-parabanic acid and methylamine, but no ammonia.

Salts.—B'HClaq. Rosettes of needles.— IBr aq. Tables. — B'₂H₂PtCl_a4aq. — B'HBr aq. B',H,PtCl,5aq. — B'HAuCl, — B',HNO, — B'HOAc.—B',H,I, . Nearly black prisms, decomposed by alcohol (Jörgenssen, Z. [2] 5, 675).—B'HAg(NO₃)₂. Silvery needles, v. sl. sol. water. - C,H,AgN₄O₂1\(\)aq.—Ba(C,H,N₄O₂)₂(dried at 105°). Got by adding theobromine to barytawater (Maly a. Andreasch, M. 4, 379). Needles

(from hot water).

Bromo-theobromine C,H,N,O2Br. From theobromine and bromine (Fischer, A. 215, 305). Crystalline powder. Sl. sol. hot water. Sol. HCl, but reppd. by water. Sol. aqueous NaOH, not in NH,. Does not give an ethoxy-derivative with alcoholic KOH (unlike bromo-caffeine). After warming with ammonia it gives with ammoniacal AgNO, a silver salt that can be converted by EtI into bromo-ethyl-theobromine C,H,EtN,O,Br, whence alcoholic KOH forms eth - oxy - ethyl - theobromine C, H, EtN, O, (OEt) [153°] as needles. Boiling HCl converts this into EtCl and oxy - ethyl - theobromine oxy - ethyl - theobromine C,H,EtN,O2(OH), whence bromine and alcohol form the di-ethyl derivative of tri-oxy-di-hydroethyl-theobromine $C_7H_6EtN_4O_2(OH)(OEt)_2[152^\circ]$ This body on treatment with hot dilute HCl methylamine and apo-ethyl-theobromine (?); and with chlorine at -10° forms hypoethyl-theobromine C,H,N,O, [142°].

Methyl-theobromine is CAFFEINE.

Ethyl-theobromine C,H,EtN,O $_{x}$ [above 270°]. Formed from C,H,AgN,O $_{x}$ and EtI (Philips, B. 9, 1308). Small crystals, m. sol. hot water. Weak base.

THEOPHYLLIN C.H.N.O. i.s.

CO NH.C:CH .NMe [264°]. Occurs in alcoholic extract of tea, from which it may be separated by means of its Ag salt (Kossel, B. 21, 2164). Monoclinic tables (containing aq, v. e. sol. warm water and very dilute ammonia. May be sublimed. When evaporated with chlorinewater it yields a scarlet residue, which is coloured violet by NH,Aq. MeI converts its Ag salt into caffeine. KClO, and HClAq yield dimethyl-alloxan.

Salt.—AgC,H,N,O, jaq. Amorph which may be crystallised from NH,Aq. Amorphous pp.,

A name given by Crum-Brown and Letts to compounds of the form CO₂H.CH₂.SR₂.OH or their anhydrides CH_2 - SR_2 where R is a hydrocarbon radicle; v. DI-ETHYL-THETINE and DI-METHYL-THETINE

THEVETIN C₃₄H₈₄O₂₄ 3aq. [170°]. S. 8 at 14°. $\alpha = -85.5°$. A poisonous glucoside occurring in the kernels of Thevetia nereifolia (Blas, J. 1868, 768; De Vrij, R. T. C. 3, 167; Warden, Ph. [3] 12, 417). Crystalline powder, v. sol. hot water and alcohol, insol. ether. Sol. acids and alkalis. Lævorotatory. Conc. H₂SO₄ gives a dirty-brown colour. Decomposed by boiling dilute HClAq into glucose and theveresin $C_{48}H_{70}O_{17}$ 2aq, a white powder [140°], v. sl. sol. cold water, sol. alkalis.

THIACETIC ACID v. THIO-ACETIC ACID. THIALDINE C.H13NS2 i.e.

NH<CHMe.S>CHMe. Mol. w. 163. [43°]. Formed by passing H2S for four hours into a solution of aldehyde-ammonia (1 pt.) in water (15 pts.), and a little NH, Aq (Wöhler a. Liebig, A. 61, 2; Hofman, A. 103, 93; Brusewitz a. Cathander, J. pr. 98, 315). Monoclinic crystals (from etheralcohol). May be distilled with steam. V. sl. sol. water, v. sol. alcohol, v. e. sol. ether. Boiling dilute H₂SO, forms thio-acetic aldehyde. KMnO₄ forms CH₂CH(SO₂K). (Guareschi, B. 11, 1204, 1600). Policia (NO₂ VO) 1384, 1692). Boiling AgNO, Aq forms aldehyde. MeI in ether forms crystalline C₆H₁₃MeNS₂I (Hofmann, A. 103, 93). I in KI forms amorphous (CHMe)₂S, (Fassbender, B. 20, 460). Salts.—B'HCl. Prisms, v. sol. water.-

B'HBr.—B'HI.—B'HNO_s.—B'H₂SO₄. Prisms.—

B'H,PO, aq. Needles, v. sol. water.

THIAZOLE N CH.S. (116.8° cor.). S.G. $\frac{17}{7}$ 1·1979; $\frac{17}{17}$ 1·1998. Formed by the action of boiling alcohol on the diazo- compound of amidothiazole, obtained from di-chloro-di-ethyl oxide and thio-urea (Hantzsch a. Popp, B. 21, 2582; A. 250, 274). Formed also from NeCCl.S. and the corresponding μ -bromo-thiazole by reduction with zinc-dust and HOAc (Schatzmann, A. 261, 12). Mobile liquid, very volatile, hygroscopic, and highly refractive. Miscible with water, alcohol, and ether. Insol. KOHAq. Smells like pyridine. Neutral in reaction. -Balts.—B',H,PtCl₂ 2aq.—B'HAuCl₄. [250°].—B'C₄H₄N₄O₇. [151°].—B'HHgCl₃. [104°].—B'HgCl₄. [204°]. Crystalline pp., sl. sol. water.

 μ -Amido-thiazole $N \ll_{CH}^{C(NH_2).S}$ Thiazo-Formed by the fine. Thiazylamine. [90°]. action of chloro-acetic aldehyde on thio-urea (Traumann, A. 249, 36). Prepared by boiling CH,Cl.CHCl.OEt (1 pt.) with thio-ures (1 pt.) and water (5 pts.). Yellowish plates, sl. sol. water, alcohol, and ether. Strong base. -B'HCl aq. Needles. -B', H, PtCl. Ye.

Acetyl derivative. Yellow tables.

[203°]. Needles. THIAZOLE-TRIAZOLE (so called)

 $CH \stackrel{S-C:N}{\sim} CH$. Formed by the action of HClAq on methyl-nitroso-imido-thiazoline (Näf, A. 265, 123). Liquid, v. sol. water. Alkaline; reduces Fehling's solution. Nitrous acid reproduces nitroso - methyl - imido - thiazoline. -B'HCl 2aq. [220°]. Needles.—B'HBr 2aq.

THIENONE is DI-THIENYL KETONE.

DITHIENYL C.H.S. i.e. C.H.S.C.H.S. [83°]. (266° cor.). Formed on passing thiophene through a red-hot tube (Nahnsen, B. 17, 789, 2197). Plates (from alcohol). Conc. H.SO. gives a reddish-brown solution with deep-green fluorescence. On warming with isatin and H₂SO, it gives a violet-blue colour.

Sulphonic acid C,H,S,(SO,H). Got by heating with H₂SO₄ (20 pts.) at 100°. Yields a

crystalline BaA', aq.

Reference.—HEXA-BROMO-DITHIENYL

THIENYL-ACETIC ACID C,H,S.CH,CO,H. [76°]. Formed from $C_4H_3S.CH(OH).CO_2H$ [115°], HIAq, and P (Ernst, B. 19, 3281). Crystalline, sol. hot water, alcohol and ether.—Ba'A, (dried at 130°). Crystals.—AgA': pp.

Reference .- OXY-THIENYL-ACETIC ACID.

THIENYL-ACRYLIC ACID C,H,SO, [138°]. C₄H₅S.CH:CH.CO₂H. Formed b₹ boiling thiophenic aldehyde with NaOAc and Ac₂O (Biedermann, B. 19, 1855). Needles (from water), sol. hot water and alcohol. FeCl, gives an amorphous orange pp.—AgA': white pp.

THIENYLAMINE v. AMIDO-THIOPHENE. THIENYL-BROMO-ETHANE

THIENYL-ETHANE.

THIENYL-BROMO-ETHYLENE v. BROMO-THIENYL-ETHYLENE

THIENYL BROMO-METHYL KETONE

C₄H₃S.CO.CH₂Br. ω-Bromo-acetothienone. Formed by adding Br (2 at.) to thienyl methyl ketone (1 mol.) dissolved in CS, (Brunswig, B. 19, 2891). Pungent yellow oil, may be distilled in vacuo. V. sol. alcohol and ether. Solidified by cold. Alcoholic NH, forms a deep-blue colour. Oxidised to (a)-thiophenic acid [126.5°]. Aniline yields C,H,S.CO.CH,NHPh [80°].

Thienyl di-bromo-methyl ketone

C₄H₃S.CO.CHBr₂. [c. 0°]. Formed from thienyl methyl ketone and Br. Pungent oil, almost insol. ligroïn.

(B)-THIENYL-CARBINOL C.H.S.CH,OH. (207° cor.). Formed by shaking (\$\beta\$)-thiophenic aldehyde with aqueous potash (Biedermann, \$B\$. 19, 639). Liquid. HCl forms C, H, S.CH, Cl, a pungent oil boiling with partial decomposition at 175°.

THIENYL CHLORO-ETHANE v. CHLORO-THIENYL-ETHANE.

THIENYL-CHLORO-ETHYLENE v. CHLORO-THIENYL-ETHYLENE.

(a)-THIENYL CHLORO-METHYL KETONE C₄H₃S.CO.CH₂Cl. [47°]. (259° cor.). Formed by chlorination of thienyl methyl ketone (Peter, B. 18, 539). Pungent white crystals. Yields (a)-thiophenic acid on oxidation.

THIENYL-ETHYL-AMINE C.H.NS $C_4H_2S.CHMe.NH_2$. (186°). Got by reducing the oxim of thienyl methyl ketone in alcoholic solution by adding sodium-amalgam and HOAc (Goldschmidt a. Schulthess, B. 20, 1700). Liquid, sol. water. Absorbs CO2 from the air. B'ĤOAc. Needles, v. e. sol. water.

Benzoyl derivative. [95°]. THIENYL ETHYL KETONE C.H.S.CO.C.H. Propiothienone. (228° cor.). Formed from propionyl chloride, thiophene, and AlCl, (Krekeler, B. 19, 677). Liquid. Hot conc. H₂SO₄ forms propionic and thiophene (a)-sulphonic acids Chromic acid mixture yields (a)-thiophenic acid Oxim C.H.S.CEt:NOH. [56°]

THIENYL DIETHYL PHOSPHINE

(C,H,S)PEt₂. (225°). Formed from thienylchloro-phosphine and ZnEt, in ether (Sachs, B. 25, 1517).

Ethylo-iodide (C.H.S)PEt.I.

Methylo-iodide (C,H,S)PEt,MeI. [122°]. White powder, v. sol. water insol. ether.

THIENYL-GLYCOLLIC ACID v. OXY-THIENYL-

(a)-THIENYL-GLYOXYLIC ACID

C₆H₄SO₃ aq i.e. C₄H₃S.CO.CO₂H. [91°]. Formed by oxidising thienyl methyl ketone with alkaline KMnO, (Peter, B. 18, 537; Biedermann, B. 19, 637; Bradley, B. 19, 2115). Formed also, together with thiophene (a) carboxylic acid, by oxidation of (α)-ethyl-thiophene by alkaline Crystals (containing aq, possibly as C.H.S.C(OH)₂.CO₂H), v. sol. warm water. Melts at 59° when hydrated. May be extracted by ether from its aqueous solution. If a drop of crude benzene, followed by H₂SO₄, be added to its solution in HOAc, a brown colour is formed, which on addition of more H2SO, passes through blue-green, green, violet, red, and bluish-violet to blue.

Reactions.-1. On heating it splits up into CO₂ and thiophenic aldehyde.—2. On heating with di-methyl-aniline and ZnCl₂ it yields the leuco-base of a green dye C, H, S.CH(C, H, NMe,)2.

PCl_s forms thiophenic chloride.

Salts. - BaA'2 aq. Slender efflorescent needles, v. sol. water.—CaA'2 2aq.—CuA'2 2aq. ZnA'₂ 2aq.—AgA' aq: amorphous pp., probably C₄H₃S.C(OH)₂.CO₂Ag.

Methyl ether MeA'. [28.5°]. Ethyl ether EtA'. (265° uncor.). Oil. Amide C, H, S.CO.CO.NH2. [88°]. Needles, almost insol. water, v. sol. alcohol and ether.

C₄H₃S.C(NOH).CO₂H. (Hantzsch, B. 24, 48). Slender white needles. Exhibits Liebermann's reaction (Bradley, B. 19, 2119).—BaA', 1; aq.—AgA': amorphous pp.— MeA'. [105°]. Converted by NaOMe and MeI [105°]. Converted by NaOMe and Mel liquid C₄H₃S.C(NOMe).CO₂Me. — EtA'. into [123°]. Needles.

Acetyl derivative of the oxim. [87°].

Phenyl-hydrazide

C₁H₂S.C(N₂HPh).CO₂H. [165°]. Deep-yellow

needles (from ether).

Reference.—NITRO-THIENYL-GLYOXYLIC ACID. THIENYL HEXYL KETONE C₁₁H₁₆SO i.e. C₄H₃S.CO.C₆H₁₉. (304° cor.). Formed from thiophene, heptoyl chloride, and AlCl, in ligroin (Schleicher, B. 19, 664). Yellow oil, yielding (a)-thiophenic acid on oxidation. Conc. H₂SO₄ forms thiophene (a)-sulphonic and heptoic acids.

Oxim [49°]. Crystals (from alcohol).

THIENYL-INDOLE C₁₂H₂NS i.e.

 $C_4H_4 < C_{NH}^{CH} > C.C_4H_3S$. [162°]. Formed by heating the phenyl-hydrazide of thienyl methyl ketone with ZnCl₂ (Brunck, A. 272, 201). Lightyellow needles, insol. water, v. sol. ether, m. sol. alcohol. Colours pine-wood, moistened with HCl, bluish-violet. Yields C₁₂H₂Br₂NS [278°] and C₂H₃CH(C₁₂H₂NS)₂ [245°]. — B'C₂H₂N₂O₂ [187°]. Dark red plates, v. sol. alcohol.

Nitrosamine C₁₂H₃(NO)NS. [241°].

DI-THIENYL KETONE (C,H,S)200. [887]. (826° uncor.). Formed from thiophene, COCl2, and AlCl. Obtained also by distilling calcium a-thiophenate (Gattermann, B. 18, 3012). Needles or plates, sol. alcohol.

Phenyl-hydraside [137°].
THIENYL MERCAPTAN (C,H,S).SH. (166°). Got by reducing thiophene (a)-sulphinic acid with zinc and HClAq (Biedermann, B. 19, 1616). Got also in small quantity from succinic acid and P_2S_5 (V. Meyer a. Neure, B. 20, 1757). Stinking oil, volatile with steam.—AgS.C₄H₃S.

Acetyl derivative (C,H,S).SAc. (231°).

Methyl ether (C,H,S)SMe. (186°).

DI-THIENYL METHANE C,H,S, i.e.

CH₂(C₄H₂S)₂. (267°). Got by adding H₂SO₄ to a solution of thiophene and methylal in HOAc (Peter, B. 17, 1345). Oil, smelling like oranges. Volatile with steam. Gives a red colour with isatin and H,SO,

(a)-THIENYL METHYL KETONE CaHaSO i.e. C, H, S.CO.CH, Acetothienone. (213° cor.). S.G. 24 1·167. Formed from thiophene, AcCl, and AlCl₃ (Peter, B. 17, 2643; Biedermann, B. 19, 636; Bradley, B. 19, 2115). Formed also from C.H₂(HgCl)S and AcCl (Volhard, A. 267, 170). 178). Liquid, smelling like acetophenone. Gives a blue colour with isatin and H2SO4. Oxidised by KMnO, to (α)-thienyl-glyoxylic and thiophenic acids. Mercuric chloride and NaOAc form the compound (CH₃.CO.C₄H₃S)HgCl₂ crystallising in white needles. Oxalic ether (1 mol.) and NaOEt give C,H,S.CO.CH,CO.CO,Et [42°], which forms large yellow crystals, converted by boiling alcoholic hydroxylamine hydrochloride into the compound C,H,S.C \(\sum_N \). O

 $CO_2Et.C < \stackrel{CH:C.C}{\sim}_N \stackrel{H_3S}{\cdot}_{O}$ crystallising in colour-

less needles [48°] (Angeli, B. 24, 232).

Oxim C.H.S(NOH). [112°]. Phenyl-hydrazide C.H.S(N2HPh). [96°]. Reference.-Bromo-, Chloro-, Iodo-, and NITRO-THIENYL METHYL KETONE.

THIENYL PHENYL - AMIDO - METHYL **KETONE** C₄H₃S.CO.CH₂NHPh. [80°]. Formed by mixing C₄H₃S.CO.CH₂Br with aniline in alcohol (Brunswig, B. 19, 2892). Plates. Yields a nitrosamine C4H3S.CO.CH2.NPh(NO) [81°], and acetyl derivative C,H,S.CO.CH,NPhAc [142°].

THIENYL PHÈNYL KETONE $C_4H_8S.CO.C_6H_5$. [55°]. (c. 300°). Formed by the action of AlCl, on a mixture of thiophene and BzCl (Comey, B. 17, 790). Formed also by heating C, H, (HgCl)S with BzCl at 100° (Volhard, A. 267, 179). Long needles (from dilute alcohol), insol. water. Gives a blue colour on heating with isatin and H₂SO₄. Yields thiophene and benzoic acid on heating with soda-lime.

Oxims C,H,S.CPh:NOH. Hydroxylamine forms two stereo-isomeric (?) oxims [93°] and [114°] (Hantzsch, B. 23, 2332; 24, 59).

THIENYL-PHENYL-METHANE

CH.Ph.C.H.S. (265°). Formed by the action of H2SO, on a mixture of thiophene and benzyl alcohol dissolved in HOAc (Peter, B. 17, 1346). Liquid with fruity smell, volatile with steam. Gives a red colour with isatin and H,SO.

Thienyl-di-phenyl-methane CHPh_x.C₄H_xS. (830°-340°). Formed from thiophene, di-phenyl-carbinol, and P2O, (Levi, B. 19, 1624). Plates (from alcohol) or needles (from ligroïn), v. sol. HOAc, alcohol, and ether. Crystallises from cold benzene in needles C₁,H₁₄SC₆H₆ [48°]. THIENYL ISOPBOPYL KETONE

C,H₃S.CO.CHMe₂. (232° cor.). Formed from isobutyryl chloride, thiophene, and AlCl₃ (Krekeler, B. 19, 675). Liquid. Yields (a)thiophenic acid on oxidation. Fuming H.SO. reacts with formation of the sulphonic acid C,H2(SO3H)S.CO.CHMe2, which yields the salts BaA'2, PbA'2xaq, and is converted by phenyl-hydrazine into C₄H₂(SO₈N₂H₄Ph)S.C(N₂HPh).CHMe₂, crystallising from hot water in plates.

Oxim C.H.S.C(NOH).CHMe. [108°]. Pearly

plates (from dilute alcohol).

THIENYL STYRYL KETONE v. STYRYL

THIENYL KETONE.

THIENYL DISULPHIDE (C4H2S)2S2. [56°] Formed by atmospheric oxidation of a solution of thienyl mercaptan in alcoholic NH₃ (Meyer a. Neure, B. 20, 1757). Needles.

SÚLPHOCYANO - METHYL THIENYL KETONE CH₂(SCy).CO.C₄H₃S. [88°]. Formed from CH₂Br.CO.C₄H₃S and KSCy in alcohol (Brunswig, B. 19, 2893). Plates (from ether), sl. sol. water and ligroin, m. sol. alcohol.

DI-THIENYL-THIOPHENE C,SH2(C,SH3)2. [147°]. (357°). V.D. 8.6. One of the products of the passage of benzene and sulphur-vapour through a red-hot tube (Renard, C. R. 112, 49). Yellowish needles (from alcohol), m. sol. benzene. H.SO, forms a rose-coloured solution, turned violet and finally blue on heating. Fuming H2SO4 at 120° forms a trisulphonic acid, which yields ($C_{12}H_3S_3(SO_3)_3)_2Ca_3$ as an amorphous powder, v. sol. water. Fuming HNO₃ at 160° forms $C_{12}H_3S_2O_4$ [313°], crystallising in white plates, v. sl. sol. benzene. Br forms $C_{12}H_4S_2O_3$ [232°], and amorphous C₁₂H₂Br₆S₃.

THIENYL TOLYL KETONE C₁₂H₁₀SO i.e.

C₆H₄Me.CO.C₄H₃S. Formed from o - toluic chloride, thiophene, and AlCl, (Ernst, B. 19,

3279). Oil. Yields an oily oxim.

THIO-. This prefix is employed to denote the replacement of O by S. Most inorganic salts of thio- acids are described under the heading of the oxy-salts; thus thio-carbonates are described under Carbonates. In some cases thio- salts are described under the heading of the characteristic element of the salts: thio-arsenates are to be found under Arsenic. Certain thio -salts are described under headings placed in their strictly alphabetical position; for instance, Thiosulphates. Thiocyanides are described as SULPHOCYANIDES.

The following addition should be made to the sub-article Thio-oxyonthophosphates (p. 147). By heating metallic chlorides or sulphides with excess of P2S5 Glatzel has prepared a number of tetrathio - oxyorthophosphates, i.e. salts of the hypothetical acid H₂PS₄(Zeit. für anorg. Chemie, 4, 186). These salts are crystalline—generally green to black in colour; they are burnt in air, giving off SO₂; some are decomposed to sulphides and P2S3, but the mercuric and arsenic salts distil unchanged when heated out of contact with air. These salts are insol. in dilute acids; they are decomposed by strong acids. G. isolated the thiophosphates of Sb, As, Bi, Cd, Cu(ous), Fe(ous), Pb, Mn, Hg(io), Ni, Ag, Tl, Sn(ous), and Zn; he sould not obtain the salts of Ca, K, Na, or Sr.

M. M. P. M.

CH2.CS.NH2. THIOACETAMIDE [108°]. Formed from acetonitrile and H.S (Bernthsen, A. 192, 46), and from acetamide and P₂S₅ (Hofmann, B. 11, 340). Monoclinic plates, v. sol. water and alcohol, m. sol. ether. HgO forms HgS. AgNO. gives Ag₂S. CuSO₄ and Pb(OAc)₂ give, on boiling, CuS and PbS. HgCl, forms a crystalline com-∞ - bromo - acetoacetic ether forms CH₃.C(NH).S.CH₂.CO.CH₂.CO₂Et [940],

finally SCMe:N CO.CH, CO.Et (239°), the ether of methyl-thiazyl-acetic acid [121°] (Steude, A. 261, 36). The ether melting at 94° is converted by warming with water into the compound CH₃.CO.S.CH₂.CO.CH₂.CO₂Et (155° at 15 mm.), whence H₂SO₄ forms an acid C₆H₈SO₂ [168°].

THIO-ACETIC ACID C2H, SO i.e. CH3.CO.SH. Mol. w. 76. (93°). S.G. 10 1.074. Prepared by distilling acetic acid with P₂S, (Kekulé, A. 90, 311; Ulrich, A. 109, 272; Kekulé a. Linnemann, A. 123, 273). Obtained also from AcCl and KSH (Jacquemin a. Vosselmann, J. 1859, 354), by heating lead acetate with Na,8,0, (Fröhde, Z. 1866, 543), and by the action of KSH on an alcoholic solution of PhSAc (Kekulé, Z. [2] 3, 196). Colourless liquid, turning yellow on keeping. Smells like acetic acid and H.S. Not solid at -17° . M. sol. warm water, v. sol. alcohol and ether. Violently attacked by fuming HNO₂. PCl₃ yields AcCl, PSCl₄, and HCl. Its neutral solutions give pps. with AgNO₃ and HgCl₂, which quickly turn black. Molten ZnCl₂ forms CMe:S.:CMe [225°], which may be crystallised from hot alcohol (Bongartz, B. 19, 2182). HCl passed into a mixture of thio-acetic acid and benzoic aldehyde forms (CH3, CO.S)2CHPh [148°], crystallising from hot ether in slender needles.

Salts.-KA'. Crystals, v. sol. water and alcohol.—NaA' 1aq. - BaA'2 3aq. - SrA'2 2aq. CaA'₂ 2aq.—PbA'₂. water or alcohol). Silky needles (from hot Quickly decomposes, with

separation of PbS.

Methyl ether MeA'. (96°). Formed by heating AcBr with Me₂S at 100° (Cahours, Bl. 25, 562; Wallach a. Bleibtreu, B. 12, 1062). Formed also from AcCl and Pb(SMe)2 (Ober-

meyer, B. 20, 2920)

(116° i.V.). Formed Ethyl ether EtA'. from AcCl and NaSEt (Saytzeff, Z. [2] 4, 642; Beckmann, J. pr. [2] 17, 461). Formed also from PhOAc and NaSEt (Seiffert, J. pr. [2] 31, 468). Colourless liquid, with alliaceous odour. Oxidised by AgMnO, to acetic acid and ethane sulphonic acid.

n-Propyl ether PrA'. (136°). Isopropyl ether PrA'. (126°). Isobutyl ether CH₂PrA'. (149°). Anhydride v. ACETYL SULPHIDE. Thio-acetic acid CH, CS.OH. Amide v. THIOACETAMIDE. Anilide CH3.CS.NHPh.

[75°]. Formed from CH₂.CCl:NPh and H₂S (Leo, B. 10, 2134). Prepared by heating acetanilide with P.S. at 100° (Hofmann, B. 11, 339; Jacobsen, B. 19, 1071). Yellowish needles (from water). Sol. NaOHAq and reppd. by acids. On oxidation with alkaline K.FeOy, it yields ethenyl-amidophenyl mercaptan. MeI forms C.H.NSMeI or (CH.,CS.NPhMe)HI [189°], which yields an aqueous solution from which AgNO₂ at once

ppts. AgI, and which is converted by AgCl into C.H., NSMcCl, whence (C.H., NSMcCl), PtCl, may be got (Bernthsen, A. 192, 56). NaOEt forms CH., C(SNa): NPh, whence MeI yields the ether CH. C(SMe): NPh (245°) (Wallach a. Bleibtreu, B. 12, 1061; 13, 529), which is decomposed by MeI at 100° into methyl thioacetate and methyl-NaOEt and EtBr yield the corresponding CH, C(SEt): NPh, a heavy oil, insol. alkalis, split up by warm HClAq into thioacetic ether and aniline. 'Ethyl-iso-thioacetanilide' yields (C₁₀H₁₈NS)₂H₂PtCl₂. Methyl anilide CH₂.CS.NMePh. [59°].

(290°). Formed from AcNMePh and P2S5 (Wallach, B. 13, 528). Monoclinic tables, v. sol.

alcohol.

o-Toluide CH3.CS.NHC7H7. [68°]. Yields CH₃.C(SEt):NC₆H₄Me (262°), an oil which reacts with bases forming amidines and EtSH.

p-Toluide [132°]. Yields oily CH₂.C(SEt):NC,H₂. (273°) (Wallach a. Wüsten,

B. 16, 147)

Xylidide OH, CS.NHC, H, Me. [80°] (G.); [95°] (Jacobsen). Formed by heating acetylxylidine with P2S, on the water-bath (Gudeman, B. 21, 2551).

ψ-Cumidide CH, CS.NHC, H, Mes. [114°].

Prisms (Jacobsen a. Elley, B. 22, 907).

(a)-Naphthalide CH₃.CS.NHC₁₀H₇. [96°]. Tables. Formed from naphthyl-acetamidine

and CS₂ (Bernthsen a. Trompetter, B. 11, 1760).

(β)-Naphthalide. [146°]. Needles or plates (Jacobsen, B. 21, 2627). Oxidised by K₂FeCy₆ to $C_{10}H_6 < N CMe$ [81°].

Tri-thio-ortho-acetic acid CH₂.C(SH)₃.

Ethyl ether CH₃.C(SEt)₃. Formed from mercaptan, CHCl₂, and NaOH at 100° (Laves, B. 25, 354). Brownish liquid. Oxidised by KMnO, and H2SO, to the trisulphone.

Benzyl ether CH_s.C(SC,H₂)_s. [46°]. Formed from benzyl mercaptan, CHCl_s, and (2 p.c.) NaOHAq (Laves). Crystals, v. sol.

CHCl, m. sol. alcohol.

THIO-ACETIC ALDEHYDE. On passing H.S into an aqueous solution of aldehyde, crude thioacetic aldehyde separates as an oil, which solidifies at -8° , and then melts at -2° . This oil [V.D. 60 (H = 1)] is a mixture of thioacetic aldehyde and its polymerides, and may be represented, in a state of vapour, as C₂H₄S + C₆H₁₂S₈.

(Pinner, B. 4, 258; Klinger, B. 9, 1893; 10, 1877; 11, 1023). On distillation, or on passing HCl through the liquid, it is converted into the polymeric tri-thio-tri-acetic aldehyde.

Tri-thio-tri-acetic aldehyde C.H., S. Mol. w. 180. [46°]. V.D. 89.7 (calc. 90) (Hofmann, B. 3, 589; Z. [2] 6, 699). Formed by passing H.S into an acid solution of aldehyde (Klinger; cf. Weidenbusch, A. 66, 152; Crafts, C. R. 54, 1279). Formed also from paraldehyde and H2S. White needles, sol. alcohol and ether. Smells like garlic. AgNO, added to its alcoholic solution forms crystalline C.H. S. 2AgNO. Chlorine forms ethylidene chloride (Pinner, A. 179, 21). KMnO, oxidises it to CH, CH(SO, H)2 and oxysulphides, including C_eH₁₂S₂O₃ [217°], which crystallises from hot water in needles (Guareschi, A. 222, 301) Zinc permanganate produces CaH128,O., which does not melt below

245°, $C_8H_{12}S_8O_2$ [112°-116°], and $C_8H_{12}S_8O_4$ [228°-231°] (cf. Baumann, B. 26, 2074).

(a)-Tri-thio-tri-acetic aldehyde CaH12S, i.e. CHMe S.CHMe S. [102°]. (247°). V.D. 6 (air = 1). Formed by the long-continued action of H₂S on a mixture of equal parts of water, conc. HClAq and aldehyde. Long prisms (from acetone). Reduced by HI to Et_2S_2 . Oxidised by KMnO, to $\text{C}_4\text{H}_{12}\text{S}_3\text{O}_4$ [284°] (S. 1 at 100°; 116 at 20°), and $C_8H_{12}S_8O_6$ or $C_2H_4 < \frac{SO_2 \cdot C_2H_4}{SO_2 \cdot C_2H_4} > SO_2$, which softens at 340° and is converted by MeI into C_pH₁₈S₂O₆ [302°]. Yields crystalline C₆H₁₂S₃AgNO₃ and C₆H₁₂S₃3AgNO₃.

(β)-Tri-thio-tri-acetic aldehyde C6H12S3. [126°]. (247°). V.D. 6.0. Formed by the action of AcCl or H_2 SO, on its (a)-isomeride. Prepared by passing H_2 S into a mixture of aldehyde (1 pt.) and alcohol previously saturated with HCl (3 pts.) (Baumann & Fromm, B. 22, 2600; 26,2074). Long needles (from alcohol). Oxidised by KMnO₄ to the same sulphones as the (a)isomeride. Reacts with MeI, yielding SMè₄I. Distillation with Cu forms ψ -butylene. Forms

crystalline C.H. S.AgNO, and C.H. S.3AgNO. (γ)-Tri-thio-acetic aldehyde C₆H₁₂S₃. [76°]. (242°). Formed, together with C₅H₆NS₃, by boiling thialdine sulphocyanide with water (Marckwald, B. 19, 1827; 20, 2817). Needles (from alcohol), insol. water, v. sol. ether and CS₂. Volatile with steam. Sol. conc. H₂SO₄ and reppd. by water. When mixed with EtI (4 pts.) and left for some weeks it suddenly changes to a crystalline mass of the (β)-isomeride. $C_eH_{12}S_3SAgNO_3$. Plates, v. sol. water. Yields Ag_2S on boiling with alcohol.

'Di - thialdehyde thiocarbimide' C₅H₉NS₃. [138°]. Formed as above. Needles, m. sol. ether. Yields C₅H₉NS₂AgNO₃ v. sol. water, and (C₅H₉NS₂)₂PtCl₄.

orthoaldehyde. Di-thio-acetic ether CH₃.CH(SEt)₂. Formed by passing HCl into a mixture of acetic aldehyde and EtSH (Baumann, B. 18, 884). Mobile oil.

THIOACETIC ANHYLRIDE v. ACETYL SUL-

THIO-ACETONE *CH3.CS.CH3. A product of the action of H2S on acetone, and formed also when CMe₂(SEt)₂ is heated above 160° (Baumann a. Fromm, B. 22, 2592). Volatile oil, with very persistent and disagreeable smell. Unstable.

Duplo-thioacetone $C_0H_{12}S_2$ i.e. $S < \frac{CMe_2}{CMe_2} > S$. V.D. 5.08 (calc. 5.11). (184° cor.). Formed by heating acetone with P₂S₃ at 125° (Wislicenus, Z. [2] 5, 324; Autenrieth, B. 20, 375). Formed also by heating tri-thio-tri-acetone at 200° for some time (Fromm a. Baumann, B. 22, 1043). Its alcoholic solution gives a white pp. with HgCl. Reduced by sodium-amalgam to isopropyl mercaptan (Spring, Bl. [2] 40, 69). Oxidised by a 5 p.c. solution of KMnO, to the disulphone SO₂ CMe₂ SO₂ [220°-225°].

Tri-thio-tri-acetone C.H, S. i.e. S CMe₂.S CMe₂ [24°]. (130° at 15 mm.). Formed, together with C₁₈H₂₈S₄ [171°], by passing H2S into a cooled mixture of acetone and

conc. HClAq (Fromm a. Baumann, B. 22, 1037, 2597). Needles (from alcohol); insol. water, v. sol. alcohol and ether. Boils, with much decomposition, at 225°-230°. Gives a white pp. with HgCl, and a yellow pp. with lead acetate. Potassium permanganate and H2SO, yield the trisulphone $SO_2 < \stackrel{CMe_2.SO_2}{Cme_2.SO_2} > CMe_2$ [302°] and

 $S < CMe_2.SO_2 > CMe_2$ [208°].

Duplo-di-thio-acetone $(Me_2CS_2)_2$. [98°]. (c. 243°). Formed from acetone and yellow ammonium sulphide in the cold (Willgerodt, B. 20, 2467). Dimetric crystals, a:c=1: 868, insol. water. Partially decomposed on distillation.

Di-thio-orthacetone. Di-ethylCMe₂(SEt)₂. Acetone-ethyl-mercaptol. (191°). Formed by passing dry HCl into a mixture of acetone (1 pt.) and mercaptan (2 pts.) (Baumann, B. 18, 887). Liquid.

Di-phenyl ether CMe, (SPh), v. Di-

PHENYL-PROPYLIDENE DISULPHIDE.

THIOACETONURAMIC ACID v. ACETONYL THIOCARBAMATE.

THIO - ACETOPHENONE C.H. CS.CH. [119.5°]. Formed by the action of alcoholic ammonium sulphide on acetophenone (Engler, B. 11, 930). Colourless leaflets or needles, insol. water, sl. sol. alcohol, v. sol. ether. May be sublimed.

THIO-ACETYL-TOLUIDINE v. Toluide of

THIO-ACETIC ACID

THIO-ALLOPHANIC ACID. Ethyl ether C₄H₈N₂SO₂ i.e. NH₂.CO.NH.CO.SEt. [180°]. Formed by heating urea with Cl.CO.SEt at 80°-90° (Peitzsch a. Salomon, J. pr. [2] 7,477). Pearly needles (from water), v. sl. sol. cold water and alcohol. Aqueous NH, at 100° forms biuret and mercaptan. Boiling baryta-water gives allophanic acid and mercaptan. Aniline yields di-phenyl-biuret and mercaptan.

Isoamyl ether NH2.CO.NH.CO.SC.H11. [176°]. Needles (Schöne, J. pr. [2] 32, 251).

AcCl yields an acetyl derivative [85°].

Phenyl ether [218°] (Gattermann, A. 244, 43).

Isothio-allophanic ether NH, CS.NH.CO, Et. [170°-175°]. Formed by the action of thio-urea on COCl.CO, Et (Peitzsch, B. 7, 896). Prisms (from

Di-thio-allophanic ether NH, CS.NH.CO.SEt. Formed from KSCy, alcohol, and HCl (Blankenhorn, J. pr. [2] 16, 358). Pearly prisms, insol. cold water, sol. hot ether. Alcoholic NH₃ at 150° forms thio-urea. Baryta-water yields mercaptan, CO2, and thio-urea

DI-THIO-AMMELIDE C₃H₄N₄S₂ i.e. C₃N₃(NH₂)(SH)₂. S. 3 at 100°. Formed by boiling pseudosulphocyanogen C.HN.S. with KSH or ammonium sulphide solution (Jamieson, A. 59, 339; Ponomareff, J. R. 8, 222). Small needles, nearly insol. cold water, alcohol and ether, v. sol. aqueous ammonium sulphocyanide. Acid in reaction, decomposing carbonates.

Reactions.-1. Yields cyanuric acid when heated with HClAq. - 2. NH3Aq forms melamine

C_sH_gN_g by heating in a sealed tube.

Salts.—NaHA"1½ aq. Tables.—KHA"2aq. White prisms, v. e. sol. water and alcohol.—MgH₂A"₂6aq.—CaH₂A"₂2aq.—BaH₂A"₂5aq.—BaH₂A"₂6aq. Monoclinic crystals, sl. sol. cold

water.—SrH, A", 4aq.—AgHA". Flocculent pp., insol. water.

Methyl ether C2N8(NH2)(SMe)2 Formed by heating C, N, (SMe), with alcoholic NH, for five hours at 100° (Hofmann, B. 18, 2756). Trimetric tables, v. sl. sol. water, v. sol. alcohol. Boiling conc. HClAq forms MeSH, cyanuric acid. and NH_s.—B'HAuCl_s. Needles.

Ethyl ether Et_sA". [112°]. Formed by

heating tri-thio-cyanuric ether with alcohol NH. at 180° (Klason, J. pr. [2] 33, 298). Trimetric prisms. Converted by acids into ammelide.

Isoamylether $(C_5H_{11})_2A''$. [82°]. Di-thio-methyl ammelide di-methyl ether Cy₃(NHMe)(SMe)₂. [175°]. Formed by heating Cy₃(SMe)₃ with aqueous NMeH₂ (Hofmann, B. 18, 2761). Needles or prisms (from dilute alcohol).

THIO-AMMELINE C.H.N.S i.e.

C(SH) N.C(NH₂) N. S. .77 at 100° (K.). Formed by adding HCl to an aqueous solution of dicyan-diamide and ammonium sulphocyanide at 100° (Rathke, B. 18, 3106; 20, 1059; 23, 1675). Formed also from C_sN_s(NH₂)₂Cl and KSH (Klason, J. pr. [2] 33, 296) and by heating ψ-sulphocyanogen with cone NH Aq at 100° (Ponomareff, C. R. 80, 1384). Needles, sl. sol. hot water, v. sol. acids and alkalis. Yields a canary-yellow pp. on boiling with ammoniacal CuSO4. Conc. HClAq at 130° yields cyanuric acid, H₂S, and NH₈. Ethylene bromide forms crystalline Conc. NH, Aq at 200° forms melamine. Boiling NaOHAq yields ammeline. Ammoniacal silver nitrate ppts. C.H.N.SAg and C₂H₂N₅SAg₂. Bromine added to a solution of thio-ammeline in HBrAq forms S₂(C₃N₃(NH₂)₂)₂, which dissolves in alkalis, and is split up by boiling with acids into ammeline, thio-ammeline, and S.

Salts.—B'HNO_s. Needles.—B'₂H₂SO₄8aq -B'₂H₂C₂O₄aq. Needles, v. sl. sol. cold water. Needles. -B'2H2SO48aq.

Methyl ether MeA'. [268°]. Formed by heating tri-methyl tri-thiocyanurate for five hours with conc. alcoholic NH, at 160° (Hofmann, B. 18, 2757). Tables (from water).—B'₂H₂PtCl₂.

Ethylether EtA'. [165°].

Isoamylether C₃H₁₁A'. [178°].

Reference. - TRI-PHENYL-THIO-AMMELINE.

THIO-ANILINE v. DI-AMIDO-DI-PHENYL SUL-

THIOANISOIC ACID (so called) C. H. SO. Got by boiling anethol with HNO, (S.G. 1.1), distilling the product and shaking the fraction 215°-245° with conc. NaHSO, Aq and some alcohol (Städeler a. Wächter, A. 116, 163). Crystalline (containing 2aq), v. e. sol. water, alcohol, and ether.—NH, A'aq.—NaA'aq. S. 15 in the cold.— BaA', 3aq. S. 8.5 in the cold.—CaA', 2aq.—MgA', 5aq.—AgA'. Plates, m. sol. water.

THIORENZAMIDE v. Amide of THIOBENZOIC

(a)-THIO-BENZOIC ACID CaHs. CO.SH. Mol.

w. 138. [c. 24°].

Formation.-1. From BzCl and alcoholic K₂S (Engelhardt a. Latschinoff, Z. [2] 4, 353).-2. By boiling EtOBz, PhOBz, and Bz₂O with alcoholic KSH.

Properties .- Yellow oil or white crystalline mass with unpleasant smell, volatile with steam. Decomposed by distillation. V. sol. alcohol and ether. Its solutions, when exposed to air, deposit crystals of Bz₂S₂. Nitric acid also oxidises it to benzoyl disulphide. A neutral solution

gives with CuSO₄ a greenish-yellow pp., turning bright red; the pp. then containing Bz₂S₂.
Salts.—KA'. Large prisms (from alcohol), v. sol. water.—NaA'.—BaA'₂xaq. Laminæ (from alcohol).—PbA'₂.—AgA'. Yellowish-white pp.

Methyl ether C₂H₂.COSMe. (232°).

Formed from Pb(SMe), and BzCl (Obermeyer, B.

20, 2922).

Ethyl ether EtA'. (243°). Formed by the action of BzCl on Pb(SEt), in presence of ether (Tütscheff, Petersb. Acad. Bull. 5, 295), and got also by boiling AgA' and EtI, and from PhOBz and NaSEt (Seiffert, J. pr. [2] 31, 471). Oil, with unpleasant smell, sol. alcohol and ether. Boiling alcoholic KOH forms KOBz and KSEt. Alcoholic KSH yields KSBz and KSEt. Oxidised by AgMnO, to benzoic and ethane sulphonic acids

(Beckmann, J. pr. [2] 17, 464).

Isoamyl ether C.H., A'. (271°).

Phenyl ether Phs.Bz. [56°]. Formed from phenyl mercaptan and BzCl (Schiller a. Otto, B. 9, 1635). Needles (from benzene).

Di-nitro-phenyl ether C,H3(NO2)2A'. [113°]. Formed from (a)-chloro-m-di-nitro-benzene, alcoholic KSH, and BzCl (Willgerodt, B. 18, 328). Needles (from alcohol).

Benzyl ether C,H,A'. [39.5°]. Formed from benzyl mercaptan and BzCl (Otto a. Lüders, Triclinic crystals, yielding B. 13, 1285). PhCH. SO H on oxidation.

p-Tolyl ether C,H,A'. [75°] (S. a. O.). Anhydride v. BENZOYL SULPHIDE.

(B)-Thio-benzoic acid CaHa.CS.OH. Formed by boiling thiobenzoic aldehyde with nitric acid (S.G. 1.3) (Fleischer, A. 140, 234). Needles (containing aq), sol. alcohol and benzene. More sol. hot water than benzoic acid. When heated strongly it blackens without melting.—BaA', 24aq. Small nodules, v. sol. water and ppd. by alcohol.

Amide C. H. CS.NH.. Thiobenzamide. [116°]. Formed by passing H.S into an alcoholic solution of benzonitrile containing a little NH, and heating on a water-bath (Cahours, C. R. 27, 329; Bernthsen, B. 10, 1241; A. 192, 49; Gabriel a. Heymann, B. 23, 157). Formed also by heating benzylamine (2 mols.) with S (1 mol.) at 180° (Wallach, A. 259, 304). Long needles (from hot water). Decomposed by HgO, yielding HgS and benzonitrile. Sodium-amalgam acting on its alcoholic solution forms amorphous thiobenzoic aldehyde, benzonitrile, benzylamine, and benzoic aldehyde. Iodine added to its alcoholic solution reacts with formation of di-benzenyl-azo-sulphim

 $C_eH_s.C \stackrel{N.S}{\sim} C.C_eH_s$, crystallising in long

colourless needles [90°] converted by boiling H,SO, into a base C₁,H₁,N₂ [71°], which yields B'₂H₂PtCl_s (Hofmann, B. 2, 645; B. 25, 1587; Wanstrat, B. 6, 335). Zinc and HCl reduce thiobenzamide in alcoholic solution to benzylamine. C,H,.C(NOH).NH, Hydroxylamine forms Ethylene bromide at 100° forms the compound C₂H₄(S.C(NH)C₆H₈)₂2HBr [233°] decomposed by boiling water into C₂H₄(SBz)₂ (Gabriel a. Heymann, B. 24, 783). Chloral (1 mol.) forms, on warming, a compound C₂H₃Cl₂NSO or CaHs.CS.NH.CH(OH).CCl. (?) crystallising

silky plates [104°], sl. sol. water, v. sol. alcohol (Spica, G. 16, 182).

Anilide C.H. CS.NHPhor C.H. C(SH):NPh. Thiobenzanilide. [102°]. Formed by the action of P.S. (1 pt.) on benzanilide (2 pts.), and by heating phenyl-benzamidine or s-di-phenyl-benzamidine OPh(NPh)(NPhH) with H₂S or with CS₂ (Bernthsen, B. 11, 503; A. 192, 31). Formed also by passing H,S into a solution of C.H. CCl:NPh in benzene (Leo, B. 10, 2133), and from benzophenone oxim and P2S3 (Dodge, A. 264, 184). Obtained by heating benzyl-aniline (1 mol.) with S (2 at.) at 220° as long as H₂S escapes (Wallach, A. 259, 301). Thin prisms (from EtOAc), nearly insol. boiling water, v. sol. alcohol and ether. V. sol. KOHAq. Converted into benzanilide by heating with dry PbO or with alcoholic potash at 150°. Yields benzenylo-amido-phenyl-mercaptan on dry distillation, on heating with S, and also on treatment in alkaline solution with K_sFeCy₆ (Jacobsen, B. 19, 1068). Hydroxylamine yields C₆H₅, C(NOH).NHPh.

o-Toluide C.H., CS.NHC, H.Me [1:2]. [86°]. Formed by melting BzNHC, H.Me with P.S. (Stieglitz, B. 22, 3160). Yellow, six-sided prisms (from benzene). Hydroxylamine in alcohol forms, on heating, C.H. C(NOH).NHC,H, [147°], crys-

tallising in needles.

p-Toluide. [129°]. Formed in like manner (Müller, B. 22, 2405), and also by the action of H₂S on C₆H₅.CCl:NC,H, (Leo, B. 10, 2134; Pfitzinger a. Gattermann, B. 22, 1065), and by heating p-tolyl-benzamidine with CS, (Bernthsen a. Trompetter, B. 11, 1759). Long yellow needles, insol. water, v. sol. alcohol, ether, and NaOHAq.

Xylide C₈H₈.CS.NHC₈H₈Me₂. [90°]. Formed by heating the benzoyl derivative of (4,2,1)xylidine with P₂S₃ (Gudeman, B. 21, 2552). Small needles (from alcohol). Oxidised by

 K_s FeCy, to oily $C_sH_2Me_2 < N > CPh$.

Di-phenyl-amide C,H,.CS.NPh2. [151]. Triclinic crystals (Bernthsen, A. 192, 37).

(a) Naphthalide v. vol. iii. p. 474. Di-thio-benzoic Acid C,H,.CS.SH. Formed by adding an alcoholic solution of K.S to C.H., CCl. (Engelhardt a. Latschinoff, Z. [2] 4, 455; Klinger, B. 15, 862). Heavy, red oil, v. sol. alcohol and ether. Its dilute ethereal solution is crimson. Resinifies when exposed to air. -PbA'2. Thin orange-red needles (from alcohol or xylene).—HgA'2. Golden plates (from alcohol). -AgA'. Brown pp., insol. water and benzene.

References. - OXY-THIO-BENZOIC ACID and AMIDO-THIO-BENZAMIDE.

(a)-THIO-BENZOIC ALDEHYDE C_eH_s .CHS. Benzylidene sulphide. Formed by passing H_2S into an alcoholic solution of benzoic aldehyde (Laurent, A. Ch. [3] 1, 292; Klinger, B. 9, 1895; 15, 863; Böttinger, B. 12, 1056), or of hydrobenzamide (Cahours, C. R. 25, 457). White amorphous powder, softening at 85°, decomposed at a higher temperature. Insol. water and cold alcohol, v. e. sol. benzene and chloro-

form. Does not combine with KHSO, or HCy. Potash-fusion yields benzyl mercaptan. heating with copper it forms CuS and s-di-phenylethylene. Alcoholic KSH on heating forms. Ph.CS.SK and di-benzyl disulphide. Thioglycollic acid forms C_eH_a.CH(S.CH₂.CO₂H)₂ [124°] |

(Bongartz, B. 21, 479).

(8)-Thio-benzoic aldehyde (C.H.OHS)x. [225°]. Formed from the (a)- isomeride by treatment with AcCl, EtI, or a little iodine in benzene (Klinger, B. 10, 1877). Prepared by passing H2S into a mixture of benzoic aldehyde and alcoholic HCl (Baumann a. Fromm, B. 22, 2604). Needles. Crystallises from benzene as (C,H,S),C,H,, and gives off its benzene at 130°-140°. V. sol. hot HOAc, m. sol. alcohol. On heating with copper-powder it yields CuS and di-phenyl-ethylene. An oily compound $(C_1H_0S)_2H_2S$ is formed, together with benzyl disulphide and di-thio-benzoic acid, by the action of alcoholic KSH on benzylidene chloride (Klinger, B. 15, 864). This compound yields benzoic acid on treatment with dilute HNO,

 (γ) -Thio-benzoic aldehyde C_sH_sOHS . [167°]. Occurs in the benzene mother-liquor in the preparation of the (β) - isomeride (B. a. F.). Small pointed needles, much more sol. benzene and chloroform than the (B)-modification, sl. sol. alcohol and ether. Its crystals do not contain benzene of crystallisation. By adding iodine to its solution in benzene it is changed to the

(β)- isomeride.

Thio-benzoic orthaldehyde CeH5.CH(SH)2. Methyl ether C.H. CH(SMe)2. Formed by passing HCl into a mixture of benzoic aldehyde and MeSH (Bongartz, B. 21, 487). Oil. Oxi-

dised by KMnO, to C.H. CH(SO CH.), [163°].

Ethyl ether C.H. CH(SEt). Oil (Bau-

mann, B. 18, 885).

p-Bromophenyl ether C_eH_s.CH(SC_eH_sBr)₂. [80°]. Silky needles (Bau-

mann, B. 18, 885). Reference.—NITRO-THIO-BENZOIC ALDEHYDE.

THIO-BENZOPHENONE C,H,.CS.C,H,.

Formed by heating CSCl₂ (5g.) with benzene (25 g.) and AlCl, (9 g.) (Bergreen, B. 21, 337). Reddish-brown oil, v. sol. ether, benzene, and hot alcohol. Decomposed by distillation. acts with hydroxylamine, forming Ph2C:NOH and with phenyl-hydrazine, with production of CPh₂:N₂HPh.

Thio - benzophenone (CSPh₂)_x. [146·5°]. Formed from CCl₂Ph₂ and K₂S (Engler, B. 11, 922). Small white needles. Converted by CrO, and HOAc into benzophenone. Does not react with hydroxylamine or phenyl-hydrazine

THIO-BENZOYL-(a)-NAPHTHYLAMINE v.

vol. iii. p. 474.

THIOBENZPINACONE C₂₄H₂₅S₂ i.e.

CPh₂H.S.S.CPh₂H.

Or CPh₂H.S.S.CPh₂H. [151°]. Formed from benzophenone by the action of alcoholic ammonium sulphide (Engler, B. 11, 922) or P₂S₅ (Japp a. Roschen, C. J. 49, 479), and, together with thiobenzophenone, by the action of KSH on C.H. CCl. C.H. (Behr, B. 5, 970). Formed also by the action of alcoholic KSH on (C₆H₅)₂CHCl, and of P₂S₅ on di-phenyl-carbinol. Slender needles (from alcohol), v. sol. CS2. In alcoholic solution it is converted by finely-divided copper into tetra-phenyl - ethane CHPh. CHPh.

THIO-BIURET C.H.N.OS. Formed by boil-carbimido-urea ('amidodicyanic acid') with ammonium sulphide (Wunderlich, B. 19, 452). Needles (containing aq). Sol. hot water and alkalis, v. sl. sol. ether. Gives a white pp

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with copper salts. By ammoniacal AgNO, it is converted back into carbimido-urea.

THIO-BUTYRIC ACID C, H, SO i.e.

CH₂.CH₂.CH₂.CO.SH. (130°). Formed by distilling butyric acid with P.S. (Ulrich, A. 109, 280). Liquid with disgusting smell, v. sl. sol. water, v. sol. alcohol.—PbA'2. Small crystals (from hot water).

Thio-isobutyric acid. Methyl ether C₂H₁₀SO *i.s.* Pr.CO.SMe. (c. 142°). V.D. 57·5. Formed from Pb(SMe)₂ and isobutyryl chloride (Obermeyer, B. 20, 2922). Oil. THIO-ISOBUTYRIC ALDEHYDE PrCHS.

(70°-90°). A product of the action of S on isobutyric aldehyde at 180° (Barbaglia, G. 18,

85). Oil.

(a)-THIO-CARBAMIC ACID NH2.CO.SH. This acid, set free by adding acids to a solution of its ammonium salt, splits up at once into COS and ammonia. The ammonium salt NH2.CO.SNH4 is prepared by passing gaseous COS into alcoholic N \hat{H}_s (Berthelot, J. 1868, 160) or into N H_s Aq at 0° (Schmidt, B. 10, 191; Kretzschmar, J. pr. [2] 7, 471). It forms colourless crystals, v. e. sol. water, m. sol. alcohol, insol. ether. The dry salt becomes yellow in air, forming ammonium sulphocyanide. The dry salt forms thio-urea when heated in a sealed tube at 135°. Thio-urea is also formed when Pb(OH)₂ is added to its cold aqueous solution. HgO forms, in the cold, ammonium cyanate (Fleischer, B. 9, 438). Water at 100° forms NH₄SH and (NH₄)HCO₃. FeCl₃ gives a red liquid, and finally a red pp. (Mulder, A. 168, 228).

Methyl ether NH...CO.SMe. [c. 98°] Formed, together with a small quantity of the (β) isomeride, by adding HCl to a solution of potassium sulphocyanide in boiling MeOH (Blankenhorn, J. pr. [2] 16, 375). Monoclinic prisms (from ether). Decomposed by alcoholic NH, into MeSH and urea. Aniline at 100° forms MeSH, s-di-phenyl-urea and NH₃.

Ethyl ether NH, CO.SEt. [102°] (Pinner, B. 14, 1083); [108°] (F.). Formed in like manner (B.) and also by passing NH_a into Cl.CO.SEt (Salomon, J. pr. [2] 7, 256) and by the action of EtBr on NH₂ CO.SNH₄ (Fleischer, B. 9, 991). Plates, v. sol. hot water, alcohol, and ether. Decomposed at 150° into mercaptan and cyanuric acid. P2O5 converts it into ethyl sulphocyanide.

Isoamyl ether CO(NH₂).SC₃H₁₁. [107°]. From CO(SC₃H₁₁)Cl and NH₃ (Schöne, J. pr. [2] 32, 247). Glittering plates (from ether). Sol. hot water, but decomposed by boiling water. Gives amorphous pps., insol. water, alcohol, and ether with AgNO₃, HgCl₂ and PtCl₄.

Reactions.-Alcoholic NH, forms urea and amyl mercaptan. - 2. Alcoholic KOH forms K₂CO₂, NH₂ and amyl mercaptan.—3. Heated with aniling it forms NH₂, HSC₂H₁₁, and diphenyl-urea.

(\$)-Thio-carbamic acid NH2.CS.OH.

Methyl ether NH, CS.OMe. [43°] . Formed by the action of alcoholic NH, on MeO.CS.SMe or on MeO.CS.SEt (Salomon, J. pr. [2] 8, 115).

Ethylether NH, CS.OEt. Xanthogenamide.

[38°]. Formed by the action of NH, on xanthogenic ether EtS.CS.OEt, on MeS.CS.OEt, and on (EtO.CS)₂S₂ (Debus, A. 72, 1; 75, 121; 82, 258; Chancel, J. 1851, 513; Salomon a. Maintz, J. pr. [2] 8, 114). Monoclinic prisms, sl. sol. water, miscible with alcohol and ether. Decomposed on distillation into mercaptan and cyanic acid. Boiling KOHAq resolves it into alcohol and potassium sulphocyanide. P₂O₅ forms ethyl sulphocyanide (Conrad, *J. pr.* [2] 10, 34). Nitrous acid passed into water containing xanthogenamide in solution forms crystalline $C_0H_{10}N_4SO_2$. CuSO, followed by HClAq ppts. $C_3H_1NOSCuCl$ as a white crystalline powder, nearly insol. water, sol. hot alcohol. Compounds of CuCl with 2, 8, and 4 mols of xanthogenamide may be obtained. KI added to a boiling alcoholic solution of (C3H7NOS)3CuCl (C₃H,NOS)₂CuI crystals of (C₃H₇NOS)₃CuI. Compounds (C₃H₇NOS)₂3CuSCy and C, H, NOSCuSCy may also be prepared (Debus). Isovaleric aldehyde in presence of HCl and alcohol forms C,Ho.CH(NH.CS.OEt),[108°] (Bischoff, B. 7, 1083). B - Iodo - propionic acid forms NH₂.CO.S.CH₂.CH₂.CO₂H [147·5°], which yields CaA'₂ Saq, BaA', 2aq, and Ag₂A" (Langlet, B. 24, 3849). B-Iodopropionic acid and Ac₂O yield sinapane propionic acid $CO < S - CH_2 > CH_2$ [159°], v. sol. hot water, converted by HgO into the mercaptide of β -thiolactic acid.

Isobutyl ether NH₂CS.OCH₂Pr. [36°].
Tables (from alcohol or ether) (Mylius, B. 5, 976; Blankenhorn, J. pr. [2] 16, 380).
Isoamyl ether NH₂CS.OC₃H₁₁. Oil

(Johnson, Č. J. 5. 242).

Di-thio-carbanic acid NH₂.CS.SH. Mol. w. 93. The ammonium salt NH₂.CS.SNH₄ is formed by passing NH₂ (from 150 pts. NH₂Cl) into 95 p.c. alcohol (600 pts.) containing CS₂ (96 pts.) (Mulder a. Bettink, J. pr. 103, 178; cf. Zeise, A. 48, 95; Debus, A. 78, 26). It is also formed by heating (NH4)2CS3. On gradually adding HClAq to a cooled conc. solution of the ammonium salt the free acid separates as colourless needles, v. sol. water, alcohol, and ether; acid in reaction. An alcoholic solution of the acid gradually decomposes into CS₂ and the NH, salt. An aqueous solution of the acid decomposes on heating into H2S and HCyS. The di-thio-carbamates also split at 100° into H.S and sulphocyanides. An alcoholic solution of iodine decomposes NH₂.CS.SNH₄, forming EtNCS and EtNH, (Hofmann, Z. [2] 5, 671). Ammonic di-thio-carbamate treated with HCl and solution of I gives crystals of the bisulphide S₂(CS.NH₂)₂ (Klason, J. pr. [2] 36, 62). FeCl₂ added to a solution of the NH, salt gives a black pp. turned white by excess. AgNOs gives a yellow pp., turning black. A solution of aldehyde forms carbothialdine NH2.CS.SN(C2H4)2. Benzoic aldehyde, isovaleric aldehyde, and acrolein form analogous bodies.

Salts .- NH, A'. Deliquescent lemon-yellow prisms. On heating with KOHAq is forms K.S. KSCy, and NH₃.—CuA'₂. Yellow powder, insol. water.—PbA'₂. White pp., blackening on boiling with water.—ZnA'₂: white pp.

Ethyl ether NH₂.CS.SEt. Mol. w. 121.

[42°]. Formed by passing H₂S under pressure

into ethyl sulphocyanide at 100° (Jeanjean, J. 1866, 501; Salomon a. Conrad, J. pr. [2] 10, 29). Trimetric crystals (from ether), insol. water, v. e. sol. alcohol. Alcoholic NH₂ or KOHAq in the cold forms mercaptan and ammonium sulphocyanide.

Isopropyl ether NH.CS.SPr. Trimetric plates (Gerlich, A. 178, 82).

Acetyl derivative of the ethyl ether NHAc.CS.SEt. [128°]. Formed by heating thio-acetic acid with ethyl sulphocyanide (Chanlaroff, B. 15, 1987). Yellow needles (from alcohol), v. sol. hot water. Decomposed by distillation into thio-acetic acid and ethyl sulphocyanide. Boiling baryta-water forms mercaptan and acetic acid. Hot dilute HClAq forms NH2CS.SEt.

Isothiocarbamic ether v. DI-ETHYL-(a)-THIO-CARBONATE. A mide.

THIO-CARBAMIDE v. THIO-UREA.

THIOCARBAMYL SULPHIDE C2H, N2S, i.e. (NH₂.CS)₂S (?) Very unstable oil, got by adding HCl to its ammonium salt. Quickly decomposes into CS₂, sulphur, H₂S, and ammonium sulphocyanide. The ammonium salt Supproblement. The alm of the action of CS₂ on alcoholic NH₃ in the cold (Zeise, B. J. 4, 98; Hlasiwetz a. Kachler, A. 166, 137). It forms colourless prisms, v. sol. water, m. sol. alcohol, sl. sol. ether. The moist salt slowly decomposes into H2S and ammonium sulphocyanide. The cupric salt CuC2H2N2S2 is canary-yellow.

Thiocarbamyl disulphide C.H.N.S. i.e. (NH₂.CS)₂S₂. Formed by the action of Cl on a solution of the ammonium salt of the preceding body, and on ammonium di-thio-carbamate (Debus, A. 73, 27). Pearly plates, v. sol. hot alcohol, insol. water. Decomposed by boiling water into CS2, ammonium sulphocyanide, and S.

THIOCARBIMIDES. Mustard oils. Compounds RN:CS. Essential oil of mustard is allyl thiocarbimide. They are formed by the action of CSCl₂ on primary amines (Rathke, A. 167, 218), and by the action of aqueous HgCl₂ on the dithiocarbamates RNH.CS.SNH₂R obtained by heating primary amines with CS2 and alcohol or ether (Hofmann, B. 1, 171; 2, 452; 7, 811; 8, 106; Rudneff, J. R. 10, 188). Methyl sulphocyanide is partially converted into methyl thiocarbimide by heating at 180°. Allyl sulphocyanide changes to the mustard oil on boiling. Thiocarbimides are also got by distilling alkyl cyanates with P2S5 (Michael a. Palmer, Am. 6, 258). The thiocarbimides are pungent oils. They are reduced by Zn and HClAq to amines and CH2S, whereas sulphocyanides yield HCy, mercaptans, and other products. Amines convert thiocarbimides into thio-ureas; thus aniline combines with phenyl thiocarbimide, forming s-di-phenyl-thio-urea. Alcoholic potash, or alcohol at 100° forms thio-carbamic ethers NHR.CS.OEt (Hofmann, B. 2, 117; Schiff, B. 9, 1316). Water at 200°, or HClAq at 100°, forms amine, CO₂, and H₂S, while with sulphocyanides it yields Et₂S, CO₂, and NH₃. Conc. H₂SO₄ forms amine and COS. HNO₃ forms amine and CO2; whereas with the isomeric sulphocyanides it forms a sulphonic acid. Hydroxylamine unites with thiocarbimides, forming oxy-thio-ureas NHR.CO.NH.OH (Tiemann, B. 22, 1939). Chlorine unites with PhN:CS, forming unstable (PhNCS)2Cl2, whence boiling water

forms (PhNCS),0 [118°], crystallising in yellow

needles (Helmers, B. 20, 786).

THIOCARBIMIDO-ACETIC ACID C.H.NSO. i.e. CH2(N:CS).CO2H. Formed by boiling thiohydantoin with HClAq (Volhard, J. pr. [2] 9, 6), and by boiling isoamyl sulphocyano-acetate with fuming HClAq (Claësson, B. 10, 1352; Heintz, A. 136, 232). Trimetric plates, v. e. sol. hot water. Oxidised by HNO, to H2SO, and oxalic acid. Weak acid.—BaA'2 aq. Four-sided prisms, sl. sol. cold water. Benzoic aldehyde and NaOHAq (1 mol.) react, forming the compound CHPh: $C(CO_2Na)$.S.CO.NH₂, which crystallises with $1\frac{1}{2}$ aq and is converted by HCl into CO.S $_{
m NH.CO}^{
m VO.S}$ C:CHPh [242°] (Andreasch, M. 10, 73).

THIOCARBIMIDO-BENZOIC ACID v. m-AMIDO-BENZOIC ACID

THIOCARBONIC ACID.

Thiocarbonates v. vol. i. p. 703. The thiocarbonic ethers that have not been treated of in former articles are described below.

Methyl thiocarbonates v. vol. iii. p. 399. Methyl CO(SMe)2. iso-dithiocarbonate (169°). Formed by warming methyl sulphocyanide with H₂SO₄ (Schmitt a. Glutz, B. 1,

166).

Ethyl thiocarbonates v. vol. ii. p. 520. Ethylene thiocarbonates v. vol. ii. p. 493.

Ethylene di-thio-di-carbonate di-ethyl C₂H_{*}(S.CO_{*}Et)₂. Formed by heating C₂H_{*}Br_{*} with KS.CO_{*}Et in alcohol (Welde, *J. pr.* [2] 15, 52). Thick oil, converted by alcoholic potash into C₂H₄(SH)₂ and KO.CO₂Et.
Isobutyl di-thio-carbonic acid CH₂Pr.O.CS₂H.

Salts.-NaA'. Formed from C.H.ONa and CS2. Yellowish white needles, v. e. sol. water and alcohol.—KA'. Needles (from alcohol). S.G. 15 1:371 (Clarke, B. 11, 1505). Chlorine gas passed into its aqueous solution forms oily (CH2Pro.CS)2S2 (Mylius, B. 5, 976).

Ethyl ether EtA'. (228°).

CH2Pr.O.CS.SCH2Pr. Isobutylether (249°). S.G. 12 1.009 (Mylius, B. 5, 975)

Isoamyl ether CH₂Pr.O.CS.SC₅H₁₁. (265°-270°) (Mylius). Isoamyl di-thio-carbonic acid

C.H., O.CS.SH. Oil, with unpleasant smell. Salts.—KA. Formed from fusel oil, KOH, and CS₂ (Balard, A. Ch. [3] 12, 807; Erdmann, J. pr. 31, 4; Desains, A. Ch. [3] 20, 505; Johnson, C. J. 5, 142). Gives a lemon-yellow pp. with CuSO₁.—PbA'₂. Plates. Iodine forms oily (C₅H₁₁O.CS)₂S₂.
Di-isoamyl s-di-thio-carbonate CO(SC₅H₁₁)₂.

(281°) (Schmitt a. Glutz).

Cetyl di-thiocarbonate v. vol. i. p. 728.

Iso-butyl-tri-thio-carbonic acid C.H.S.CS.H. Formed from C4H₂SNa and CS₂ (Mylius, B. 6, 316). The Na salt crystallises in yellow needles, v. sol. alcohol and ether.

Isobutyl ether (C,H,S),CS. (287°). Got by heating aqueous K₂CS₃ with isobutyl iodide.

Yellow oil with faint odour.

Di-isoamyl-tri-thiocarbonate $(C_aH_{11}S)_2CS.$ (247°). S.G. 88 (Husemann, A. 126, 297). Oil, with unpleasant smell.

 $(C_3H_5S)_2CS$ tri-thio-carbonate Di-allyl S.G. 94. Formed from allyl **∢170°−175°).** todide and Na,CS, in the cold (Husemann, A. 126, 269). Pungent yellow oil, with offensive odour.

THIOCARBONYL - ACETOACETIC ETHER CS:CAc.CO.Et. [156°-162°]. Formed from thio-carbonyl chloride and acetoacetic ether (Bergreen, B. 21, 847). Yellow needles, sl. sol. ether and ligroïn.

THIO-CARBONYL-DI-AMIDO-DI-BENZOIC

ACID v. m-AMIDO-BENZOIC ACID.

THIOCARBONYL - BENZOYL - ACETIC ETHER CS:CBz.CO₂Et. [164°]. Formed from CSCl₂ and benzoylacetic ether (Bergreen, B. 21, 351). Yellow needles, sol. alcohol-chloroform.

THIO-CARBONYL CHLORIDE v. Sulpho-

chloride of CARBON.

Thio-carbonyl tetrachloride v. PER-CHLORO-METHYL-MERCAPTAN.

THIO - CARBONYL - ETHYL - o - AMIDO-PHENOL $C_6H_4 < NEt_0 > CS.$ [112°]. 300°). Formed from ethyl-o-amido-phenol and CSCl₂ (Seidel, *J. pr.* [2] 42, 449, 457). Insol. water and dilute NaOHAq, m. sol. cold alcohol. HClAq at 170° forms ethyl-amido-phenol, CO. and H_2S . PCl_s yields $C_0H_4 < N > CCl$.

THIO - CARBONYL - METHYL - AMIDO-PHENOL $C_6H_4 < NM_{\odot} > CS.$ [128°]. 300°). Formed by the action of CSCl, on methylo-amido-phenol (Seidel, J. pr. [2] 42, 452). Needles, insol. water, sl. sol. cold alcohol.

THIO - CARBONYL - DI - (β) - NAPHTHYL -THIO - UREA $C_{10}H_7.N < \frac{CS}{C(NC_{10}H_7)} > 8.$ [152°]. Formed from di-(β)-naphthyl-thio-urea and CSCl₂ in benzene (Freund a. Wolf, B. 25, 1466). Stellate groups of needles, m. sol. hot benzene. Melts at 152° with evolution of gas, forming a solid melting at 224°

THIO-CARBONYL-DI-p-TOLYL-THIO-UREA $C_1H_1.N:C < \frac{N(C_1H_1)}{S} > CS.$ [109°]. Formed from di-p-tolyl-thio-urea [172°] and CSCl, in ether (Freund a. Wolf, B. 25, 1465). Yellowish needles, v. sol. hot alcohol and ether. Converted by HgO into carbonyl-di-tolyl-thio-urea [112°].

THIOCHRONIC' C,H,S,O,, ACID C_s(OH)(SO₄H)(SO₈H)₄. Formed by adding tetrachloro-quinone to a conc. solution of K.SO. (Hesse, A. 114, 313; Graebe, A. 146, 40). Yellow trimetric prisms, v. sol. hot water, insol. alcohol. Coloured brownish-red by FeCl. Reduces AgNO. Converted into hydroquinone disulphonic acid by heating with water at 135°, or by boiling with HClAq.

Euthiochronic acid C.H.S.O. i.e. C_e(OH)₂O₂(SO₂H)₂. Di-oxy-quinone disulphonic acid. Formed by boiling thiochronic acid with KOHAq. Deliquescent yellow needles, v. sol. water and alcohol. Reduced by tin and HClAq to $C_0H_0S_2O_{10}$, which yields $Na_2A'' 2aq$ and $K_2A'' 2aq$, which are re-oxidised in aqueous solution by air to euthiochronates.—Na,A'raq (dried at 150°).—K,A'r 2aq. Minute lemon - yellow prisms, v. sol. water.—K₂HA'vaq. Small orange-red four-sided prisms.—Ba₂A'v4aq. Ochre-yellow pp.—Ag,A'v: amorphous pp.

(a) - THIO - CINNAMIC ACID C.H.SO i.e. C.H.CH.CH.CO.SH. Formed from cinnamoyl chloride and Pb(SEt), the resulting oily ether

being boiled with alcoholic KSH (Engelhardt, Z. [2] 4, 859).—KA'. Crystalline.

(β)-Thio-cinnamic acid. Amids C₂H₇S.NH₂. [112°]. Formed by passing H₂S into a solution of cinnamic nitrile in alcoholic NH, (Rossum, Z. 1866, 362; Krüss, B. 17, 1768). Golden plates.
(a)-THIO-CINNAMIC ALDEHYDE (C.H.S).

[167°]. Formed by passing H2S into oil of cinnamon dissolved in alcoholic HCl (Baumann a Fromm, B. 24, 1452). Crystalline, v. e. sol. benzene, sl. sol. alcohol.

(β)-Thio-cinnamic aldehyde (C_9H_8S)₃. [213°]. Formed at the same time as the (a)-isomeride. Prisms, v. sl. sol. cold benzene and alcohol.

Thio - cinnamic orthaldehyde. Phenylether C.H.CH:CH:CH:CH(SPh), [81°]. Formed by passing dry HCl into a mixture of cinnamic aldehyde and phenyl mercaptan (Baumann, B. 18, 885). Needles (from ligroin).

p-Bromo-phenyl ether CHPh:CH.CH(S.C,H,Br)₂. [107°]. Needles (from

alcohol or ether).

Gly colly l'ether CHPh:CH.CH(S.CH, CO₂H), [143°]. Formed from cinnamic aldehyde and thioglycollic acid (Bongartz, B. 21, 481). Plates (from water). Zinc-dust in alkaline solution converts it into CHPh:CH.CH, S.CH, CO, H [77°], crystallising

from dilute alcohol in plates.

TRI-THIO-CITRIC ETHER C₃H₅O(CO.SEt)₂. Formed from tri-phenyl citrate and NaSEt in presence of ether (Seiffert, J. pr. [2] 31, 470).

Oil, smelling like mercaptan.

CH:CH THIO-COUMARIN C.H. . [101°].

Formed by heating equal parts of coumarin and P₂S₅ at 120°; the yield is 50 p.c. Also by heating o-coumario acid with P2S3. Long yellow needles, v. sol. alcohol, ether, and benzene; insol. water. Sublimable. By heating with aqueous alcoholic KOH it is reconverted into coumarin. Unlike coumarin it readily reacts with hydroxylamine and with phenyl-hydrazine (Tiemann, B. 19, 1661).

THIO-CRESOL v. TOLYL MERCAPTAN.
THIO-CROCONIC ACID C.H.2O.S, probably $SC < \stackrel{C}{<} \stackrel{C(OH).CO}{<}$. Formed by saturating a warm

solution of potassium croconate treated with 2 mols. of HCl, with H2S.—BaA" 2aq: brownishyellow amorphous pp., v. sol. dilute HCl, which after some time changes to reddish-brown crystals with violet reflex, insol. dilute HCl (Nietzki a. Benkiser, B. 19, 299).

THIO-CUMINIC ACID. Amide C10H13NS i.e. C₃H₁, C₆H₄, CS.NH₂. Formed by passing H₂S into a solution of cuminic nitrile in alcoholic NH₄ (Czumpelik, B. 2, 185). Needles, v. sol. hot alcohol. Converted in alcoholic solution by iodine into C₂₀H₂₂N₂S [45°] crystallising in prisms (Wanstrat, B. 6, 322)

THIODICYANDIAMIDINE v. GUANYL THIO-UREA

THIOCYANIC ACID v. SULPHOCYANIC ACID. THIOCYANO. v. SULPHOCYANO.

DI-THIO-CYANURIC ACID Cy. (SH)2(OH) aq. Preparation.—An alcoholic solution of KCyS is mixed with the calculated quantity of HCl, filtered from KCl, and evaporated at 40°. The residue is extracted with strong NH.Aq and BaCl, is added to the filtrate. The barium salt gradually separates in globular crystals (Klason, J. pr. [2] 33, 121). Yield, very small.

Properties.—White scales (from hot water). Sol. NaOHAq. A solution of the potassium salt gives pps. with AgNO₃, HgCl₂, and Pb(OAc)₂.

Reactions.—1. Conc. HCl at 180° gives cyanuric acid and H₂S.—2. Cold KMnO₄Aq and hot HNO, Aq form cyanuric acid.—3. Iodine forms (HOCy₈)₂S₄ as a white crystalline powder.

Salts.—KH₂A'''. Silvery mass of microscopic prisms.—BaHA'''2aq.

Tri-thio-cyanuric acid Cy₃(SH)₃. Formed by warming powdered C₅N₃Cl₂ (1 mol.) with conc. aqueous KSH (4 mols.) (Hofmann, B. 18, 2196; Klason, J. pr. [2] 33, 116). Minute prisms, when ppd. from dilute solutions; amorphous, from conc. solutions. Nearly insol. hot water, alcohol and ether. Not decomposed at 200°. FeCl, gives no colour in solutions of the acid, but in conc. solutions of the K salt it gives a white pp. and a yellow solution.

Reactions.-1. HClAq at 130° gives cyanuric acid and H₂S.—2. Alkaline KMnO₄ at 20° gives cyanuric acid.—3. Warm HNO₃ forms cyanuric acid.—4. At 360° it gives off CS₂ and some

HCyS and leaves melem $C_aH_aN_{10}$. Salts.— $K_aA'''3aq$.— $K_aH_aA'''_46aq$.— NaH_2A''' . Crystals, v. sol. water.—BaHA''' 3aq. Yellow crystals, ppd. by addition of BaCl₂ and NH₃Aq.—BaH₄A'''₂2aq. Prisms, m. sol. water.—SrHA''' 5aq. -CaHA" 5aq. Prisms, m. sol. water.

Methyl ether Me, A". [189°]. Got by heating methyl sulphocyanide with a few drops of HClAq at 100° (Hofmann, B.13, 1351; 18, 2197, 2755, 2774; 19, 2065). Hexagonal crystals. An alcoholic solution of ethylamine at 100° forms Cy_s(NHEt)(SMe)₂ [114°] while at 140° it yielde Cy_s(NHEt)₂SMe [84°]. Alcoholic NH, forms, in like manner, Cy₃(NH₂)(SMe)₂ and Cy₃(NH₂)₂SMe.

Ethyl ether Et, A". [27°]. (350°). Formed from Cy, Cl₃ and NaSEt. Colourless tablets (from HOAc) converted by alcoholic NH₃ at 180° into $\text{Cy}_{\text{s}}(\text{NH}_2)(\text{SEt})_2$ [112°] (crystallising in trimetric forms; a:b:c=669:1:391, v. sol. ether), accom-

panied by $Cy_*(NH_2)_2(SEt)$ [165°], which is insole ther (Klason, J. pr. [2] 33, 298). Amylether $(C_5H_{11})_3A'''$. Oil. Yields $Cy_*(NH_2)(SC_5H_{11})_2$ [82°] and $Cy_*(NH_2)_2SC_5H_{11}$ [178°].

Phenyl ether Ph.A". [97°]. Prisms (from HOAc).

p-Tolyl ether $(C_7H_7)_8A'''$.

THIO DI-ETHYL-ANILINE v. TETRA-ETHYL-

DI-AMIDO-DI-PHENYL SULPHIDE.

CH-CEt THIO-ETHYL-COUMARIN

[94°]. Formed by heating (a)-ethyl-coumarin with P₂S₅ at 120° (Aldringen, B. 24, 3462). Yellowish-red plates, sol. alcohol. May be sub-

THIO-ETHYLENE GLYCOL v. ETHYLENE MERCAPTAN.

THIO-FORMIC ACID H.CO.SH (?) [120°]. Formed by the action of H2S on lead formate (Wöhler, A. 91, 125; Limpricht, A. 97, 861), but not from P₂S_a and formic acid (Hurst, A. 126, 68). Small transparent crystals (from alcohol) with alliaceous odour, insol. water. May be sublimed. According to Nicol (Pr. E. 10, 425) a solution of thioformic acid, got from PbA'₂ and H₂S, rapidly decomposes, yielding formic acid.

Salts .- H.CO.SK, Formed from CHCl, and alcoholic K₂S (Nicol, T. E. 29, 531). Needles, sol. water and alcohol, nearly insol. dilute alcohol. AgNO, added to its aqueous solution forms a white pp., turning black on standing. HNOs yields S and formic acid. HgO also gives formic

acid.—(H.CO.S)₂Pb.

Anilide H.CS.NHPh. Thioformanilide. [137°]. Formed from dry phenyl carbamine and H_2S (Hofmann, B. 10, 1095, 1238). Formed also by passing dry H2S over di-phenyl-formamidine NPh:CH.NHPh at 150° (Bernthsen, A. 192, 35). Prepared by warming formamidine (5 pts.) with P_2S_3 (3 pts.), the yield being 60 p.c. of the theoretical (Hofmann, B. 11, 338). Thin plates (from water), partially decomposed on distillation into H2S and phenyl carbamine. Its solution tastes bitter. Decomposed by hot KOHAq into aniline, H2S, and formic acid. Heated in a sealed tube at 180° it yields $C_{14}H_{12}N_{2}S$ [140°] v. sol. alcohol (Nicol, B. 15, 211). NaOEt and EtBr sol. alcohol (Nicol, B. 15, 211). yield oily CH(SEt):NPh (c. 235°) (Wallach a. Wüsten, B. 16, 145)

o-Toluide H.CS.NHC,H, [96°]. Got by heating formyl-toluide with P.S. (Senier, C. J.

47, 762).

p-Toluide. Yellow needles. Xylide HCS.NHC₆H₃Me₂. [105°]. Small needles (from alcohol) (Gudeman, B. 21, 2549).

Thio-orthoformic acid. Ethyl ether CH(SEt)_s. Formed by boiling chloroform with aqueous NaSEt (Gabriel, B. 10, 186; Claesson, J. pr. [2] 15, 174). Oil, with nasty smell, partly decomposed on boiling. Oxidised by HNO₂ to ethane sulphonic acid. Fuming HClAq yields formic acid and mercaptan

Benzyl ether CH(SC,H,). [98°]. Formed from NaSC, H, and chloroform (Dennstedt, B.

11, 2265). White crystals.

Phenyl ether CH(SPh), [39.5°]. Trimetric tables. Oxidised by potassium permanganate to PhS.CH(SO_2Ph)₂ [176°] (Laves, B. 23, 1416; 25, 347).

Reference.—CHLOROFORMIC ACID.
THIO-FORMIC ALDEHYDES

Thio-formic paraldehyde (CH₂S)_s. Methyl-sulphide. Mol. w. 188. [216°]. V.D. 5.08 ene sulphide.

Formation.—1. By heating Na₂S with methylene iodide (Husemann, A. 126, 293). -2. By reducing CS2 with Zn and HClAq (Girard, A. 100, 806).—8. By the action of H2S on formic aldehyde or trioxymethylene.

Preparation.—By saturating a mixture of formic aldehyde (2 vols.) and conc. HClAq (4 vols.) with H_2S (Baumann, B.23, 67).

Properties.—Prisms, smelling like onions. Insol. cold water, sl. sol. alcohol and ether, m. sol. benzene and CS₂. May be sublimed. Oxidised by KMnO₄ to a trisulphone and also C,H,S,O, crystallising in colourless needles.-Forms: C.H.S.2AgNO, — C.H.S.AgNO, 2aq. — C.H.S.HgCl.—(C.H.S.), 2PtCl. — (C.H.S.), 2PtCl. — (C.H.S.), 2PtCl. (Hofmann, B. 2, 152; 3, 584; Girard, C. R. 70,

Thio-formic orthaldehyde.

Ethyl ether CH₂(SEt)₂. (184°) (C.); (180°) (F.). S.G. ²⁰ 987. Formed from CH₂I₂ and NaNEt (Claesson, J. pr. [2] 15, 176). Liquid | 124°, 146°, 148°, 128°, 180°, and 162° respec-

with unpleasant smell. Oxidised by HNO, to ethane sulphonic acid. KMnO, and H₂SO₄ yield CH₂(SO₂C₂H₂)₂ (Fromm, 4, 253, 155).

Thio-formic metaldehyde (CH₂S)_n. [176°]. Formed, as a white amorphous substance, on warming an aqueous or alcoholic solution of hexamethylene-amine saturated with H.S (Wohl, B. 19, 2344). Insol. most solvents. Decomposed when heated above its melting-point.

Di-thio-formic paraldehyde (C.A.s.o), aq. Dithioglyceric aldehyds. [82°]. (180°-185°). Formed by passing H2S into a solution obtained by electrolysis of dilute glycerin (Renard, A. Ch. [5] 17, 307). Amorphous waxy mass, sol. warm

water, insol. alcohol and ether.

(α)-THIO-FURFURALDEHYDE (C,H,SO). [128°]. Formed, together with the (β)-isomeride, by passing H2S into a solution of furfuraldehyde (10 g.) in alcohol (100 c.c.) and HClAq (20 c.c.) at -5° (Baumann a. Fromm, B. 24, 3593). Crystals, v. e. sol. alcohol and chloroform, insol. water.

(β)-Thio-furfuraldehyde (C₅H₄SO)₃. [229°]. Needles, v. sl. sol. alcohol, v. sol. chloroform.

Thio-furfuraldehyde $(C_4H_0SO)_n$ [n=20 to 24]. Mol. w. (by Raoult's method) 2182. Formed by mixing an alcoholic solution of furfuraldehyde with alcoholic ammonium sulphide (Cahours, A. 69,85; B.a.F.). White crystalline powder, softens at 80° and is melted at 91°. Thiofurfuraldehyde is decomposed by warm aniline, yielding furfuraldehyde and H_2S (Schiff, B. 19, 2153).

THIO-GLYCERIN v. GLYCERIN.

THIO-GLYCOL v. ETHYLENE MERCAPTAN. ACID CH2(SH).CO2H. THIO-GLYCOLLIC

Sulphydro-acetic acid.

Formation.—1. By reducing the compound ClSO2.CHCl.COCl with tin and HClAq (Siemens, B. 6, 659).—2. By adding OH₂Cl.CO₂H (1 mol.) to conc. KSHAq (2 mols.) (Claesson, A. 187, 113).—3. By the action of H₂S on glyoxylic acid in presence of Ag₂O (Böttinger, A. 198, 215).-4. By boiling thiohydantoin with water (Andreasch, B. 12, 1385).-5. Together with HSCy, by the decomposition of rhodanic acid C₄H₄NS₂O with baryta-water (Ginsberg a. Bondzynski, B. 19, 113).

Properties.—Oil, miscible with water, alcohol,

and ether. Decomposes when quickly heated. Very dilute FeCl, gives a fugitive indigo-blue colour, changed by a few drops of NH, Aq to dark red and then to violet. Excess of FeCl, oxidises it to di-sulphido-di-acetic acid S₂(CH₂.CO₂H)₂, which is also readily formed by atmospheric oxidation of the alkaline solution. A solution of the K salt added to CuSO, gives a bluish-black pp. reduced by further addition of the K salt to the white cuprous salt Cu₂(S.CH₂CO₂H)₂. HCl passed through a mixture of thioglycollic acid and acetoacetic ether gives rise to crystalline CO₂Et.OH₂.CMe(S.CH₂.CO₂H)₂ [96°] (Bongartz, B. 21, 485). Acetyl-propionic acid forms, in B. 21, 485). Acetyl-propionic acid forms, in like manner, CO₂H.CH₂.CH₂.CMe(S.CH₂.CO₂H). [154°]. Aldehydes react, forming the compound CRH(S.CH₂.CO₂H)₂, compounds which are produced either on standing, on warming, or on treatment with ZnCl₂or dry HCl (Bongartz, B.19, 1981; 21, 479). Such compounds obtained by using acetic, benzoic, cinnamic, o-oxy-benzoic, and o-, m-, and p-nitro-benzoic aldehydes melt at 108°,

tively. Thioglycollic acid reacts with ketones in presence of ZnCl, or HCl, forming acids ORB'(S.CH, CO,H),; such compounds obtained from acetone, acetophenone, and benzophenone

from acetone, acetophenone, and benzophenone melt at 135°, 139°, and 164° respectively.

Salts. — KHA"aq (from water). — KHA"
(from alcohol). Decomposes K₂CO₃, forming CH₂(SK).CO₂K.—BaA" 3aq: crystalline pp.—Ball, A"₂: gummy.—CdA".—Hg(S.CH₂.CO₂H)₂.

Needles or flat prisms, v. sol. hot water and alcohol.—Hg₂(S.CH₂.CO₂),BaH₂. Minute crystals.—Hg₃(S.CH₂.CO₂),AlH₃.—Hg₂MnH₂A"₄.—HgPbA"₂.—HgA".—PbA".—Bi(HA")₃. [c. 33°].

Yellow pp.—Cu₂H₂A"₂. White pp.—Cu₂BaA"₂.—Ag₂A",—Ag₂(NH₄)A"(NO₃).—Ag₂HgA"₂.

Ethul sther EtA' Formed by boiling

Ethyl ether EtA'. Formed by boiling thioglycollic acid with alcohol and a little H₂SO₄ (Claesson, A. 187, 116). Oil with nasty smell. Slowly decomposed by boiling into H₂S and S(CH2.CO2Et) Alcoholic HgCl2 ppts. ClHg.S.CH₂.CO₂Et, converted by excess of the ether into Hg(S.CH₂.CO₂Et)₂ (Wislicenus, A.

146, 145),

Ethyl derivative CH2(SEt).CO2H. Formed by mixing CH2Cl.CO2Et with NaSEt and alcohol (Claesson, B. 8, 120). Liquid, v. sol. alcohol and ether, m. sol. water. Decomposed by heat, but may be distilled with steam.—

KA'.—BaA'₂.—CaA'₂.—MgA'₂ & aq. —ZnA'₂ & q.

—CdA'₂ & q. [85°].—CoA'₂ & qq. [90°].—NiA'₂ & q.

—CuA'₂ & q. —AgA'aq. Crystalline pp.

Ethyl ether of the ethyl derivative

CH₂(SEt).CO₂Et. (188°). S.G. \$\frac{1}{2}\$ 1.0469. Oil.

Converted by Et1 at 120° into crystalline

CH₂(SEt₂I).CÖ₂Et.

Ethyl ether of the isoamyl derivative CH2(SC,H11).CO.Et. (230°). S.G. 4 ·9797. Oil.

Phenyl derivative v. Phenyl-thio-gly-

COLLIC ACID.

Benzyl derivative CH2Ph.S.CH2CO2H. [59°]. Formed from CH2Cl.CO2H and benzyl mercaptan (Gabriel, B. 12, 1641). Flat tables. Yields EtA' (275°-290°) and the amide CH₂Ph.S.CH₂.CONH₂ [97°] crystallising in flat plates.

Phenyl-phenyl derivative C.H.Ph.S.CH. CO.H. [170°]. Crystals.

Amide CH2(SH).CO.NH2. A product of the passage of H2S through an alcoholic solution of chloro-acetamide containing a little NH, (Schulze, Z. 1865, 73). Groups of small prisms.

Ethyl derivative of the amide [44°]. CH₂(SEt).CONH₂. Formed from CH₂(SEt).CO₂Et and ammonia. Thin prisms. Reference.-NITROSO-THIO-GLYCOLLIC ACID.

THIO-GLYCOLLIC ORTHALDEHYDE. Ethyl derivative of the ether CH₂(SEt).CH(OEt)₂. (169°). Formed from CH₂Cl.CH(OEt)₂ and NaSEt (Autenrieth, B. 24, 162). Volatile in steam.

Phenyl derivative of the ethyl ether CH₂(SPh).CH(OEt)₂. (278°). Got in like manner.

THIO-HYDANTOÏC ACID C,H,N,SO, i.e. NH,.CS.NH.CH,.CO,H or NH₂·C(NH).S.CH₂·CO₂H. Formed by heating sodium chloro-acetate in aqueous solution with thio-urea (Maly, A. 189, 880). Crystalline powder, al. sol. water, v. sol. acids and alkalis. Decomposed by heat. Converted into thiohydantoin by warming with Na₂CO₂Aq or HClAo.

THIOHYDANTOÏN C,H,N,SO i.s.

CS NH.CH₂ or C NH.CH₂.

Formation.-1. By heating thic-urea with chloro-acetic acid or chloro-acetamide (Volhard, A. 166, 883; Mulder, B. 8, 1264; Maly, A. 168, 133; B. 10, 1853; Claesson, B. 10, 1852).—2. By heating thioglycollic acid (2 mols.) with cyanamide (1 mol.) in aqueous solution (Andreasch, B. 13, 1421; M. 1, 442).

Preparation.—Thio-ures (50 g.) dissolved in water (500 c.c.) is mixed with chloro-acetic acid (62 g.) in water (50 c.c.), heated to 90°, cooled, and mixed with the calculated quantity of NaOH

(Andreasch, M. 8, 414).

Properties.—Long needles (from hot water), insol, alcohol and ether. Decomposes about 200°. Since it cannot be desulphurised by HgO, Liebermann (A. 207, 132) suggests that it does

not contain the group CS.

Reactions.—1. Chlorine passed into its cooled solution in HClAq forms thio-oxy-hydantoin (Kramps, B. 13, 788).—2. Br forms di-bromothio-hydantoin.—3. KClO₈ and HCl yield C₂H₃KN₂SO₃, crystallising in monoclinic tables, S. 1.7 at 22°; 23.3 at 100°, converted by nitrous acid into SO₂H.CH₂.CO₂H, and by baryta-water into sulpho-acetic acid and urea (Andreasch, B. 13, 1423; M. 4, 131).—4. Baryta-water forms thioglycollic acid and dicyandiamide. — 5. Boiling HClAq forms thiocarbimido-acetic acid.

Salts.—Ag,O,H,N,SO.—B',H,SO, Plates.
—B'HNO,.—B'HCl.—B'H,C,O, aq. Prisms.—
B',H,PtCl,.—B'C,H,N,O,. Minute yellow needles.
References.—DI-BROMO- and NITROSO-THIO-

HYDANTOÏN.

DI-THIO-HYDROQUINONE $C_0H_4(SH)_2$. [98°]. Formed by reducing benzene p-disulphonic chloride with tin and HClAq (Körner a. Monselise, G. 6, 142). Formed also from (C_eH₄(NH₂))₂S₂ by diazotisation, treatment with potassium xanthate, and saponification of the product with alcoholic potash (Leuckart, J. pr. [2] 41, 205). Six-sided plates, slowly oxidised by air.

THIO-ISATYDE v. ISATYDE.

THIO-LACTIC ACID CH_s.CH(SH).CO₂H.

a-Sulphydro-propionic acid.

Formation.—1. From a-chloro-propionic scid and KHS (Schacht, A. 129, 1; Lovén, J. pr. [2] 29, 368).—2. By the action of H₂S on silver achloro-propionate or silver pyruvate (Böttinger, A. 188, 320; B. 9, 404, 804, 1061; 11, 1561; 18,

Preparation.—By saturating a solution of pyruvic acid with H2S, and adding conc. HClAq

and zinc (Lovén).

Properties.—Syrup, miscible with water, alcohol, and ether. May be distilled in vacuo. Its odour is unpleasant. FeCl, gives a transient indigo colour, and then oxidises it to sulphidodipropionic acid. CuSO, gives a violet solution; a smaller quantity of CuSO, ppts. the cuprous salt. Cobalt acetate and air gives a brown colour.

Salts.—BaA'₂ (at 180°). Gummy mass.— Hg(S.CHMe.CO₂H)₂.— Hg(S.CHMe.CO₂K)₃ xaq. Hg(S.CHMe.CO₂)₂Ba 2½aq.— AgS.CHMe.CO₂H.

-Bi(S.CHMe.CO₂H)₂. - Pt(S.CHMe.CO₂H)₂. CuS.CHMe.CO₂H. Yellow pp.-PbC₂H₄O₂S. Ethyl other EtA'. Oil. Yields

Cus. CHMe. CO. Et as a yellowish powder.

THIO-DILACTYLIC ACID v. Sulphido-di-

PROPIONIC ACID.

THIO-MALIC ACID C, H, SO,. Formed from bromo-succinic acid and aqueous KSH at 110° (Carius, A. 129, 6). Deliquescent mass, oxidised by dilute HNO, to sulpho-succinic acid.—BaA". Ag₂A". Bulky pp., readily blackening.

THIO-MESITOL v. TRI-METHYL-PHENYL MER-

THIO-DI-METHYL-ANILINE TETRAv. METHYL-DI-AMIDO-DI-PHENYL SULPHIDE.

THIO-METHYL-COUMARILIC ETHER

[91°]. Formed from methyl-coumarilic ether and P2S. (Hantzsch, B. 19, 2400). Yellow needles (from alcohol), v. sol. ether.

THIO-(a)-METHYL-COUMARIN C, H SO i.e. $C_eH_4 < \stackrel{CH:CMe}{O-CS}$. [122°]. Formed by heating (a)-methyl-coumarin with P2S, at 120° (Aldringen, B. 24, 3460). Yellow needles, insol. water, v. sol. alcohol. Converted by alcoholic potash into (a)-methyl-coumarin.

THIO - METHYL - DI - PHENYL - AMINE

METHYL-IMIDO-DI-PHENYL SULPHIDE.

THIO-METHYL-URACIL C₅H₆N₂SO CS<NH.CMe>CH (?). S. .054 at 21°. Formed by the action of alcoholic potash on the needles which separate after some days from a solution of thio-urea (1 mol.) and acetoacetic ether (1 mol.) in alcohol to which a little HCl has been added (Nencki a. Sieber, J. pr. [2] 25,72; List, A. 236, 1; Behrend, B. 19, 219). Plates, v. sl. sol. ether. Decomposed at 280°.

Reactions.—1. The K salt reacts with EtI forming C₃H₃N₂SOEt [145°], and with MeI forming C₄H₅N₂SOMe [220°], which yields C₄H₅AgN₂SO.—2. Chloro-acetic ether forms C₃H₃N₂SO(CH₂.CO₂Et) [143°], which yields C₅H₃N₂SO(CH₂.CO₂H) [204°]. — 2. Br forms methyl-uracil, and finally di-oxy-methyl-uracil. Cl acts in like manner, forming C₅H₈Cl₂N₂O₅.—8. On warming with lead hydrate in strongly alkaline solution it is converted into methyl-uracil. Conc. HClAq at 150°, conc. NH, Aq at 150°, and Ac,O at 185° also form methyl-uracil.

Salts. — $C_5H_4Ag_2N_2SO$. — $C_5H_4CuN_2SO$.

— (C₃H₁N₂SO)₂Hg. — C₃H₃N₂NaSO 2aq. — C₄H₁N₁KSO 3aq. — THIONAMIC ACID. H. Rose (P. 33, 235; 42, 415) found that SO₂ and NH₃ combined in equal volumes, when dry NH, was mixed with excess of dry SO, to form a yellowish white solid, which quickly decomposed in solution, or moist air, giving (NH₄)2SO₄ and ammonium salts of other sulphur oxyacids. Rose concluded that the substance was SO2.NH2 from the fact that it was formed by combining equal volumes of the two gases; if this empirical formula is accepted the compound may be SO, NH, H or SO.OH.NH, When dry SO, was mixed with excess of dry NH, Rose found that 1 vol. SO, combined with 2 vols. NHa; the product—an amorphous, white solid -may be SO.NH. NH, or SO.ONH. NH. M. M. P. M.

THIONAMIDE. By passing NH, into cooled S₂Cl₂ Schiff (A. 102, 111) obtained a white solid which was decomposed by warm water to (NH₄)₂SO₃Aq. The solid contained NH,Cl; Schiff regarded it as a mixture of this salt with SO(NH2)2; he did not attempt to separate the supposed thionamide from NH₄Cl.

M. M. P. M.

(a)-THIO-NAPHTHOIC ACID.

[126°]. Amide C10H7.CS.NH2. Formed from (a)-naphthoic nitrile and alcoholic ammonium sulphide at 80°-35° (Hofmann, B. 1, 40; Bamberger, B. 21, 54). Crystals, v. sol. alcohol. Tetrahydride of the amide

 OH_2 : OH_2 : OH_3 a solution of tetrahydride of (a)-naphthonitrile in alcoholic ammonia saturated with H2S for some days at 30°-40° (Bamberger a. Bordt, B. 22, 629). Dark-yellow oil.

(β)-Thio-naphthoic acid. Amide C₁₀H₂. [149°]. Formed from (\$\text{\delta}\$)-naphthonitrile and ammonium sulphide at 35°-40° (Bamberger a. Bockmann, B. 20, 1115). Needles, v. sol. alcohol. Dissolves in hot water, being slowly converted into the nitrile.

THIO-NAPHTHOL v. NAPHTHYL MERCAPTAN. THIO-DI-NAPHTHYL-AMINE v. IMIDO-DI-

NAPHTHYL SULPHIDE.

THIONATES. Salts of the thionic acids (q. v., p. 698). Four series of thionates are known; a fifth probably exists: M2S2O6 = dithionates; M₂S₄O₆ = trithionates; M₂S₄O₆ = tetrathionates; M₂S₄O₆ = pentathionates; and pro-The thionates bably $M_2S_6O_6 = hexathionates$. are formed by the action of alkalis on the liquid obtained by passing H2S into saturated SO2Aq; also by the oxidation of sulphites, thiosulphates, and mixtures of these, and in other reactions (cf. THIONIC ACIDS, p. 698). For general qualitative reactions of thionates and thiosulphates v.

Debus (C. J. 53, 298).

DITHIONATES, M₂S₂O₆, M²S₂O₆, and M²S₂O₆, (Hyposulphates.) The alkali salts are formed by the interaction of a dilute solution of I in KIAq and dilute NaHSO,Aq (Spring a. Bourgeois, Bl. [2] 46, 151; Sokoloff a. Mattscheffski, B. 14, 2058); also by the reaction of KMnO, Aq on thiosulphates in acetic acid solution, or on sulphites or trithionates (Hönig a. Zatzek, M. 4, 738; Fordos a. Gélis, J. Ph. [8] 36, 113). The Ba salt is formed by passing SO₂ into water holding MnO, in suspension, filtering, and adding BaOAq (for more details v. DITHIONIC ACID, p. 698); most of the other dithionates are obtained by the interaction of BaS2O.Aq and sulphates; several have been prepared by digesting H₂S₂O₆Aq with metallic hydroxides. The dithionates are sol. water; the salts of the alkali and alkaline earth metals are not readily decomposed either in solution or as solids; solutions of most of the other salts are decomposed by heat, giving off SO₂ (v. Klüss, A. 246, 179, 284; Geuther, A. 226, 232).

Dithionates give off SO, when heated to redness, leaving normal sulphates; thus M28,0 = SO₂ + M₂SO₄; this reaction is characteristic; so also is the reaction of H,SO,Aq or HClAq with boiling solutions of dithionates, whereby H₂SO₄Aq is formed and SO₂ is given off, without

separation of S.

Ammonium dithionate (NH₄)₂S₂O₆ Obtained by decomposing the Ba salt in solution by (NH₄)₂SO₄Aq, fiftering and crystallising. S.G. 1.704 (Topsoë, C. C. 4, 76). Klüss (A. 246, 179, 284) gives formula 2Am, S.O. aq, and says the salt crystallises in monoclinic needles. Very sol. water; insol. alcohol. Aqueous solution may be boiled without change. Forms double salts with AmCl (Fock a. Klüss, B. 24, 3017); also with the dithionates of Al, Cd, Co, Cu, Fe, Mn, Ni, and Zn (K., l.c.)

Barium dithionate BaS₂O₆. 2aq. Formed by passing SO₂ into water at c. 0° holding coarselypowdered MnO2 in suspension, till the MnO2 is dissolved $(MnO_2 + SO_2 + Aq = MnSO_4Aq$; and also $MnO_2 + 2SO_2 + Aq = MnS_2O_6Aq$), filtering if mecessary, adding BaOAq to alkaline reaction (MnS₂O₈Aq + BaO₂H₂Aq = BaS₂O₈Aq + MnO₂H₂), filtering from MnO₂H₂, evaporating, and recrystallising from water (cf. Stas., Chem. Proport. 117). White prisms; S. 24.75 at 18°, 90.9 at 100°, 100.6 at $102^{\circ} = b.p.$ of saturated solution; insol. alcohol. The crystals effloresce in air, losing 2H₂O; a hydrate with 4H₂O is obtained by slow evaporation (v. Heeren, P. 7, 55; Gay-Lussac, A. Ch. [2] 10, 312; Baker, C. N. 36, 203; cf. Spring a. Bourgeois, Bl. [2] 46, 151. For crystering and supplies the supplies of the supplies o talline form v. Senarmont, J. 1857. 142; Rammelsberg, P. 58, 295; von Lang, W. A. B. 45 [2] 27). Schiff (A. 105, 239) described double salts with MgS₂O₆ and Na₂S₂O₆ (cf. Kraut, A. 118, 95); Bodländer (Chem. Zeit. 14, 1140) described a double salt with Rb₂S₂O₆; and Klüss (A. 246, 179, 284) double salts with $Ag_2S_2O_4$ and TlaS2O

Many of the other dithionates are prepared

from the Ba salt.

Potassium dithionate K2820st Prepared from a solution of the Ba salt by adding an equivalent quantity of K2SO4 in solution, warming, filtering, and evaporating; also by boiling ing, filtering, and separating K₂SO₂Aq with MnO₂ filtering, and separating from K₂SO₄ by fractional crystallisation (K₂SO₄). White, hexagonal crystals; unchanged in air; decrepitates when heated and then gives off SO₂ and leaves K₂SO₄; S.G. 2-277 (Topsoe, Bl. [2] 19, 246). The crystals rotate the plane of polarisation of light; an aqueous solution is optically inactive (Pape, P. 139, 224). S. c. 6 at ordinary temperature, c. 66 at 100°; insol. alcohol (Heeren, P. 7, 55).

Sodium dithionate Na₂S₂O₆. 2aq. Prepared similarly to the K salt. Also formed, according to Bunte (B. 7, 646), by heating Na.Et.S.O. for some time at 100° (Et.S. distils off). Large, lustrous, rhombic prisms; unchanged in air; S.G. 2·175 at 11° (Baker, C. N. 86, 203). S. 47·6 at 16°; 90·9 at 100°; insol. alcohol. Solution is slowly reduced by sodium-amalgam to Na₂SO₂Aq (Spring, B. 7, 1161). Kraut (A. 117,

97) described a hydrate with 6H2O.

Dithionates of the following metals have Dithonates of the following metals have also been described (the figures indicate the references): Al (1) (8); Be (8); Bi (3); Cd (1) (2) (4); Ca (1) (4) (5); Cr (8); Co (1) (2) (8) (4); Cu (8); Fe (1) (8) (4); Pb (1) (4) (5); Li (2) (4); Mg (1) (4) (5); Mn (4); Hg (1) (2) (8); Ni (2) (4) (5); Rb (6); Ag (1) (2) (4) (5); Sr (2) (4); Tl (8); Sn (8); U (8); Zn (1) (2) (4).

(1) Heeren, P. 7, 72, 171; (2) Rammelsberg,

P. 58, 298; (3) Klüss, A. 246, 179, 284; (4) Topsoë, C. C. 4, 76; (5) Baker, C. N. 36, 203; (6) Piccard, J. pr. 86, 456.

TRITHIONATES M'2S,O6 and M'1S,O6. The alkali salts are formed by digesting conc. MHSO₃Aq with S at 50° - 60° (6MHSO₃Aq + 2S = 2M₂S₃O₃Aq + M₂S₃O₃Aq + 3H₂O)(Pelouze, A.Ch. [3] 79, 85); also by heating dilute MHSO₂Aq, or keeping the solution for a long time in a closed tube (Saint-Pierre, C. R. 62, 632); by passing SO_2 into $M_2S_2O_3Aq$ $(2M_2S_2O_3Aq + 3SO_2)$ = 2M₂S₃O₄Aq + S); by passing SO₂ into M₂SAq (Chancel a. Diacon, J. pr. 90, 35; Rathke, J. pr. 95, 11); by the interaction of I and a mixture of M₂SO₃ and M₂SO₃ in solution $(M_2SO_3Aq + M_2S_2O_3Aq + I_2 = M_2S_3O_6Aq + 2MIAq)$ (Spring, B. 7, 1157); by the interaction of boiling water on AgKS₂O₃ or HgNa₂(S₂O₃), whereby K₂S₃O₆Aq or Na₅S₂O₆Aq is formed, along with Ag₂S or HgS (Spring, *l.c.*); by reacting on M₂SO₄Aq with S₂Cl₂ (Spring, *B.* 6, 1108); by decomposing M₂S₂O₂Aq by a non-oxidising acid, e.g. by H₂SO₄ (Spring a. Levy, Bull. de l'Acad. roy. de Belge, 42, 108); by heating $(NH_4)_2SO_4$ with P_2S_3 , Spring (B. 7, 1158) obtained a little $(NH_4)_2S_3O_8$, along with much $(NH_4)_2S_2O_8$, some NH, polysulphides, and P compounds; K2S2O8 is also obtained from Wackenroder's solution.

The trithionates are soluble in water; they very readily decompose; heated with water they give off SO₂, deposit S, and form sulphates in solution. Addition of excess of CuSO₄Aq to solution of a trithionate causes ppn. of CuS and

evolution of SO₂ (Spring, B. 6, 1108).
SELENOTRITHIONATES M₂SeS₂O₆ are described as dithio-triselenates under Selenates (p. 434). Potassium trithionate K2S8O8. A conc. solu-

tion of K2S2Os is formed by dissolving the salt in 8 pts. water and 1 pt. alcohol till the alcohol separates from the aqueous solution; SO_2 is passed into this solution at $25^{\circ}-80^{\circ}$, solid $K_2S_2O_3$ being added from time to time, till the liquid is yellow and smells of SO2; the crystals that separate are dissolved in water at 60°-70°, the solution is filtered from S, mixed with eight times its volume of 84 p.c. alcohol, warmed gently, and allowed to crystallise (Plessy, A. Ch. [3] 20, 162; for other methods of preparation v. Langlois, A. Ch. [2] 79, 77; Rathke, J. 1864. 164; Chancel a. Diacon, C. R. 56, 710). White rhombic needles (Baker, C. N. 36, 203). sol. water, insol. alcohol. Aqueous solution gives K_2SO_4Aq , SO_3 , and S when heated. Debus (C.J.58, 818) found that $K_2S_2O_4Aq$ slowly changes at the ordinary temperature, giving K2SO4Aq, K₂S₄O₂Aq, and K₂S₃O₃Aq without separation of S. According to C. s. D. (l.c.), K₂S₄O₃Aq interacts with K₂S₂Aq to give only K₂S₂O₃Aq; with H₂S the products are K₂SO₄Aq, K₂S₂O₃Aq, and S (Debus, C. J. 58, 329). Sodium-amalgam reacts readily, producing KNaS₂O₃ and KNaSO₃ (Spring, R. 7.1151) B.7,1161).

The other trithionates that have been isolated are those of Ba (Kessler, P. 74, 253); Pb (K., l.c.; Chancel a. Diacon, C. R. 56, 710); Na (K., l.c.; Rathke, J. pr. 95, 18; Villiers, C. R. 106, 1856); and Zn (Fordos a. Gélis, A. Ch. [8] 22, 66)

TETRATHIONATES MI S,O, and MIS,O, The alkali salts are formed by triturating M.S.O. with I, adding a little water and pouring into

alcohol (von Klobukoff, B. 18, 1869; Eliasberg, B. 19, 322; Fock s. Klüss, B. 28, 2429); the Pb salt is formed by the prolonged interaction of I and PbS₂O₃, also by the interaction of H₂SO₄Aq and a mixture of PbS2O2 with PbO2 (Chancel a. Diacon, J. pr. 90, 55); tetrathionates are also formed, according to Spring a. Levy (Bull. de l'Acad. roy. de Belge, 42, 108), by the interaction of thiosulphates and ferrous or cuprous salts, KMnO,Aq, hypochlorites, or KClO, dissolved in H,SO,Aq. K,S,O, is obtained by neutralising Wackenroder's solution by KC,H,O, (v. Debus, C. J. 53, 278; cf. Curtius a. Henkel, J. pr. [2] 37, 137). The tetrathionates are e. sol. water, but insol. alcohol; aqueous solutions are gene-

rally decomposed on evaporation. Potassium tetrathionate K₂S₄O₄. Formed by adding I, little by little, to conc. K₂S₄O₄Aq till a permanent reddish-brown colour is produced, dissolving the crystals that separate in warm water, filtering from S, adding alcohol till the pp. that forms is re-dissolved, and allowing to crystallise (Kessler, P. 74, 253). Spontaneous decomposition of conc. K₂S₅O₆Aq gives K₂S₄O₆ and S (Debus, C. J. 53, 311). Perfectly dry K₂S₄O₆ can be kept unchanged in a dry atmosphere; but in ordinary air, after a time, SO2 is formed; an aqueous solution of K2S4O6 slowly changes to K₂S₂O₆Aq, K₂S₂O₆Aq, K₂SO₄Aq, and SO₂ (D., *l.c.*). BrAq reacts to form H₂SO₄Aq, S, and KBr; but if the BrAq is added very gradually the S that is formed combines with residual $K_2S_4O_6$ to form $K_2S_5O_6$; similarly, if a little K_2SO_4 is added to K₂S₄O₆Aq, and H₂S is then passed in, K2SO4 and S are formed, and the S combines with unchanged K₂S₄O₆ to form K₂S₅O₆ (D., l.c. p. 815). Sodium-amalgam forms KNaS₂O₈Aq (Spring, B. 7, 1161).

The other tetrathionates that have been isolated are those of Ba (Lewes, C. J. 39, 69; Fordos a. Gélis, A. Ch. [3] 22, 66; Curtius a. Henkel, J. pr. [2] 37, 137); Cd (Kessler, P. 74, 253); Cu (Chancel a. Diacon, C. R. 56, 510); Fe (F. a. G.); Pb (K.; C. a. D.); Na (K.; von Klobukoff, B. 18, 1869); Sr (K.); and Zn (F.

PENTATHIONATES M¹₂S₄O₆ and M¹¹S₄O₆. Wackenroder, in 1845 (A. 60, 189), asserted that the liquid formed by passing H₂S into SO₂Aq contained pentathionic acid H₂S₅O₆; other chemists confirmed his results, and prepared a few salts of the acid (v. Lenoir, A. 62, 253; Fordos a. Gélis, A. Ch. [3] 22, 66; Kessler, P. 74, 257; Ludwig, Ar. Ph. [2] 85, 9; Chancel a. Diacon, C. R. 56, 710). At a later time the existence of pentathionates was denied, chiefly by Spring (B. 12, 2254; 13, 924; 15, 2618), but upheld by Kessler (B. 13, 424; v. also Takatmatsu a. Smith, B. 13, 1976). In 1881 Lewes (C. J. 39, 68) seemed to have isolated BaS₆O₆ and K2S5O6, but Spring failed to confirm his results. Shaw, however (C. J. 43, 357), repeated Lewes's experiments, and obtained K.S.O. In 1888 Debus (C. J. 53, 278) made an elaborate investigation of Wackenroder's solution, and prepared K₂S₃O₆ therefrom. Curtius a. Henkel (J. pr. [2] 37, 187) assert that Wackenroder's solution contains acid tetrathionates, in which the ratio of S to Ba is greater than 4S:Ba, and that these have been mistaken for pentathionates. But Debus's experiments seem to have settled the dispute in favour of the existence of pentathionates. Ammoniacal AgNO Aq gives very distinct brown colouration, followed by a black pp. with pentathionates; this reaction distinguishes them from salts of other S oxyacids (v. Debus,

l.c., p. 297). Potassium Potassium pentathionate 2K₂S₂O₄. Saq. Debus (C. J. 53, 294) prepared this salt from Wackenroder's solution (v. p. 698) by adding 16.66 g. KC₂H₃O₂ (previously fused) to 48 c.c. of the solution of S.G. 1.348; the KC, H,O, was dissolved in as little water as possible acidulated with a few drops of acetic acid; the solution was placed in a large flat dish in a current of air until the liquid had evaporated; the crystalline residue (weighing 26 g.) was repeatedly pressed between filter-paper, and then dissolved in 50 c.c. water + 1 c.c. H_2SO_4 , at 40° ; the minute quantity of S that separated (c. 005 g.) was filtered off, and the filtrate was allowed to crystallise in a flat dish. The crystals that separated (18.75 g.) were a mixture of K₂S₄O₆ and K₂S₅O₆; they were dried by placing them on blotting paper, and the crystals of pentathionate were picked out (5.75 g. were obtained), crystallised twice from water acidulated with a little H₂SO₄Aq, and dried over H₂SO₄. The crystals of pentathionate form four-sided rhombic plates, or six-sided star-like plates; those of tetrathionate form six-sided prisms, with pyramids only on one end, and with the side on which they rest much developed. Lewes (C. J. 39, 75) described three salts: $K_2S_3O_6$, $K_2S_3O_6$. H_2O_7 , and $K_2S_3O_6$. $2H_2O_7$; Debus's salt was $2K_2S_3O_6$. $3H_2O_7$. According to Fock a. Klüss (B. 23, 2428), this is the only salt that exists (cf. D., l.c., p. 297). The salt can also be obtained by passing H₂S into K₂S O₆Aq acidulated with H₂SO₄Aq (D., l.c., p.

Crystals of 2K2S3O3.8H2O begin to decompose, when kept in a stoppered bottle, after a month or so, giving eventually a yellow, pulpy mass of K2S4O6, S, and water; the decomposition is caused by water in the interstices of the crystals. By powdering finely, washing with dilute alcohol, and placing over H₂SO₄, Debus (l.c. p. 295) kept the salt unchanged for two or three years. 2K₂S₅O₆.3H₂O dissolves in c. 2 three years. 2K₂S₅O₆.3H₂O dissolves in c. 2 parts water, forming a clear, neutral solution, which soon decomposes with separation of S and formation of K₂S₂O₂Aq (D., *l.c.*, p. 811); addition of a drop of dilute KOHAq to the solution causes separation of S (p. 291); H₂S ppts. much S, and K2S,Oa and K2S2Os are formed in the solution (p. 828); SO₂ produces K₂S₂O₂Aq and K.S.O.Aq (p. 882); the salt is insoluble in alcohol.

Barium pentathionate BaS, O, . 3aq; obtained by Lewes (C. J. 89, 68) by partially neutralising Wackenroder's solution by BaOAq, crystallising over H₂SO₄, pouring off from BaS₄O₆, and again crystallising.

Copper pentathionate CuS, O, 4aq; obtained by Debus (l.c., p. 800) in blue, prismatic crystals, by adding Cu(C₂H₂O₂)₂Aq to Wackenroder's solution and allowing to evaporate. Debus probably obtained sinc pentathionate, but not in a pure state (l.c., p. 299).
HEXATHIONATES. By adding KC_H_O_Ag

to the mother-liquor from K₂S₄O₆ and K₂S (v. supra), evaporating in a current of air, dissolving in water with a little H2SO, Aq, allowing to evaporate in portions, and collecting the crystals separately, Debus (C. J. 53,301) obtained a salt in which the ratio of S to O was almost 68:60, as required by potassium hexathionate.

M. M. P. M.

THIONESSAL v. Tetra-phenyl-thiophene. Thionic acids. The acids $H_2S_xO_6$, where x=2, 3, 4, 5, and probably 6, are generally called thionic acids, and are distinguished as di-, tri-, &c., thionic acid. The acids are known only in aqueous solutions; most of these solutions readily oxidise to H₂SO₄Aq. Dithionic acid is produced by the action of certain oxidisers on H₂SO₅Aq; dithionates are formed by oxidising sulphites and thiosulphates; trithionates are formed when mixtures of sulphites and thiosulphates are oxidised, also when SO2Aq interacts with alkali thiosulphates; tetrathionates result by the action of certain oxidisers on thiosulphates. These processes may be represented diagrammatically as follows (the actual reactions are much more complex):

(1) 2H₂H₂O₃Aq + O = H₂S₂O₄Aq + H₂O; (2) 2M₂SO₃Aq + O = M₂S₂O₄Aq + M₂O; M₂S₂O₄Aq + 3O = M₂S₂O₄Aq; (3) M₂SO₂Aq + M₂S₂O₃Aq + O = M₂S₃O₄Aq + M₂O; $2M_2S_2O_3Aq + 3SO_2Aq = 2M_2S_3O_6Aq + S;$

(4) $2M_2S_2O_4Aq + O = M_2S_4O_6Aq + M_2O$. After passing H_2S for a long time into nearly saturated H_2SO_4Aq , the solution contains H₂S₂O₆, much H₂S₂O₆, H₂S₂O₆, and probably H₂S₂O₆, along with H₂SO₄, dissolved colloidal S, and a little S in suspension. This solution is known as Wackenroder's solution (Wackenroder, A. 60, 189 [1845]). It has been examined by many chemists; a very thorough investigation was made by Debus in 1888 (C. J. 53, 278; where references will be found to other memoirs). The solution examined by Debus was prepared by passing a slow current of H2S into 480 c.c. nearly saturated SO2Aq at a little above 0° for two hours, keeping in a stoppered bottle for forty-eight hours at the ordinary temperature, passing in H2S for two hours, after forty-eight hours again passing in H2S, and repeating these treatments until the liquid no longer smelled of SO, after standing for a couple of hours at the ordinary temperature (about two weeks were required before the action was completed). The liquid thus obtained contains a large quantity of S, partly as a hard brittle solid, partly as a soft and gummy mass, and partly in suspension in minute particles which pass through filter paper; the liquid remains milky after filtration, and does not become clear on standing for two or three weeks (regarding the variety of S in solution, v. Sulphur, Allotropic forms of, p. 608). Debus (l.c., p. 348) thinks that the product of the direct interaction of SO₂Aq and H₂SA₂ is H₂S₂O₆ (8SO₂Aq + H₂SA₄ = H₂S₄O₆Aq); that part of this H₂S₄O₆ reacts with excess of SO₂Aq to form H₂S₄O₆Aq and H₂S₄O₅Aq), and that part of the H₂S₄O₆Aq + H₂S₄O₅Aq), and that part of the H₂S₄O₆Aq reacting with H₂S₄O₅Aq forms H₂S₄O₆Aq or three weeks (regarding the variety of S in (? H₂S₁O₂Aq + H₂S₂O₃Aq = H₂S₂O₄Aq + H₂SO₄Aq) (D., l.c., p. 333); that the H₂S reacts with

H₂S₄O₂Aq to form H₂O and S, part of the S combining, as it is formed, with H₂S₄O₂Aq to form

H.S.O.Aq, H.S.O.Aq and H.S.O.Aq, and part

remaining in solution as colloidal S. If H.S is passed into Wackenroder's solution, more $H_2S_4O_6$ is formed; and if the passage of $H_2S_4O_6$ is continued until all reaction ceases the thionic acids are decomposed, and the final products are S and H_2O , so that the equation $SO_2Aq + 2H_2SAq = 3S + 2H_2OAq$ correctly represents the interaction of SO_2Aq with H_2SAq so far as the final products are concerned (v. D., l.c., p. 351).

Solutions of the thionic acids more or less rapidly decompose, the final products being H₂SO₄Aq, SO₂Aq, and S; oxidisers such as Cl or HNO₂Aq—in some cases air is sufficient convert them into H2SO.Aq; reducers, such as Na amalgam, produce H₂S₂O₂Aq, sometimes also H₂SO₃Aq, and in some cases H₂SAq. None of the acids has been isolated apart from water; salts of all are known (v. Thionates,

p. 695). The acids are all dibasic.

Regarding the constitution of the thionic acids v. Mendeléeff, B. 3, 870; Michaelis, A. 170,

31; Spring, B. 6, 1108; Debus, C. J. 53, 351.

DITHONIC ACID H₂S₂O₆Aq. (Hyposulphuric acid.) A solution of this acid is formed by suspending coarsely-powdered MnO₂ in water, and passing in SO₂ while the liquid is kept at c. 0°; filtering, adding BaOAq to alkaline reaction, and filtering from BaSO, and MnO₂H₂; evaporating, crystallising out BaS₂O₆, 2aq, and decomposing this salt in solution by the equivalent quantity of H2SO, Aq; filtering from BaSO, and evaporating in vacuo till a liquid of S.G. 1.347 is obtained $(MnO_2 + 2SO_2Aq = MnS_2O_6Aq)$; $MnO_2 + SO_2Aq = MnSO_4Aq$) (Spring a. Bourgeois, Bl. [2] 46, 151). If evaporation is continued beyond this concentration, H_2SO_4Aq and SO_2 are formed; the same decomposition occurs when H₂S₂O₆Aq H₂S₂O₆Aq is evaporated by heat. is reduced by Zn and HClAq to H₂SO₂Aq at 0° (Otto, A. 147, 187); it is oxidised to H₂SO₄Aq by Cl, HNO₈, KClO₃ and HCl &c.; it slowly oxidises in air (cf. Dithionates, p. 695). Thomsen (Th. 2, 259) gives the following heats of formation of $H_2S_2O_6Aq := [S^2,O^3,Aq] = 211,080$; [2SO²,O,Aq] = 68,920; [2SO²Aq,O] = 53,520; for the heat of oxidation of $H_2S_2O_6Aq$ to H_2SO_4Aq $(S_2O_3Aq + O = 2SO_3Aq)$ [S'O'Aq,O] he gives =73,740.

Terthionic acid $H_2S_sO_sAq$. A dilute solution of this acid is formed by decomposing K₂S₂O₆Aq by the equivalent quantity of HClO₄Aq or H₂SiF₆Aq, and filtering from KClO₄ or K₂SiF₆ (Langlois, C. R. 10, 461; 62, 842; Kessler, P. 74, 250). The dilute solution cannot be evaporated, even in vacuo, without partial decomposition, with formation of SO₂ and S; when heated it rapidly changes to H₂SO₄Aq, SO₂ and S. The dilute solution is colourless and odourless, and has a sour and bitter taste (cf. TRITHIONATES, p. 696). The K salt, from which the acid solution is obtained, may be prepared by dissolving K2S2O3 in water containing one-eighth part of alcohol, adding the salt till the alcohol separates from the aqueous solution, passing in SO₂, with addition from time to time of a little K₂S₂O₃, temperature being kept at 25°-80°, till the liquid is yellow and smells of SO₂, setting aside till crystals separate, and re-crystallising from warm water (filtering from separated S) (Plessy, A. Ch. [3] 11, 182).

TETRATHIONIC ACID H₂S₄O₆Aq. PbS₄O₆Aq is decomposed by the equivalent quantity of H₂SO₄Aq (not by H₂S, which reacts with H₂S₄O₄), and the filtrate from PbSO₄ is concentrated on the water-bath, and then in vacuo over H₂SO₄ (Fordos a. Gélis, C. R. 115, 920). PbS_4O_6 may be prepared by pouring a solution of 2 pts. $Na_2S_2O_3.5H_2O$ in much warm water into a dilute warm solution of 8 pts. lead acetate, and thoroughly washing the ppd. PbS₂O₃ with warm water, adding water enough to partially dissolve the PbS₂O₃, then adding 1 pt. I, and letting stand for some days, with frequent shaking, filtering from PbI₂, evaporating and crystallising (2PbS₂O₃ + 2I + Aq = PbS₄O₅Aq + PbI₂).

H₂S₄O₆Aq is colourless and odourless; it has a sour taste. Dilute H₂S₄O₆Aq may be boiled without change, but the conc. solution decomposes to H₂SO₄Aq, SO₂, and S. Dilute HClAq or H₂SO₄Aq does not interact with cold H₂S₄O₆Aq; ngso, and does not interact with cold H_2 SO, and in on warming H_2 S is given off; HNO_2Aq or Cl rapidly oxidises H_2 SO, and to H_2 SO, and (cf. Tetrathionates, p. 696). Thomsen (Th. 2, 262) gives the following data for H.F.: [S', 0', Aq] = 192, 430; $[2SO^2, S^2, O, Aq] = 50, 270$; $[2SO^2Aq, S^2, O] = 34, 870$; $[2SO^2Aq, O] = 53, 490$.

PENTATHIONIC ACID H2S,O,Aq, and HEXA-THIONIC ACID $H_2S_6O_6Aq$. These acids very probably exist in Wackenroder's solution (v. supra, p. 698), as salts of H₂S₅O₆, and probably also of H₂S₆O₆, have been obtained from that solution; solutions of the acids have not been isolated (v. Pentathionates, p. 697, and Hexathionates, p. 697). M. M. P. M.

THIONINE. The group $\underset{C_e}{\text{S}} \xrightarrow{C_e} \underset{H_s(\text{NH}_2)}{\overset{C_e}{\text{H}_s}} > \text{NH}$

 $C_{s}H_{s}(NH_{2}) < \sum_{N}^{S} > C_{s}H_{s}:NH$ (Bernthsen a. Goske, B. 20, 931).

THIONOL $C_{12}H_7NSO_2$ i.e. $S \longrightarrow C_6H_4(OH).N-$ Formed by heating imido-di-phenyl sulphide with dilute H₂SO₄ at 150°-160° (Bernthsen, A. 230, 188). Brownish-red powder, which acquires green lustre on rubbing. Insol. cold water, sol. acids and alkalis, forming a violet solution. Yields di-oxy-imido-diphenyl sulphide (q.v.) on reduction. $-B'_2H_2SO_4$. Green needles. -B'BaO (dried at 100°). Plates with green lustre.—B'Ag $_2$ O (dried at 100°). Brown amorphous pp.

THIONOLINE v. OXY-AMIDO-IMIDO-DIPHENYL

THIONURIC ACID CAH, N. SO, i.e.

CO NH.CO CH.NH.SO.H. Formed by the simultaneous action of NH, and SO2 on alloxan in aqueous solution (Liebig a. Wöhler, A. 26, 268, \$14,881). Formed also by warming nitroso-barbituric acid with ammonium sulphite (Baeyer, A. 127, 210). Crystalline mass of needles, v. sol. water. Decomposed by boiling water into uranil mirror.—(NH₄)₂A"aq. Four-sided tables, sl. sol. cold water. At 200° it yields (NH₄)₂SO₄ and xanthinine (Finck, A. 132, 298).—(NH₄)HA". White needles.—CaA". Prisms.—PbA" aq.

THIONYL-m-AMIDO-BENZOIC ACID

80:N.C,H,.CO,H.

Methyl ether MeA'. [57°]. (212° at 100 mm.). Formed from methyl m-amido-benzoate and SOCl, (Michaelis, A. 274, 250). crystalline mass, v. sol. ether and benzene. Ethyl ether EtA'. Oil.

THIONYL-DIAMIDO-DI-PHENYL-ETHYL-ENE C₂H₄(C₆H₄.N:SO)₂. [202°]. Formed from di-amido-stilbene and SOCl, (Michaelis, A. 274, 265). Red needles, m. sol. benzene and CHCl_s.
THIONYL-AMIDO-PHENOL. Ethyl de-

rivative C.H. (OEt).N:SO. [32°]. (220° at 200 mm.). Formed from p-amido-phenetol and SOCl₂ (Michaelis, A. 274, 246). Light-yellow needles, v. sol. ether. Slowly decomposed by

THIONYL-AMYLAMINE SO:NC.H.,. (87° at 60 mm.). Formed from amylamine (103°) in ether and SOCl, (Michaelis, A. 274, 191). Oil, decomposed by water.

THIONYL-ANILINE C₆H₅N:SO. (200°). S.G. 15 1.236. Formed from aniline in benzene solution and $SOCl_2$ (Michaelis, B.23,8480;24,745). Liquid with peculiar odour, sol. alcohol. Converted by alkalis into aniline and sulphite. Chlorine forms tri-chloro-aniline [78°]. Conc. HClAq also forms aniline and SO₂. The compound C₆H₂Br₂N:SO [75°] is formed from tri-bromo-aniline in benzene and SOCl₂. The compounds [1:2]C₈H₄Cl.N:SO [-8°], [1:3] C₈H₄Cl.N:SO (233°) and [1:4] C₈H₄Cl.N:SO [36°] (287°) are formed in like manner. o-, m-, and p- Thionyl-bromo-anilines melt at 4°, 32°, and 61° respectively (Michaelis, A. 274, 221). Thionyl-tetra-bromo-aniline melts at 78°. Thionyl-p-iodo-aniline [54°] and [4:2:1] $C_8H_2I_2N$:SO [74°] are also crystalline.

Reference.—NITRO-THIONYL-ANILINE. THIONYL BROMIDE SOBr, v. SULPHUR OXY-

BROMIDES, p. 617.
THIONYL-ISOBUTYLAMINE C.H. NSO i.e. CHMe₂.CH₂.N:SO. (116°). Formed from SOCl₂ and isobutylamine in ether (Michaelis, A. 274, 191). Liquid, with pleasant smell.
THIONYL CHLORIDE SOCI, v. SULPHUB

oxychlorides, p. 617. THIONYL-CUMIDINE C.H.Pr.N:SO. (158°). Got from SOCl₂ and the cumidine obtained from isopropyl-benzene by nitration and reduction (Michaelis, A. 274, 239). Oil, not decomposed by boiling water.

Thionyl- ψ -cumidine $C_6H_2Me_3.N:SO.$ [-10°].

(246°). Oil, v. sol. alcohol and ether.

THIONYL-ETHYLAMINE EtN:SO. Formed by adding SOCl2 to an ethereal solution of ethylamine cooled below 0° (Michaelis, B. 24, 756).

THIONYL-MESIDINE C.H.Me.N:SO. [-11°]. (241°). S.G. 14 1·121. Volatile with steam (Michaelis, A. 274, 240).

THIONYL-METHYLAMINE MeN:SO. (59°). Liquid (Michaelis, A. 274, 187).

THIONYL-METHYL-ANILINE

SO(C₆H₄.NHMe)₂. [154°]. Formed from SOCl₂ and methyl-aniline in ethereal solution in presence of AlCl. (Michaelis a. Godchaux, B. 23, Colourless needles, v. sol. alcohol. 3019). Yields a nitrosamine SO(C.H.NMe.NO)2. [171°].

THIONYL-(a)-NAPHTHYLAMINE $C_{10}H_TN:SO.$ [33°]. (226° at 100 mm.). Formed from (a)-naphthylamine in benzene and SOCl. (Michaelis, A. 274, 253). Reddish yellow needles. slowly converted by water into naphthylamine sulphite.

Thionyl- (β) -naphthylamine [53°]. Needles. The thionyl derivative of bromo-(\$)-naphthylamine [63°] melts at 118°

THIONYL-o-NITRO-ANILINE C.H. N. SO. i.e. C₆H₄(NO₂).N:SO. [62°], the m-compound melts at 63°, while the p-isomeride melts at 70° (Michaelis, A. 274, 225)

THIONYL-NITRO-MESIDINE CoH10N2SO3 i.e. C₆HMe₈(NO₂).N:SO. [77°]. Got from nitromesidine [74°] (Michaelis, A. 274, 241). Light yellow needles, v. e. sol. benzene.

Thionyl-di-nitro-mesidine. [127°]. Formed

from di-nitro-mesidine [193°]

THIONYL-NITRO-NAPHTHYLAMINE C₁₀H₆(NO₂).N:SO. Nitro-naphthylamines [191°] and [119°] yield thionyl derivatives melting at

69° and 135° respectively (Michaelis, A. 274, 258). THIONYL-NITRO-TOLUIDINE C, H, N, SO, i.e. O.H.Me(NO.).N:SO. The compounds from (2,1,4)-nitro-toluidine [75°] and (3,1,4)-nitro-toluidine [114°] melt at 44° and 89° respectively (Michaelis, A. 274, 232)

DI-THIONYL-PHENYLENE-DIAMINE

 $C_eH_4(N:SO)_2$. The m- and p- compounds melt at 44° and 116° respectively (Michaelis, A.274,261). THIONYL-PHENYLETHYL-AMINE

PhCH2.CH2.N:SO. (172° at 25 mm.). Liquid, decomposed by water into SO2 and base. passed into its ethereal solution forms amorphous PhCH₂.CH₂.NH.SO₂H, whence benzoic aldehyde yields PhCH₂.CH₂.N(SO₂H).CHPh.OH [114°] (Michaelis, B. 26, 2167).

THIONYL-PHENYL-HYDRAZINE v. Action

of SOCl₂ on Phenyl-hydrazine.

THIONYL-PHENYL-PROPYLAMINE

CH₂Ph.CH₂.CH₂.N;SO. Formed from phenyl-propylamine hydrochloride and SOCl₂ (Michaelis a. Jacobi, B. 26, 2161). SO₂ passed into its ethereal solution forms the amorphous compound NH(SO₂H).C₂H₆Ph, which reacts with benzoic aldehyde forming SO2H.N(CHPh.OH).C3H6Ph

THIONYL-PROPYLAMINE Prn:SO. (104°). Smells like bleaching-powder (Michaelis, A. 274,

190).

THIONYL-TOLUIDINE C.H.Me.N.SO. Formed from toluidine and SOCl₂. The o-compound is liquid (184° at 100 mm.). The m-compound is liquid (184° at 100 mm.). The m-compound is liquid (220°). The p- isomeride [7°] (c. 224°) may be crystallised by cooling (Michaelis a. Herz, B. 23, 8482; 24, 753; A. 274, 231). The thionyl derivatives of (3,1,4)- and (5,1,2)-bromo-toluidines melt at 47° and 50° respectively.

THIO-OBCIN C₆H₃Me(SH)₂. [35°]. Formed by reducing C₆H₃Me(SO₂Cl)₂ with tin and HClAq (Gabriel, B. 12, 1640).—PbC,H₆S₂.

THIONYL-XYLIDINE C.H.Me.N:SO. The (1,2,4)-compound [-9°] (181° at 20 mm.), (1,3,4)-compound (288°), and (1,4,2)-isomeride [-8°] (119° in vacuo) are all oily (Michaelis, A. 274, 287). Thionyl-fluoro-xylidina (144° at 45 mm.) is also an oil.

THIO-OXALIC ACID. Ethylether

CeH. SO, i.e. COEt.CO.SEt. (217° cor.). S.G. 2 1.145. Formed by the action of Cl.CO.CO2Et on mercaptan (Morley a. Saint, C. J. 43, 400). Colourless liquid with faint odour like garlic. Decomposed by KOHAq into mercaptan, alcohol, and oxalate. Alcoholic KOH (1 mol.) forms EtSH and CO₂Et.CO₂K. Water slowly decom-poses it, forming oxalic acid, alcohol, and mer-

Dry NH, forms mercaptan and captan. CO₂Et.CONH.

(β)-Thio-oxalic acid NH2.CS.CO2H. The salt C2H2KNSO, is formed by the action of cold conc. KOHAq on the ethers. It crystallises in needles, v. sol. water and decomposed by evaporating. The free acid is unstable.

Amide of the methyl ether

NH. CS.CO. Me. [86°]. Formed by passing dry H.S into Cy.CO. Me (Weddige, J. pr. [2] 7, 79; 9, 133; 10, 193). Light-yellow crystals, sol. water, alcohol, and ether.

Amide of the ethyl ether NH₂CS.CO₂Et. Mol. w. 133. [64°]. Formed in like manner. Lemon-yellow crystals. Re-converted by boiling with alcohol and Pb(OH)2 into cyanoformic ether. Methylamine and ethylamine form the crystalline compounds CO(NHMe).CS.NH2 and CO(NHEt).CS.NH, respectively.

Amide of the isobutyl ether NH₂.CS.CO₂C₄H₉. [58°]. Lemon-yellow crys-

tals, sl. sol. water.

Di-amide CO(NH₂).CS.NH₂. Formed from CO₂Et.CS.NH₂ and alcoholic ammonia (Weddige, J. pr. [2] 9, 187). Lemon-yellow needles (from alcohol), sl. sol. water.

Di-thio-oxalic acid. Amide

CS(NH2). CS(NH2). Hydrogen rubeanide. Mol. w. 120. Formed, together with yellow Cy.CS.NH₂, by combination of cyanogen with H₂S (Wöhler, P. 3, 177; Völckel, A. 38, 315). Formed also by passing cyanogen into an alcoholic solution of NaSH and ppg. by HCl (Wollner, *J. pr.* [2] 29, 129; Ephraim, *B.* 22, 2305). Prepared by passing H,S into an ammoniacal solution of CuSO₄ which has been decolourised by KCy (Formanck, B. 22, 2655). Small yellowish-red crystals, sl. sol. water, sol. alcohol and ether. Dilute KOHAq forms oxalic acid, conc. KOHAq forms KOy and KCyS. Boiling HClAq forms oxalic acid. MeI and MeOH at 150° form SMe₂I. Alcoholic hydroxylamine hydrochloride (2 mols.) forms $C_2(NH_2)_2(NOH)_2$ and H_2S . Benzoic aldehyde, acting on a boiling solution, forms a compound [209°] which is apparently

CHPh < S > C.C < S > CHPh (Ephraim, B. 24,

1027).—PbC₂H₂N₂S₂: orange pp. (Wallach, B. 13, 528). Yields PbS on boiling with water.

[133°]. Anilide CS(NHPh).CS(NHPh). Formed from oxanilide by successive treatment with PCl, and H2S (Wallach, B. 13, 527). Golden plates.

Di-methyl-di-amide CS(NHMe).CS(NHMe). [140°]. Formed from thiocyanic acid, alcohol, and methylamine (Wallach, A. 262, 360). Yellow crystals, sl. sol. alcohol. Converted by NaOEt and propyl chloride into C(NMe)(SC,H₇).C(NMe)(SC,H₇) (170°-175°).

Di-ethyl-di-amide CS(NHEt).CS(NHEt).[58°]. Formed in like manner. Yellowish-red crystals, insol. water. Converted by NaOEt and EtI into C(NEt)(SEt).C(NEt)(SEt) (126°-129°).

Di-amyl-di-amide CS(NHC,H10).CS(NHC,H10). [60°]. Formed from thiocyanic acid, alcohol, and amylamine (Wallach, A. 262, 862). Red prisms.

Di-bensyl-diamide CS(NHCH,Ph).CS(NHCH,Ph). [115°]. Formed from thiocyanic acid, alcohol, and benzylamine. Yellowish-red crystals, sl. sol. warm alcohol.

Hexa-thio-ortho-oxalic acid. Ethyl ether C₂(SEt)₆. Formed from C₂Cl₅ and NaSEt (Claesson, J. pr. [2] 15, 212). Heavy oil. THIO-OXY- v. SULPHYDRO-.

THIO-PARABANIC ACID. Methyl derivative v. METHYL-THIO-PARABANIC ACID. .

Allyl derivative C.H.N.SO. i.e. CS NH.CO CO. Oxalyl-thio-sinamine. [90°]. Formed by warming the dicyanide of

allyl-thio-urea with dilute H2SO, (Maly, Z. 1869, 260). Lemon-yellow needles, m. sol. cold water.

THIOPHENE C,H,S i.e. SCH:CH (84° cor.). V.D. 3·0 (calc. 2·9). S.G. 2·2 1·062; ½ 1·0884; ½ 1·0705 (R. Schiff, B. 18, 1601). S.V. 84·9. R_∞ 41·4 (Nasini a. Scala, G. 17, 70). H.C.p. 670,900 (Berthelot a. Matignon, C. R. 111, 10; Bl. [3] 4, 252). H.F.-14900. Critical temperature 317° (Pawlewski, B. 21, 2141). Critical pressure 47.7 atmospheres. Occurs in coal-tar benzens to the extent of about 5 p.c. (V. Meyer, B. 16, 1471).

Formation.—1. In small quantity by passing acetylene through boiling sulphur (V. Meyer a. Sandmeyer, B. 16, 2176).—2. By heating succinic anhydride with P2S3.—3. By heating sodium succinate with P₂S₃, the yield being 50 p.c. of the theoretical amount (Volhard a. Erdmann, B. 18, 454). It is not formed by heating s-di-bromo-succinic acid with P2S3 or P2S3 (Ossipoff, J. R. 20, 245).—4. By heating erythrite with P₂S₈ (Paal a. Tafel, B. 18, 688).— 5. From its (a)-carboxylic acid, which is obtained by heating mucic acid with BaS (Paal a. Tafel, B. 18, 456).—6. In small quantities by passing Et₂S through a red-hot tube (Meyer, B. 18, 217).-7. In small quantities by passing ethylene, coal gas, or ligroin-vapour over heated iron pyrites.—8. In small quantities by heating P2S, with crotonic acid, with n-butyric acid, with

Isolation .- 20 pts. commercial 'pure' benzene is shaken for four hours with 1 pt. of H₂SO₄. The acid is diluted with two or three times its weight of water and treated at once with steam. Nearly pure thiophene passes over. Yield, about 3½ p.c. by volume of the H₂SO, used (Schulze, B. 18, 497).

paraldehyde, or with ether (Meyer).

Properties .- Liquid with faint odour, not decomposed below 335°. Not solid at -20° . Greatly resembles benzene. Oxidised by HNO, with great violence. On shaking with isatin and H₂SO₄ it yields a blue colouring matter (indophenine). Not affected by boiling with sodium. Benzene that contains thiophene gives a bright-blue colour to H2SO4 containing nitrous acid. Thiophene (10 g.) shaken with alcohol (100 g.), cold saturated HgCl, solution (1,000 g.), and NaOAc (70 g.) dissolved in water (130 g.) forms a white pp. containing C₄H₅(HgCl)S, which crystallises from alcohol in white plates [183°], and C,H₂(HgCl)₂S, a white powder, insol. alcohol (Volhard, A. 267, 172). The compound C4H3(HgCl)S is converted by AcCl into a-thienyl methyl ketone and by I into di-iodo-thiophene. Thiophene combines with CHPh, and with dibromo-di-(\$)-naphthyl oxide (Liebermann, B. 26, 858).

Reactions.—1. Chloral and H_2SO_4 yield $CCl_2.CH(C_4H_2S)_2.$ —2. BzCl and $AlCl_2$ form thienyl phenyl ketone.—3. Phenyl-glyoxylic acid and H₂SO, form C₁₂H₂NSO₂, which dissolves in chloroform with crimson, and in H₄SO₄ with purple colour.—4. Cl and Br yield chloro- and bromo- derivatives.—5. H₂SO, forms a brown solution containing a sulphonic acid, and slowly decomposes the thiophene, forming H2S, SO2 and an amorphous pp.—6. H₂SO₄ and KNO₅ form a compound called 'nitroso-dithiophene'

 $\rm HO.N < \stackrel{O_4H_2S}{C_4H_2S} > O$ (?), ppd. by water as a dark insoluble powder (Liebermann, B. 20, 3231).—7. Passed with POl, through a red-hot tube it forms C₄H₃S.PCl₂ (218°) which is decomposed by water into C₄H₃S.P(OH)₂ [70°] and is converted by Cl into C₄H₃S.PCl₄, which is converted by SO₂ into C,H,S.POCl, (259°) and by water into C,H,S.PO(OH), [159°] (Sachs, B. 25, 1514).—8. Diazobenzene chloride and AlCl, forms (B)phenyl-thiophene [57°] (Möhlau a. Berger, B. 26, 2001).

References .- AMIDO-, BROMO-, DI-BROMO-DI-NITRO-CHLORO-, TRI-CHLORO-NITRO-, IODO-, IODO-NITRO-, and NITRO- THIOPHENE.

THIOPHENE (a)-CARBOXYLIC ACID

 $C_bH_4SO_2$ i.e. $S < \stackrel{CH}{\underbrace{CH}} \stackrel{CH}{\underbrace{CH}} CH$. Thiophene 'B'carboxylic acid. [126.5°]. (260° cor.). H.F. 85,400. H.C.p. 591,900 (Stohmann, J. pr. [2] 43, 12). Formed by adding sodium-amalgam to a mixture of Cl.CO₂Et with (a)-iodothiophene, with di-iodo-thiophene, or with di-bromo-thiophene (Nahnsen, B. 17, 2192; 18, 2304; Bonz, B. 18, 2306). Formed also by heating mucic acid with BaS for 6 hours at 205°, the yield being 12 p.c. of the theoretical (Paul a. Tafel, B. 18, 456). Obtained by oxidation of (a)-ethyl-thiophene (Schleicher, B. 18, 3017), of (α)-propyl-thiophene (Ruffi, B. 20, 1740), of thienyl methyl ketone (Peter, B. 17, 2645; 18, 542), and of thienyl ethyl ketone (Ernst, B. 20, 518) with alkaline KMnO4.

Properties.-Slender white needles (from water) or long flat needles (by sublimation), v. sol. hot water, sl. sol. cold, v. e. sol. alcohol and ether, m. sol. chloroform, sl. sol. ligroin. Gives a blue colour on warming with isatin and H₂SO₄ (indophenine reaction). Yields thiophene on distilling with lime. Yields a di-bromo-derivative [211°].

Salts.—BaA'₂ 2aq. Small crystals. S. 22·19 at 14·5°.—CaA'₂ 3aq. S. 18·5 at 18·5°.—ZnA'₂ (dried). S. 14 at 15°.—PbA'₂ (dried) ·5 at 18·5°.—AgA': plates or needles. S. ·195 at 11°.

Ethyl ether EtA'. (218° cor.). S.G. 22 1.1155. Liquid, smelling like benzoic ether.

(190° Chloride C.H.S.COCL Liquid, smelling like benzoyl chloride.

A mide C.H.S.CONH₂. [174°] (Meyer, A. 236, 210). Prisms, m. sol. ether. Phenyl thiocarbimide forms C4H4S.CO.NH.CO.NHPh. [206°]

Anilide C.H.S.CONHPh. [140°]. Formed by the action of phenyl cyanate on thiophene in presence of AlCl, (Leuchart a. Schmidt, B. 18,

2340). Iridescent plates.
Amidoxim C.H.S.C(NOH).NH. Formed by the action of hydroxylamine on the nitrile (Meyer, A. 236, 213). Long prisms (from

Nitrile (192°) (D.); (200° cor.) (Peter, B. 18, 543). Formed by distilling thienyl-glyoxylic acid with aqueous hydroxylamine (Douglas, B. 25, 1311).

Tetrahydride SCH(CO₂H).CH₂. [51°]. H.F. 114,900. H.C.p. 700,400. Formed by reducing the acid, in alkaline solution, with sodiumamalgam (Ernst, B. 19, 3278; 20, 518). Plates or needles (from ligroin), v. e. sol. water and alcohol, v. sol. ether. Volatile with steam. alcohol, v. sol. ether. Volatile with steam. Much decomposed on distillation. Reduces ammoniacal AgNO₃. Salts.—CaA'₂ 3aq. Crystalline, sol. water. — AgA'. M. sol. water. (c. 206°). Methyl ether MeA'. Ethylether EtA'. Liquid.

SCH:CH (γ)-Thiophenic acid. [136°]. S. 44 at 17°. Formed by the action of alkaline KMnO, on (3)-methyl-thiophene (Muhlert, B. 18, 3003) or (β) -ethyl-thiophene, the yield being about 8 p.c. (Damsky, B. 19, 3284). Needles (from water), sol. ether, votatile with steam. Gives the indophenine reaction.—CaA'2 \frac{1}{2}aq. S. (of CaA'2) 7.92 at 14.5°.—BaA'2 (dried). S. 11.54 at 17°.—AgA'. Needles or plates, v. sl. sol. Aq. Amide C,H,S.CONH₂. [178°]. Converted by phenyl cyanate into C,H,S.CO.NH.CO.NHPh.

Thiophene (β) -carboxylic acid

[2060]

Thiophene 'a'-carboxylic acid. (258° cor.). This is a mixture of the (a)- and (β) - acids. It is got by saponifying the mixed nitrile which is formed by distilling a mixture of potassium thiophene (a) and (β) sulphonates with KCy (V. Meyer, B. 16, 2174; 18, 2315; Bonz, B. 18, 2308). Formed also by oxidising a mixture of (α) -methyl-thiophene (2 pts.) and (β) methyl-thiophene (3 pts.) and by slow evaporation of an aqueous solution of the mixed thiophene carboxylic acids in the cold (Egli, B. 18, 548; V. Meyer, B. 19, 2891; A. 236, 221). Volatile with steam. Can be separated into the (a)and (β) - acids by crystallisation.

Thiophene dicarboxylic acid

S<CH=CH C(CO₂H):C(CO₂H). [270°]. Formed by oxidising the corresponding di-methyl-thiophene with alkaline KMnO₄ (Grünewald, B. 20, 2586). Formed also by oxidising (3)-methyl-thienyl methyl ketone (Gerlach, A. 267, 153). Long needles (from water), v. e. sol. ether. Not volatile with steam. Resorcin at 200° forms a fluorescein which exhibits yellowish-green fluorescence in alkaline solution. - PbA". -- BaA". Crystals, al. sol. cold water.—Ag₂A". White pp. Methyl ether Me₂A". [59.5°]. Plates.

Thiophene dicarboxylic acid S<C(CO,H):CH

Formation.-1. By oxidation by alkaline KMnO, of di-methyl-thiophene from coal tar (Messinger, B. 18, 567), from (α)-ethyl-thienyl methyl ketone, from (a)-ethyl-thiophene (a)-carboxylic acid (Schleicher, B. 18, 3020), from methyl-thienyl methyl ketone (Demuth, B. 18, 3025), and from (a)-methyl-thiophene carboxylic acid (Levi, B. 19, 656).—2. By saponifying its nitrile, which may be got by fusing potassinm thiophene disulphonate with KCy (Jackel, B. 19, 191).—3. By saponifying its ether which is got by the action of sodium-amalgam on a mixture of di-bromo-thiophene and ClCO₂Et (Bonz, B. 18, 2306).

Properties.—Crystalline powder, v. sl. sol. cold water, sol. ether. Not melted at 300°. Does not give the fluorescein reaction.

Salts. — BaA"aq. Crystalline, v. sl. sol. cold water.—CaA" 3aq.—Ag,A": white pp.

Methyl ether Me,A". [145°] (J.); [147°] (S.); [151^c] (D.). White needles from alcohol. Ethyl ether Et₂A". [47^c] (J.); [52^c]
(S.). Long needles (from alcohol). Nitrile C₄H₂SCy₂. [92^c]. Crystals.

Tetrahydride SCH(CO₂H).CH₂CH(CO₂H).CH₂ cor.]. Formed by reducing the acid in alkaline solution by sodium-amalgam (Ernst, B. 19, 3275). Plates, v. sol. water, sol. ether. Reduces hot ammoniacal AgNOs. Hot conc. H₂SO₄ evolves CO (1 mol.).—BaA": scales.—Ag₂A": white powder.—Me₂A". Oil.

Thiophene dicarboxylic acid

SCH C.CO₂H. Formed by oxidation of the corresponding di-methyl-thiophene by alkaline KMnO, (Zelinsky, B. 20, 2022). Needles, sl. sol. cold water. Decomposes and partially sublimes at 280°, v. sol. hot water.—Ag₂A".

Methyl ether Me₂A". [121°]. Plates. Ethyl ether Et₂A". [36°].

Thiophene tri-carboxylic acid C₄SH(CO₂H), Formed by oxidation of di-methyl-thienyl methyl ketone with KMnO. (Messinger, B. 18, 2302).

Methyl ether Me.A". [118°] Plates, v.

sol. alcohol and ether.

References .- Bromo- and Iodo- THIOPHENE CARBOXYLIC ACID and NITRO-THIOPHENIC ACID.

THIOPHENE GREEN C21H22N2S. The leucobase C₄SH₃.CH(OH)(C₆H₄NMe₂)₂ is got by heating thiophenic aldehyde with di-methyl aniline, ZnCl2, and a little alcohol at 100° (Levi, B. 20, 513). It crystallises in needles [93°], and is readily oxidised to thiophene green C₄SH₃.C(OH)(C₆H₄NMe₂)₂, which is an oil and forms the salts (C₂H₂₂N₂S)₃ZnCl₂ 2aq, B'H₂SO₄, B'₂SH₂CO₄ 2aq, and B'2C₆H₄N₃O₇, crystallising in scales with coppery lustre.

Leuco-thiophene green v. Tetra-methyl-Di-AMIDO-DI-PHENYL-THIENYL-METHANE.

THIOPHENE (a)-SULPHINIC ACID C₄H₃S.SO₂H. [67°]. Formed by the action of zinc-dust on an alcoholic solution of thiophene sulphonic chloride (Weitz, B. 17, 800). Needles, v. sol. water, alcohol, and ether. Gives the indophenine reaction.—BaA', 2aq. White plates, v. sol. water.—ZnA', 3aq.—ÂgA'. Crystalline pp. THIOPHENE 'α'-SÜLPHONIC ACID White plates,

C.H.S.SO.H. Formed by shaking thiophene (20 g.) dissolved in ligroin (2,000 c.c.) with H₂SO₄ (800 c.c.) for 2 hours (V. Meyer a. Kreis, B. 16, 2172; Biedermann, B. 19, 1615). Deliquescent mass. Yields thiophene on distillation. It is doubtless a mixture of the (a) and (3) acids.
—NaA'aq.—BaA'₂Baq.—CaA'₂.—PbA'₂aq. Very -NaA'aq.-BaA'23aq.-CaA'2.soluble powder.-AgA' Saq. Plates (Weitz, B. 17, 796).

Ethyl ether EtA'. Oil. Chloride C, H, S.SO, Cl. [28°]. (above 200°). Oil or colourless crystals.

Ami de C,H,S.CONH, [142°] (Langer, B. 17, 1568). Needles (from water).— C.H.S.CO.NHAg. Pearly scales.

Anilide C.H.S.CONHPh. [96°]. Thiophene (B)-sulphonic acid

·CH:ÇH Formed by the action of s<CH:C.SO.H. sodium-amalgam on di-bromo-thiophene (β)sulphonic acid, which is got by sulphonation of di-bromo-thiophene (Langer, B. 17, 1567; 18, Obtained in like manner from iodothiophene sulphonic acid, and from tri-bromo-thiophene sulphonic acid (Rosenberg, B. 18, 1776). Crystalline solid, v. sol. water. Gives a blue colour with isatin and H2SO4.—BaA'2

Chloride C,H,S.SO,Cl. [43°]. Plates, v. sol. ether, insol. ligroïn.

Amide $C_4H_3S.SO_2NH_2$. [148°]. Tables. Thiophene disulphonic acid $C_4H_2S(SO_3H)_2$. Formed by reduction of di-bromo-thiophene 'β' - di - sulphonic acid by sodium - amalgam (Langer, B. 18, 554, 1114; Rosenberg, B. 18, 3030). Crystalline, v. sol. water.—BaA" $2\frac{1}{2}$ aq. 3030). Crystalline, v. sol. water.—BaA

Chloride C,H,S(SO,Cl),. [149]. Scales. Amide C,H,S(SO,NH,),. [c. 280]. Needles. Thiophene disulphonic acid C,H,S(SO,H),. Formed by reducing iodo-thiophene (aB)-disulphonic acid with sodium-amalgam (Langer, $B. \, \bar{1}8, 560).$

A mide $C_1H_2S(SO_2NH_2)_2$. [142°]. Plates.

Gives the indophenine reaction.

Thiophene disulphonic acid C4H2S(SO3H)2 Formed from lead thiophene sulphonate and fuming H₂SO₄ (Jackel, B. 19, 185, 1066). Crystalline, v. sol. water and alcohol.—K₂A"aq: prisms, v. sol. water.—Na,A" 3aq. Needles, v. sol. water.—BaA" 3aq. Flat prisms, sl. sol. water.—CuA"4aq. Blue needles.—Ag₂A".

Chloride C₄H₂S(SO₂Cl)₂. [77.5°]. Needles

(from ether).

Amide C₄H₂S(SO₂NH₂)₂. [211.5]. References.—Di-Bromo-, Iodo- and Nitro-THIOPHENE SULPHONIC ACID.

THIOPHENIC ACID v. THIOPHENE CARB-

OXYLIC ACID.

THIOPHENIC ALDEHYDE C,H,S.CHO. (198° cor.). S.G. 21 1-215. Formed by distilling (a)-thienyl-glyoxylic acid (Biedermann, B. 19, 637, 1853). Formed also by the action of H2S on CHCl $\stackrel{\text{CH}_1\text{CH}_2}{\text{CO}}$ (Hantzsch, B. 22, 2838). Oil, smelling like benzoic aldehyde. Oxidised by air to thiophene (a)-carboxylic acid.

Oxim C,H,S.CH(NOH). Thiophene synal-doxim. [128°]. Needles (from ether). Converted by NaOHAq into thiophene carboxylic nitrile (Hantzsch, B. 24, 47). Yields an acetylderivative [75°-80°]. NaOMe and MeI in MeOH yield C₄SH₃.CH<0 [120°], orystallising in bables: converted by dilute H.SO, into the aldehyde and (B)-methyl-hydroxylamine (Goldschmidt a. Zanoli, B. 25, 2588). Phenyl cyanate in ether forms C4SH2CH:NO.CO.NHPh [70°], crystallising in needles

Oxim C,H,S.CH:NOH. Thiophene anti-aldoxim. Formed by the action of hydroxyl-amine hydrochloride, NaOHAq and Na₂CO, on the aldehyde (Goldschmidt a. Zanoli, B. 25, 2590). Converted by phenyl cyanate into C,SH₂,CH:NO.CO.NHPh [144°], whence N2OH

forms C₄SH₃.CH:N.NHPh [46°]. o-Tolyl cyanate forms C₄H₃S.CH:NO.CO.NH.C₇H, [66°].

C.SH.CH:N.HPh. Phenyl hydrazide

[134.5°]. Needles (from alcohol). THIOPHENIC IMIDO-ETHYL ETHER

C₄H₃S.C(OEt):NH. The salt B'HCl is formed by the action of dry HCl on the nitrile of thiophene carboxylic acid (Douglas, B. 25, 1312). It is crystalline, and is converted by hydroxylamine into C.H.S.C(OEt):NOH [67°].

THIOPHENINE is AMIDO-THIOPHENE.

THIOPHENOL v. PHENYL MERCAPTAN.

THIOPHENURIC ACID C.H.NSO. [172°]. Excreted in the urine of rabbits after injection of sodium thiophene (a)-carboxylic acid (Jaffé a. Levi, B. 21, 3458). Prisms, resembling hippuric acid (from water), v. sl. sol. ether, v. sol. alcohol. Split up by boiling baryta-water into glycocoll and thiophene (a)-carboxylic acid.—BaA', 2aq. Needles, v. sol. water.—CaA', xaq. - AgA'.

THIO-PHENYL-ACETIC ACID. Amide C₈H₈.CH₂.CS.NH₂. [98°]. Formed by heating phenyl-acetamide (3 pts.) with P₂S₅ (2 pts.), and also by allowing a solution of phenyl-acetonitrile in alcoholic ammonium sulphide to stand for some days (Bernthsen, B. 11, 503; A. 184, 292). Trimetric crystals (from alcohol), decomposed on

distillation into H.S and C.H.CH.CN.

Reactions.—1. Boiling NH.Aq forms phenylacetamide. -2. HgO forms phenyl-acetonitrile. 3. Zn and HClAq form phenyl-ethyl-amine (Colombo a. Spica, B. 8, 821). — 4. Bensoic addehyde forms oily C₂₃H₂₂N₂S₂(?)—5. Sodium-amalgam forms Ph.CH₂CH₂NH₂ and a compound C₂₄H₂₇NS₂ [108°].—6. Iodine added to a hot alcoholic solution forms C₁₅H₄NS₂ [42°].— 7. MeI forms (CH,Ph.C(NH).SMe)HI [139°] which gives rise to (C₂H₁₁NS)₂H₂PtCl₂.—8. EtI forms (C₁₀H₁₃NS)HI [116°], which is v. e. sol. water, and yields, on treatment with NaOHAq, the free base CH₂Ph.C(NH).SEt, which is an oil that quickly decomposes into mercaptan and phenyl-acetonitrile. The salts (C₁₀H₁₂NS)HCl [c. 120°], B'₂H₂PtCl₅ [below 130°], and B'HBr are crystalline .- 9. Aniline forms the compound CH2Ph.C(NH2):NPh.

THIO'- PHENYL - ALLOPHANIC ACID v. PHENYL-THIO-ALLOPHANIC ACID.

THIO - DI - PHENYL - AMINE v. IMIDO-DI-

PHENYL SULPHIDE. THIO - PHENYL - CARBAMIC ACID v. PHENYL-THIO-CARBAMIC ACID.

THIO - DI - PHENYL KETONE v. THIO-BENZOPHENONE.

THIO - DI - PHENYL - METHYL - AMINE v.

METHYL-IMIDO-DI-PHENYL SULPHIDE.
DI-THIO-PHTHALIC ACID C_eH₄(CO.SH), The salt K2A" is formed by dissolving KSH (2 mols.) in an alcoholic solution of phenyl phthalate (1 mol.). The free acid splits up at once into H₂S and thiophthalic anhydride (Schreder, 'B. 7, 706).

(a) - Di - thio - isophthalic acid. The K salt, formed by the action of KSH on phenyl

isophthalate, crystallises in yellow needles.

Ethyl ether Et₂A". Formed by adding

water to the hydrochloride of C.H. (C(NH).SEt). (Luckenbach, B. 17, 1435). Small needles.
(β) - Di - thio - isophthalic acid. An Amide

C.H. (CSNH2)2. [200°]. Formed by heating isophthalic nitrile with alcoholic ammonium sulphide (L.). Needles, sol. hot alcohol. Reconverted into the nitrile by lead acetate. Boil-

ing KOHAq forms isophthalic acid.

THIO.PHTHALIC ANHYDRIDE C.H. 802.i.s. C.H. CO.S. [114°]. (284°). Formed by the action of KSH on phthalyl chloride, or on phenyl phthalate (Graebe a. Zschokke, B. 17, 1175). Formed also by the action of As. S. on phthalyl chloride (Rayman, Bl. [2] 47, 898). Plates, v. sol. hot alcohol, ether, and chloroform, al. sol. water. Volatile with steam. Sol. alkalis and converted by long boiling with alkalis into phthalic acid. Ammonia forms phthalimide. Resorcin and H. SO. give a thiofluorescein, the alkaline solution of which is deep red with green fluorescence.

THIO-PHTHALIDE C₆H₄C_{CO}S. [57°]. Formed by heating C₆H₄Cy.CH₂:SCy with conc. HClAq for 5 hours at 180° (Day a. Gabriel, B. 28, 2480). Thin needles (from dilute alcohol). THIO-PHTHALIMIDINE C₆H₄NS i.e.

C_cH₄C(NH) S or C_cH₄Cy.CH₂SH. [62°]. Formed by warming o-cyano-benzyl sulphocyanide with H₂SO₄ at 65° (Day a. Gabriel, B. 23, 2481). Formed also by adding KSHAq to o-cyano-benzyl chloride in alcohol. Needles, which darken in air, v. sol. alcohol and ether. Boiling NaOHAq forms thiophthalide. KSH added to its alcoholic solution forms C_{1c}H_{1c}S₂, crystallising from nitrobenzene in needles with green lustre.—B'HCl. Needles.—B'₂H₂PtCl₂. Orange-yellow prisms.—B'HI. Needles.—Piorate: slender needles.

THIOPHTHENE C_eH₄S₂ i.e. CH—C—CH. (225° cor.). Formed by distilling citric acid (1 pt.) with P₂S₂ (2 pts.), adding NaOHAq, and distilling with steam (Biedermann a. Jacobson, B. 19, 2444). Oil, not solid at -10°. Gives the indophenine reaction. — B'C₂H₂N₂O₃. [138°]. Yellow needles, v. sol. alcohol and benzene. Reference.—Tetha-bromo-thiophthene.

tejerence.—IETRA-BROMO-THIOPHTHENE.
THIOPHYLLINE C,H₂N₁O₂. [264°]. Occurs, in small quantity, in extract of tea (Kossel, B. 21, 2164). Crystals (containing aq), more sol. water and alcohol than theobromine, v. sol. NH₄Aq. Forms salts with acids. On evaporation with chlorine-water it leaves a scartic residue, turned violet by NH₂Aq.—C,H,AgN₄O₂. Amorphous, converted by MeI into caffeine C,H,MeN₁O₂ [229°].

(a) - THIO-PROPIONIC ACID C.H.OS. Methyl ether CH.CH.CO.SMe. (120°). V.D. 51·23. Formed from propionyl chloride and lead mercaptide in the cold (Obermeyer, B. 20, 2922).

(8)-Thio-propionic acid Et.CS.OH. The salt NaA'aq formed by boiling propionitrile with alcoholic ammonium sulphide is crystalline

(Dupré, Bl. [2] 29, 804).

Amide EtC(NH).SH. [42°]. Formed by heating propionamide (5 mols.) with P₂S₄ (1 mol.) in benzene for a short time (Hubacher, A. 259, 229). Yellowish plates, v. e. sol. benzene. THIO-ISOPEOPYL-COUMARIN C₁₂H₁₅SO i.e.

THIO-ISOPROPYL-COUMARIN C₁₂H₁₂SO s.c.

O₂H₁C₁C₁C₂H₇). [81°]. Formed by heating

Isopropyl-coumarin with P₂S₂ at 120° (Aldringen,

B. 24, 8463). Reddish-yellow needles, sol. hot alcohol, sl. sol. ether.

THIO - PRUSSIAMIC ACIDS. By heating ammonium sulphocyanide there is formed, besides thio-urea, dithiodiprussiamic acid $C_cH_1N_oS_a$ a yellowish-white powder, sl. sol. hot water; thiodiprussiamic acid $C_cH_oN_oS_a$ morphous, m. sol. hot water; and ammonium dithiotriprussiamate $C_oH_{13}N_{16}S_a$, which crystallises from water. The first acid gives no colour with FeCl₃, the two other bodies give a red colour. The lead salts of these acids dissolve in hot water.

THIOTETRAPYRIDINE, so called,

C₂₀H₁₈N₄S(?). [155°]. Formed by heating nicotino (1 pt.) with sulphur (6 pts.) at 150° till the mass is chrome-green (Cahours a. Etard, Bl. [2] 34, 452). Yellow prisms (from alcohol).—B'2HCl.—B'HHgCl₂.—B'H.PtCl₃: yellow amorphous pp.

B'HHgCl_s.—B'H.PtCl_s: yellow amorphous pp.
THIO - PYRÔCATECHIN v. OXY-PHENYL

MERCAPTAN.

THIO-PYROMUCIC ACID. Amide C₄H₃O.CS.NH₂. [127°]. Formed by passing H₂S into a solution of furfuronitrile in alcohol-ether (Douglas, B. 25, 1314). Yellowish crystals, sl. sol. cold water, v. sol. alcohol.

THIO-PYRUVIC ACID. The compound (C₃H₄O₂S)C₃H₄O₃ [87°] separates as a powder when H₅S is passed into pyruvic acid (Böttinger, A. 188, 325). It is v.e. sol. water and is converted by boiling HIAq into CH₃.CH(SH).CO₂H. Boiling water forms H₂S and thiopyruvic acid.

DI-THIO-RESORCIN C₆H₄(SH)₂. Mol. w. 142. [27°]. (243°). Formed by reducing the sulphochloride C₆H₄(SO₂Cl)₂ with tin and HClAq (Patschke, J. pr. [2] 2, 418; Körner, G. 6, 140). Crystals, smelling like some geraniums. Pb(OAo)₂ added to its alcoholic solution ppts. orange C₆H₄S₂Pb, whence McI forms C₆H₄(SMe)₂ (278°) (Obermeyer, B. 20, 2927).

THIO-RUFIC ACID C₁₀H₁₄S₃O₄. Formed by the action of CS₂ on a mixture of NaOEt and acetoacetic ether (Norton a. Oppenheim, B. 10, 701). Pearly orange scales, nearly insol. water.—NaA': brick-red needles.—CaA'₂: red needles.

THIO SALICYLIC ACID v. OXY-THIO-BENZOIG ACID.

THIOSINAMINE v. ALLYL-THIO-UREA.

THIO.SUCCINIC ACID. Anhydride C₂H₄CO₂O. 'Sulphosuccinyl.' [31°]. (225°). Formed by adding succinyl chloride dissolved in benzene or CS₂ to an aqueous solution of Na₂S. Got also by distilling succinic acid (100 g.) with P₂S₃ (100 g.) (Auger, A. Ch. [6] 22, 333). White crystals. Its aqueous solution gives off H₂S on heating. Phenylhydrazine forms CO₂H.C₂H₄.C(SH):N.NHPb [120°] and C₂H₄CO₂C(N₂HPh) O [216°], which on distillation changes to succinyl-phenyl-hydrazine [155°]. Phenyl-hydrazine also produces C₂H₄(CO.N₂H₂Ph)₂ [219°] (Zanetti, C. C. 1889, 668). Sodium-amalgam forms C₄H₅NaSO₂, converted by phenyl-hydrazine acetate into CO₂H.C₂H₄.CH:N₂HPh [c. 175°].

Di - thio - succinic acid C₂H₄(CO.SH)₂. The salt K₂A'', got by boiling succinyl-phenol with KSH in alcohol, crystallises in minute needles, v. sol. water, alcohol, and ether (Weselsky, B. 2, 518). Its aqueous solution is decomposed by spontaneous evaporation. Acids-

split it up into H2S and thiosuccinic anhydride

(v. supra).

Ethyl ether C₂H₄(CO.SEt)₂. (271°). Formed from di-phenyl succinate (42 g.) and sodium mercaptide (30 g.) in dry ether in the cold (Seiffert, J. pr. [2] 31, 469). Oil, smelling of garlic.

THIO-SUCCINURIC ACID C, H, N, SO, . [211°]. Formed by heating succinic anhydride with thiourea at 140° (Pike, B. 6, 1105). Yellowish powder, insol. alcohol and ether, sl. sol. water and HOAc. Decomposed by alkalis into thio-urea and

succinic acid.

(a) - DI - THIO - TEREPHTHALIC ACID $C_8H_8S_2O_2$ i.e. $C_8H_4(CO.SH)_2$. Formed by the action of alcoholic KSH on di-phenyl terephthalate (Schreder, B. 7, 708). Amorphous, v. sl. sol. alcohol.

(β) - Di - thio - terephthalic acid. Amide C_cH₄(CS.NH₂)₂. [263°]. Formed by heating terephthalic nitrile with alcoholic ammonium sulphide (Luckenbach, B. 17, 1430). Yellow amorphous powder, insol. alcohol. Boiling with

lead acetate reproduces the nitrile.

THIOSULPHATES. Salts of the hypothetical thiosulphuric acid $H_{1}S_{2}O_{3}$. These salts were called hyposulphites before the isolation of the true hyposulphites $(M_{2}S_{2}O_{4}, p. 592)$. $Na_{2}S_{1}O_{3}$. 5aq is still, however, commonly known as sodium hyposulphite. Thiosulphates are formed by boiling normal alkali sulphites in solution with S; $(M_{2}SO_{3}Aq + S = M_{2}S_{0}Aq)$; caustic alkali solutions and milk of lime also give thiosulphates by boiling with S. Thiosulphates are produced by the action of air, SO_{4} , or $K_{2}Cr_{2}O_{4}Aq$ on the alkali and alkaline earth polysulphides in solution $(M_{2}S_{3}Aq + 3O = M_{2}S_{2}O_{4}Aq + 3S$; $2M_{2}S_{3}Aq + 3SO_{2}Aq = 2M_{2}S_{2}O_{4}Aq + 9S$; $2M_{2}S_{3}Aq + 4M_{2}Cr_{2}O_{4}Aq + H_{2}O$

 $2m_2S_3Aq + 4m_2O_2O_3Aq + H_2O$ = $5M_2S_2O_2Aq + 4Cr_2O_3 + 2MOHAq$ (Döpping, A. 46, 172). Spring (B. 7, 1157) by heating (NH₄)₂S₀O₄, with P₂S₅ got a large quantity of (NH₄)₂S₂O₃, along with (NH₄)₂S₂, and also some (NH₄)₂S₂O₄, and P compounds; and by the interaction of H₂S and SO₂NH₂.ONH₄ he obtained (NH₄)₂S₂O₄. A little H₂S₂O₄Aq is said to be formed by the interaction of steam and molten

S (Myers, A. 129, 351).

The alkali salts, also the Ca and Sr salts, are readily soluble in water; many of the others are insoluble or only slightly soluble; the insoluble salts are often obtained from solutions of the alkali salts by double decomposition. Thiosulphates are decomposed by heating strongly; alkali salts leave a mixture of sulphate and sulphide; other salts also give off H₂S and S. Aqueous solutions of thiosulphates evolve H₂S when boiled, forming sulphates, or sulphides if the metal of the thiosulphate is ppd. by H₂S.

Aqueous solutions of thiosulphates are decomposed by dilute mineral acids, with separation of S and evolution of SO₂. Salts of the acid used are formed, and H₂S₁O₂Aq, which then decomposes to H₂SO₂Aq and S. This reaction has been studied by several chemists (v. Berthelot, C. R. 108, 971; Vaubel, B. 22, 1686, 2703; Vortmann, B. 22, 2307; Foussereau, C.R. 104, 1842). Colefax (C. J. 61, 176) in 1892 examined the interaction of Na₂S₂O₂Aq and acids; he found that when the products of the change are not removed the interaction stops before all the Vox. IV.

H₂S₂O₂Aq is decomposed. C. determined the effects on the velocity and the limit of the reaction of concentration, increasing the relative mass of the acid used, the nature of the acid, temperature, and the presence of SO₂ in the solution before acidification (reference to older memoirs are given by C.).

Thiosulphates are converted into tetrathionates by the action of I; oxidisers such as HNO₄, KClO₂ and HClAq, and KMnO₄Aq produce sulphates. According to Hönig a. Zatzek (M. 4, 738), KMnO₄ in solution in acetic acid produces

dithionates.

For the general qualitative reactions of thiosulphates compared with those of thionates v. Debus (C. J. 53, 298).

SELENOTHIOSULPHATES M, SeSO₃ are described as Thioselenates, under Selenates (p. 434).

as Thioselenates, under Selenates (p. 434).

Constitution of thiosulphates. The formula
SO₂-OM.SM is generally adopted for the normal
thiosulphates of monovalent metals (Odling,
C. N. 29, 153; cf. Schorlemmer, C. J. [2] 7,
254; Williamson, C. J. [2] 7, 259). Some
chemists, considering the resemblances between
thiosulphates and sulphites, have suggested the

formula SO (v. Blomstrand, Die Chemis

der Jetztzeit [Heidelberg, 1869], 258; Buchanan, B. 3, 485; Michaelis, G. O. [5th ed.] 1, 745). The formation of thiosulphates by boiling sulphites with S is readily represented by either formula, according as sulphites are regarded as SO₂.OM.M or SO.OM.OM. As the balance of evidence is in favour of the formula SO2.OM.M, it is likely that the salts formed by adding S to the sulphites are SO_{2.}OM.SM. The production of (NH₄), \$2O₃ by the interaction of (NH₄), \$O₄ and P₂S₃ (Spring, B. 7, 1158) also confirms the formula SO_{2.}OM.SM, inasmuch as a thiosulphate is produced in this reaction from a sulphate (SO₂.ONH₄.ONH₄ giving SO₂.ONH₄.SNH₄). Spring's synthesis of H₂S₂O₃ from SO₃ and H₂S—for the actual reaction carried out, viz. formation of (NH1)2S2O2 by first combining SO, and NH,, and then interacting with H₂S, is equivalent to the synthesis of H₂S₂O₂ (v. S., l.c.)—is also in keeping with the formula SO₂.OM.SM. By warming EtBr with Na₂S₂O₃Aq, Bunte (B. 7, 646) obtained the salt EtNaS O. This salt is decomposed by dilute acids giving EtSH and SO...OH.ONa; when heated it gives off Et,S₂ and SO₂, leaving Na,SO₄. These reactions indicate that EtNaS₂O₃ is SO₄Na.EtS; hence the formula of Na₂S₂O₃ is probably SO₃Na.NaS. One of the Na atoms in Na S₂O₂ is then probably in direct union with an atom of S; and as EtNaS,O. easily gives SO. OH.ONa, along with EtSH, it is likely that the formula of this salt is SO₂ONa.SEt, and hence that the formula for Na₂S₂O₃ is SO₂.ONa.SNa. The formula SO. OH.SH makes possible the existence of two isomeric salts MM'S,O, viz. SO, OM.SM' and SO. OM'.SM; two such salts are known where M Na and M' = K. (v. Potassium Thiosulphate, p. 706).

Ammonium thiosulphate (NH₄)₂S₂O₅. Obtained by the interaction of (NH₄)₂CO₅ and CaS₂O₅A₆, filtering and evaporating (Rammelsberg, P. 56, 298); white, rhombic, deliquescent leaflets, crystallising with xH₂O. Fock a. Klüss (B. 22, 3099; 23, 534) obtained the anhydrous

salt by reacting on SO₂Aq with (NH₄)₂S₂, evaporating at a low temperature and finally over H₂SO₄, and thoroughly drying between filter paper (cf. Arppe, A. 96, 118). The anhydrous salt is not deliquescent (F. a. K.). Double salts are described by F. a. K. (l.c.; Double salts are described by F. a. K. (L.C.; also B. 23, 1753): (1) with OdS₂O₂; (2) with MgS₂O₂; (3) with K₂S₃O₃, the salt obtained was SO₂.OK.SNH₄. F. a. K. could not obtain SO₂.ONH₄.SK; (4) with Cu₂S₂O₃ and Cu₃I₂, also with Cu₂I₂ and NH₄I (Brun, C. R. 114, 667); (5) with PbS₂O₄ (Rammelsberg, P. 56, 808); (6) with HgS₂O₃ (R., L.c.).

Barium thiosulphate BaS₂O₄.aq. A white pp. obtained by mixing Ba(C₂H₄O₂)₂Aq and Na.S.O.Ag. and completing ppn. by alcohol:

Na.S.O.Aq, and completing ppn. by alcohol; loses all water at 170°, at red heat gives BaS, BaSO, BaSO, and S (H. Rose, P. 21, 437;

Rammelsberg, P. 56, 295)

Double salts. (1) With CdS₂O₂ (Fock a. Klüss, B. 23, 1753); (2) with Cu₂S₂O₂ (Cohen, C. J. 51, 38); (8) with PbS₂O₂ (Rammelsberg, P. 56, 808).

Cadmium thiosulphate CdS₂O₃. 2aq (F. a. K., B. 23, 534); double salts with (1) Na₂S₂O₃, (2) K.S.O., (3) SrS.O. (F. a. K., B. 23, 1753; Vortmann a. Padberg, B. 22, 2637).

Calcium thiosulphate CaS₂O₂. 6aq. Formed by mixing hot conc. solutions of CaCl, and Na.S.O., evaporating under 60°, separating NaCl, which crystallises first, and allowing to cool (Kessler, P. 74, 282). Also obtained by heating CaSO, and S with water, or by boiling milk of lime with S, passing in SO₂, and crystallising below 60° (Herschel, N. Ed. P. J. 1, 8, 296; 2, Forms large, clear, six-sided, triclinic prisms, that effloresce in air at 40°; S. 100 at 3° (H., l.c.). Conc. solution gives CaSO, and S by heating above 60°. Rammelsberg (P. 56, 808) describes a double salt with PbS,O.

Cobalt thiosulphate CoS.O. 6aq (Rammelsberg, P. 56, 898; Fock a. Klüss, B. 22, 3310). A double salt with Na.S.O. is described by Vortmann a. Padberg (B. 22, 2637).

Copper thiosulphates. According to von Hauer (W. A. B. 13, 448) an acid cuprous salt Cu.H.(S₂O₃), is obtained, in microscopic, goldenyellow needles, by adding conc. CuSO,Aq to Na,S,O,Aq till the liquid is deep yellow, and gently Na₂,O₂,Aq tin the liquid is deep yellow, and gently warming, washing the pp. with water and alcohol, and drying over H₂SO₂. Double salts of Cu₂S₂O₂: (1) with BaS₂O₃ (Cohen, C. J. 51, 38); (2) with K₂S₂O₃ (C., Lc.; Rammelsberg, P. 56, 321); (3) with Ag₂S₂O₃ (C., Lc.); (4) with Na₂S₂O₃ (Lenz, A. 40, 99; Siewert, J. 1866; 257; Kessel, R. 10, 1877, 2000; Schätter Lenz, 70, 240. Vert B. 10, 1677, 2000; Schütte, J. pr. 70, 249; Vortmann, M. 9, 165).

Iron thiosulphate. The ferrous salt FeS.O. 5aq was obtained by Vortmann a. Padberg (B. 22, 2637) by rubbing BaS.O. with conc. FeSO, Aq (the salts being present in the ratio of equal numbers of molecules), filtering from BaSO, adding a mixture of alcohol and ether, and allowing the green, oily liquid that separated to stand for some days. Small green crystals; very sol. water; solution is decomposed by boiling, with separation of FeS. A double salt with Na,S,O, is described by V. a. P. (i.e.). For accounts of former attempts to prepare pure ferrous thiosulphate v. Fordos a. Gélis (J. Ph. [8] 4, 888); Berthelot (A. Oh.

[5] 2, 58); Vogel (J. pr. 8, 102); Koene (P. 68, 245, 481); Rammelsberg (P. 56, 306).

No ferric thiosulphate has been isolated. Lead this sulphate PbS₂O₃. Obtained by adding Na₂S₂O₂Aq to Pb(NO₂)₂Aq, or Pb(C₂H₃O₃)₂Aq, and washing the pp. with cold water. A white solid; very sl. sol. water; sol. solutions of alkali thiosulphates (v. Pape, J. 1864. 57; Letts, C. J. [2] 8, 424). For reaction with POl₃—giving PbOl₂, HCl, SO₂, SOCl₂, POCl₃ and probably PSOl₃—v. Buchanan (B. 3, 485). Rammelsberg (P. 56, 308) has described double salts with the thiosulphates of NH, Ba, Ca, K (v. also Spring, B. 7, 1157), and Na (v. also Vortmann a. Padberg, B. 22, 2637).

Lithium thiosulphate Li.S.O. Saq (v. Fock a.

Klüss, B. 22, 3096)

Magnesium thiosulphate MgS2O2.6aq. By passing SO, into MgSAq till colourless, boiling MgSO, in SO,Aq with S, or adding BaS,O,Aq to MgSO Aq and filtering, a solution is obtained which on evaporation at a low temperature gives small, white, prismatic crystals of this salt. Loses 8H₂O at 170°, and gives off SO₂ and S at red heat, leaving MgSO₄ and MgO (Herschel, N. Ed. P. J. 1, 8, 296; 2, 154; Rammelsberg, 150° P. 56, 303; Fock a. Klüss, B. 22, 8310). Double salts (1) with (NH₄)₂S₂O₃, (2) with K₂S₂O₃ are described by R. (l.c.; v. also Kessler, P. 74, 256; also F. a. K., B. 23, 584).

MnS.O. 5aq. thiosulphate Manganese Formed similarly to FeS.O₃. 5aq (Vortmann a. Padberg, B. 22, 2637). A double salt with Na.S.O₃ is described by V. a. P. (l.c.).

Mercury thiosulphates. Neither the mercurous nor the mercuric salt has been isolated. Addition of a thiosulphate to solution of a mercurous salt ppts. Hg,S; with a mercuric salt, a compound of HgS with the mercuric salt employed is said to be formed (Rammelsberg, P. 56, 314). Double salts of mercurous thiosulphate with (1) (NH₄)₂S₂O₃, (2) Cu₂S₂O₃, (8) Na₂S₂Ô₃ are described by Rammelsberg (*l.c.*); and a double salt of mercuric thiosulphate with K2S2O2 by Vortmann a. Padberg (B. 24, 1351).

Nickel thiosulphate NiS.O. 6aq (Rammelsberg, P. 56, 309; Fock a. Klüss, B. 22, 3310). For a double salt with Na.S.O. v. Vortmann

a. Padberg (B. 22, 2637).

Platinum thiosulphate. Not isolated; but Schottländer (A. 140, 200) describes a double

salt PtS,O,.3Na,S,O,.10aq.

Potassium thiosulphate K.S.O. Obtained by boiling conc. K2SO, with excess of S; or by adding hot K2Cr2O,Aq to hot K2S,Aq, little by ittle, waiting after each addition till the separated Cr₂O₃ is pure green in colour (2K₂S₃Aq + 4K₂Cr₂O₄Aq + H₂O = 5K₁S₂O₄Aq + 2KOHAq + 4Cr₂O₃); (Döpping, A. 46, 172; Kessler, P. 74, 274). In

either case the filtered solution is evaporated to the crystallising point, and the crystals of K₂S₂O₃.xH₂O are heated to c. 180°. By evaporating at 30°, thin, four-sided, deliquescent prisms are obtained of the hydrated salt 3K₂S₂O₂.H₂O (Rammelsberg, P. 56, 296; Kessler, l.c.; Fock a. Klüss, B. 22, 3096). This hydrate is very sol. water, but insol. alcohol; it loses all water at c. 180° ; heated more strongly it decomposes to K_2S_2 and K_2SO_4 (B., *l.c.*). Other hydrates are described by D. a. P., *l.c.* (v. also Plessy, A. Oh.

[8] 11, 182; F. a. K., l.c.). The final products of the action of SO₂ on K₂S₂O₃Aq are K₂S₂O₄Aq, K₂S₂O₅Aq, and K₂S₂O₅Aq (Debus, C. J. 58, 343).

Double salts.—(1) with (NH₂)₂S₂O₅ (Fock a. Klüss, B. 23, 534); the salt obtained was SO₂.OK.SNH₄, the isomeric salt SO₂.ONH₄.SK could not be isolated; (2) with Bi₂(S₂O₃) (Carnot, C. R. 83, 838); (3) with CdS₂O₃ (F. a. K., B. 23, 534); (4) with Cu₂S₂O₃ (Cohen, C. J. 51, 88); (5) with HgCy₂ (F. a. K., B. 24, 1351); (6) with HgS₂O₃ (F. a. K., l.c.); (7) with PbS₂O₃ (Rammelsberg, P. 56, 308; Spring, B. 7, 1157); (8) with Ag,S₂O₃ (Cohen, C. J. 51, 38; Schwicker, B. 22, 1728, describes KAgS₂O₃.NH₃). (9) With Na₂S₂O₃ two salts are obtained: (i.) SO₂.OK.NaS. 2aq, by the interaction of Am₂S₂Aq and NaKSO₃, 2aq; (ii.) SO₂.SK.ONa. 2aq, by the interaction of Am₂S₃Aq and KNaSO₃ (c. SULPHITES, p. 587); the first salt melts at c. 57° and has S. = 218·7 at 15°, heated with EtBr it gives SO₂.OK.SEt; the second salt melts at c. 62° and has S. = 105·3 at 15°, heated with EtBr it gives SO₂.ONa.SEt(Schwicker, B. 22, 1728).

Silver thiosulphate Ag₂S₂O₃. This salt has probably not been isolated pure. On adding dilute AgNO₃Aq to K₂S₂O₄Aq a white pp. is formed, and is dissolved on shaking; addition of more AgNO₂Aq produces a greyish, flocculent pp., which is a mixture of Ag₂S₂O₃ and Ag₂S. By treating this pp. with NH₄Aq, filtering from Ag₂S, exactly neutralising the filtrate by HNO₃Aq, and quickly drying the white flocculent pp. that formed between filter paper, Herschel (N. Ed. P. J. 1, 26, 398; 2, 154) obtained fairly pure Ag₂S₂O₃. A white powder; slightly soluble water; solution has a sweet taste; decomposes

readily to Ag₂S and H₂SO₄Aq.

Double salts. Ag,0 and Ag salts generally dissolve readily in solutions of alkali thiosulphates, forming double salts, either of the form 2AgMS₂O₃, aq, or of the form Ag₂M₄(S₂O₃), 2aq (Herschel, l.c.). (1) With (NH₄)₂S₂O₃; by adding excess of AgOl to (NH₄)₂S₂O₃Aq, and washing the white crystalline pp., 2Ag(NH₄)S₂O₃, aq is obtained. Using less AgCl and ppg. the solution by alcohol produces Ag₂(NH₄)₄(S₂O₃), 2aq (H., l.c.). (2) With Na₂S₂O₃; addition of AgNO₃Aq, drop by drop, with constant shaking, to Na₂S₂O₃Aq, till a permanent pp. begins to form, followed by addition of alcohol, ppts. Ag₂Na₄(S₂O₃), 2aq; the salt is washed with alcohol, and dried in vacuo (Lenz, A. 40, 94); when the mother-liquor from this salt is allowed to evaporate 2AgNaS₂O₃-aq is deposited (Herschel, l.c.; cf. Lenz, l.c.). The second of these salts is much less soluble in water than the first; it dissolves in Na₂S₂O₃Aq, with formation of Ag₂Na₄(S₂O₃)₃. 2aq. Both salts are decomposed by boiling their aqueous solutions, with formation of Ag₂S. (3) With K₂S₂O₃; two salts are known similar to the sodium salts. Schwicker (B. 22, 1788) describes the compound AgKS₂O₃NH₄. Herschel (l.c.) obtained double salts with OaS₂O₃ and with SrS₂O₃.

Sodium thiosulphate Na₂S₂O₃. 5aq.

Formation.—1. By boiling Na₂SO₃Aq with S.—2. By passing SO₃ into solution of sodium polysulphides.—3. By the action of SO₂ on sodium waste suspended in water, whereby CaS₂O₃Aq is formed; followed by addition of Na₂SO₄, filtration from CaSO₄, and evaporation.

Preparation.—Na, SO, and is dissolved in a little water, the solution is boiled with flowers of sulphur for a considerable time, and filtered; the clear, colourless liquid is evaporated to the crystallising point, and the crystals that separate are purified by recrystallisation from water.

Properties.—Large, white, monoclinic prisms; S.G. 1.736 at 10° (Kopp, J. 8, 45; cf. Schiff, J. 12, 41; Smith, Am. J. Pharm. 58, 148). Melts at 45° in the water of crystallisation (Kopp, J. 8, 45); at 48.5° (Tilden, C. J. 45, 268). The crystals are dehydrated by keeping in vacuo over H.80, or by heating to 100°. S. 49.75 at 0°, 69.44 at 20°, 104.16 at 40°, 192.3 at 60° (Kremers, P. 99, 50). An aqueous solution of the salt is easily supersaturated; thus Schiff (A. 118, 350) dissolved 171 parts Na,S,O,. 5aq =108.9 parts Na₂S₂O₄) in 100 parts water at 19.1°, and Kremers obtained a solution of 217.4 parts Na₂S₂O₃ in 100 parts water at the ordinary temperature. According to Baumhauer (J. pr. 104, 148), a supersaturated solution may be cooled to -10° without crystallising; addition of a crystal of the salt causes rapid crystallisation, with the evolution of much heat (cf. Parmentier a. Amat, C. R. 98, 735). Solution of Na₂S₂O₃. 5aq in water is accompanied by the disappearance of much heat; Rüdorff (B. 2, 68) says that 110 parts Na₂S₂O₂.5aq dissolving in 100 parts water lower the temperature from 10.7° to -8° , i.e. through 18.7° . Schiff (A. 113, 350) gives the following data for S.G. and composition of aqueous solutions of Na₂S₂O₂.5aq:-

•	•				•
8.G. of	P.o.	P.c.	B.G. of	P.o.	P.c. Na.S.O.
1.0052	,S,O,. 5aq 1	0.637	1.1440	8,0,.5aq 26	16·564
1.0105	2	1.274	1.1499	27	17.201
1.0158	8	1.911	1.1558	28	17.838
1.0211	4	2.584	1.1617	29	18.475
1.0264	5	3.185	1.1676	80	19.113
1.0317	6	3.822	1.1738	81	19.750
1.0370	7	4.459	1.1800	32	20.387
1.0423	8	5.096	1.1862	33	21.024
1.0476	9	5.733	1.1924	84	21.661
1.0529	10	6.371	1.1986	85	22.298
1.0639	11	7.008	1.2048	86	22.935
1.0584	12	7.645	1.2110	87	28.572
1.0695	13	8.282	1.2172	38	24.209
1.0751	14	8.919	1.2234	39	24.846
1.0807	15	9.556	1.2297	40	25.484
1.0863	16	10.198	1.2362	41	26.121
1.0919	17	10.830	1.2427	42	26.758
1.0975	18	11.476	1.2492	43	27.395
1.1031	19	12.105	1.2558	44	28.032
1.1087	20	12.742	1.2624	45	28.669
1.1145	21	13.379	1.2690	46	29.306
1.1204	22	14.016	1.2756	47	29.948
1.1263	23	14.658	1.2822	48	80.580
1.1322	24	15.290	1.2888	49	81.218
1.1381	25	15.927	1.2954	50	81.855
2 2001		20 0001			J_ 000

Na₂S₂O₃.5aq is insol. alcohol.

According to Jochum (C. C. 1885. 642), a trihydrated salt Na.8.0, 3aq is obtained by adding alcohol to a saturated solution of the ordinary salt, removing the oily liquid that seps rates, and adding to it 98 p.c. alcohol. According to Parmentier a. Amat (C. R. 98, 735), Ns.8.0, 5aq is obtained crystallised in needles, malting at c. 32°, by placing a supersaturated

solution of the ordinary salt in a freezing mixture in a closed vessel.

Reactions.—1. Heat decomposes Na2S2O2; the change begins at c. 220° with separation of the change begins at c. 220° with separation of S, at c. 400° Na₂SO₄ and Na₂S₅ are formed, and at a higher temperature more S is given off, and Na₂SO₄ and Na₂S remain.—2. Na₂SO₅Aq interacts with iodine in KIAq to form NaIAq and Na₂S₄O₆Aq; according to Villiers (C. R. 108, 402) some Na₂S₃O₆ is also formed. With a mixture of Na.SO, and Na.S.O. in solution I produces NaIAq, Na.SO.Aq and Na.S.O.Aq (Colefax, C. J. 61, 198, 1083).—3. Na₂S₂O₃ is oxidised by potassium permanganate solution; in alkaline solutions the oxidation is complete to Na,SO,Aq according to Hönig a. Zatzek (M. 4, 788); Glaser (M. 6, 329) says that the oxidation is also complete in neutral solutions; Luckow (Fr. 32, 53) represents the oxidation, on boiling, either in neutral, acid, or alkaline solution as producing Na₂SO₄ and Na₂S₂O₅.—4. Acids interact with Na₂S₂O₅Aq, forming Na salts of the acids, and H₂S₂O₃Aq, which then decomposes to H₂SO₃Aq and S; when the products are not removed from the solution the reaction stops before all the H₂S₂O₃ is decomposed (Colefax, C. J. 61, 176; and v. beginning of article THIOSULPHATES, p.

Combinations.—1. With water (v. supra, Properties).—2. With various thiosulphates to form double salts. Double salts with CdS₂O₈, CoS₂O₃, FeS₂O₃, PbS₂O₃, MnS₂O₃, NiS₂O₃, Tl₂S₂O₃, and ZnS₂O₃ are described by Vortmann a. Padberg (B. 22, 2637); Fock a. Klüss (B. 23, 1753) also describe that with CdS₂O₈. Rammelsberg (P. 56, 308) prepared salts with PbS₂O₃ and Hg₂S₂O₃. For a double salt with HgS₂O₃ v. Schwicker (B. 22, 1728). For the double salts with K2S2O3 v. Potassium thiosulphate, double salts, No. (9) (p. 707), and for those with Ag₂S₂O₃ v. SILVER THIOSULPHATE, double salts, No. (2) (p. 707). Schottländer (A. 140, 200) describes a double salt 3Na,S,O,.PtS,O,. 10aq.
Strontium thiosulphate SrS,O,.5aq.

pared like the Ba salt (Herschel, N. Ed. P. J. 1, 26, 398; 2, 154; Kessler, P. 74, 274). Fock a.

Klüss (B. 22, 3310) describe a salt SrS₂O₃, aq.

Thallium thiosulphate Tl₂S₂O₃. A white, crystalline pp., formed by adding Na₂S₂O₃Aq to fairly conc. TIOHAq (Crookes, C. J. 17, 136; Hebberling, A. 134, 11). For a double salt with Na,S₂O, v. Werther (J. pr. 92, 150); Vortmann a. Padberg (B. 22, 2637); also Jochum (C. C. 1885. 642)

Zine thiosulphate ZnS₂O₄. Prepared by double decomposition from ZnSO₄Aq and BaS₂O₂Aq (v. Rammelsberg, P. 56, 295; Fordos a. Gélia, J. pr. 29, 291). For a compound with NH, v. B. (l.c.). For a double salt with Na,S,O, v. Vortmann a. Padberg (B. 22, 2687).

М. М. Р. М. THIOSULPHURIC ACID H.S.O.Aq. acid almost certainly exists in the liquid obtained by adding a dilute weak acid to dilute Na₂S₂O₂Aq, the acid and salt being mixed in equivalent quantities; the H.S.O.Aq soon begins to decompose to H.SO.Aq and S. Landolt (B. 16, 2958) has made an extensive investigation into the conditions and time of existence of H2S2O, in solution. Thomsen (Th. 2, 256) gives the thermal data; $[SO^2Aq,S] = -9810$; $[SO^2,S,Aq] = -1610$. For the salts of H₂S₂O₃ v. Thiosulphates. M. M. P. M.

THIOTOLENE v. Methyl-thiophene. THIO-o-TOLUIC ACID. Amide

O.H.Me.CS.NH2. [88°]. Formed from o-toluic

nitrile (Gabriel a. Heymann, B. 24, 786). Thio-p-toluic acid. Amide. [168°]. Formed by passing H₂S into an alcoholic solution of p-toluic nitrile (Paterno a. Spica, B. 8, 441). Yellowish needles. Zinc and HClAq yield C.H.Me.CH,NH

THIO-TOLUIDINE v. DI-AMIDO-DI-TOLYL SUL-

THIO-p-TOLYL-DI-PHENYL-DI-GUAN-IDINES v. DI-PHENYL-DI-GUANIDO-DI-TOLYL SUL-PHIDE and DI-PHENYL-DI-CARBIMIDO-TETRA-AMIDO-TETRA-TOLYL DISULPHIDE.

THIO-TOLYL-DI-PHENYL-THIO-UREA v. DI-PHENYL-DI-THIOURAMIDO-DI-TOLYL SULPHIDE.

THIO-URAMIDO-BARBITURIC ACID

 $C_bH_eN_4SO_s$ i.e. $NH_2.C(NH).S.CH < CO.NH > CO.$ Formed by the action of thio-urea on an aqueous solution of chloro-, bromo-, or di-bromo-barbi-turic acid (Mulder, B. 12, 2309; Trzcinski, B. 16, 1057). Formed also by heating alloxan with thio-urea and alcoholic SO, at 100° (Nencki, B. 4, 722; 5, 451). Slender needles, insol. water, sol. hot HClAq. Converted by boiling KOHAq into thio-dialuric acid $C_4H_4N_2SO_2$ $1\frac{1}{2}aq$ (?) which yields KA'aq. H_2SO_4 ($\frac{1}{2}$ pt.) at $150^\circ-160^\circ$ forms uro-sulphinic acid $C_3H_4N_4SO_2$.

THIO-URAMIDO-BENZENE p-SULPHONIC ACID. The salt NH₂·CS.NH.C₆H₄·SO₃K, formed by heating potassium sulphocyanide with amidobenzene p-sulphonic acid at 140°, crystallises from dilute alcohol in aggregates of thin needles (Pellizzari, A. 248, 156).

m-THIO-URAMIDO-BENZOIC ACID

NH₂.CS.NH.C₆H₄.CO₂H. [187°]. Formed from m-amido-benzoic acid and potassium sulpho-cyanide (Arzruni, B. 4, 406). Formed also by allowing a solution of m-cyanamido-benzoic acid in colourless ammonium sulphide to stand for 24 hours (Traube, B. 15, 2118). Needles. Converted by ammoniacal AgNO, into muramido-benzoic acid. Not coloured by FeCl,

p-THIO-URAMIDO-BENZYL-THIO-UREA NH₂.CS.NH.C₆H₄.CH₂.NH.CS.NH₂. p- ω -di-thiuramido-toluene. [176°]. Formed by heating p-amido-benzyl-amine hydrochloride (1 mol.) with potassium sulphocyanide (2 mols.) in aqueous solution (Amsel a. Hofmann, B. 19, 1289). White needles.

THIO-URAMIDO-BUTYRIC ACID. Anhy-NH:C<S-CHEt. [200°]. dride from a-bromo-n-butyric acid and thio-urea (Andreasch, M. 8, 419). Needles, v. sol. hot water.

Thio-uramido-iso-batyric acid. Anhydride NH:C $<_{\rm NH.CO}^{\rm S-OMe_2}$. C₂H₈N₂SO i.6. [242°]. Formed from bromo-isobutyric acid and thiourea (Andreasch, M. 8, 410). Plates, sl. sol. Oxidised by HCl and KClO, to urea and a-sulpho-iso-butyric acid.

o-THIO-URAMIDO-CINNAMIC ACID [236°-239°]. NH₂.CS.NH.C₆H₄.CH:CH.CO₂H. Got by heating the sulphocyanide of o-amido-cinnamic acid C₀H₄(NH₂.HSCy)OH:CH.CO₂H (152°) at 115° (Rothschild, B. 23, 3342). V.

sol. hot HOAc and NH,Aq.

a-Thio-ψ-uramido-cinnamic acid C₁₀H₁₀N₂SO₂ NH:C(NH₂).S.C(CO₂H):CHPh. Formed from benzoic aldehyde, thiohydantoïn, and NaOHAq (Andreasch, M. 8. 421). Needles (from alcohol), decomposed by alkalis.

β-THIO-ψ-URAMIDO-PROPIONIC ACID

NH:C(NH₂).S.CH₂.CH₂.CO₂H. Formed by heating \$-iodo-propionic acid with thio-urea and water and a little NaOH (Andreasch, M. 6, 831). Small pyramids (containing 2aq), sl. sol. cold water and cold alcohol. Decomposed by baryta into cyanamide and CH2(SH).CH2.CO2H. KClO3 and HCl yield β-sulpho-propionic acid.

ωp-DI-THIO-URAMIDO-TOLUENE v. THIO-

URAMIDO-BENZYL-THIO-UREA. DI-p-THIO-URAMIDO-DI-TOLYL-o-SUL-PHIDE [2:1:4] S(C₆H₂Me.NH.CS.NH₂)₂. Thiop-tolyl-di-thiurea. [121°]. Formed by evaporating to dryness an aqueous solution of di-pamido-di-tolyl sulphide, hydrochloride and ammonium sulphocyanide (Truhlar, B. 20, 669). White amorphous powder. Sol. alcohol and benzene, sl. sol. ether.

THIÓ-UREA CSN₂H₄ *i.e.* CS(NH₂)₂ or HS.C(NH₂):NH. Mol. w. 76. [170°]. S. 9 in

the cold (Volhard, J. pr. [2] 9, 13).

Formation.—1. By heating dry ammonium sulphocyanide for two hours at 170°, cooling to 100°, adding an equal weight of water at 80°, filtering, and allowing to crystallise (Emerson Reynolds, C. J. 22, 1; Claus, A. 179, 113; Smolka a. Friedreich, M. 10, 90).—2. By the action of tin and HCl, or of HI, on persulphocyanic acid C₂H₂N₂S₃ (Glutz, A. 154, 39).— 3. By passing dry H₂S into an ethereal solution of cyanamide (Baumann, B. 6, 1375).-4. By allowing a solution of cyanamide, mixed with yellow ammonium sulphide, to stand for a day (Baumann, B. 8, 26).-5. From cyanamide and thioacetic acid. - 6. Together with urea by the action of ammonia on COS (Berthelot, C. R. 94, 1069).

Properties.—Silky trimetric needles, melting at 170⁵. After one fusion it melts at 149⁶ (Prätorius, *J. pr.* [2] 21, 141). V. sol. water, nearly insol. alcohol and ether. Tastes bitter, not coloured by FeCl_s. According to Armstrong (C. J. Proc. 3, 2), the molecule of thio-ures is

C282N4H8

Reactions.—1. Water at 140° forms ammonium sulphocyanide. Alcohol at 100° acts in like manner (Haller, Bl. [2] 45, 706).—2. An alcoholic solution of nitrous ether quickly converts it into ammonium sulphocyanide. - 3. Prolonged heating at 170° also converts it into ammonium sulphocyanide. At 180° it slowly forms guanidine sulphocyanide and ammonium tri-thio-carbonate. Nitrous acid gives off nitrogen.-4. KOH at 100° slowly forms NH, KSH, K2CO2, and a little KCyS. Alcoholic potash acts in like manner (Haller, C. R. 102, 974). H.SO, and HCl act in the same way. -5. Aqueous KMnO, forms urea and other products (Maly, M. 11, 277).—6. Oxidising agents (e.g. H₂O₂) K₂Cr₂O₇, KNO₂, KMnO₄, and iodine) form S₂(C(NH₂):NH)₂, which has an insoluble nitrate (Storch, M. 11, 452).—7. Ammoniacal AgNO₃ forms ures and Ag₂S.—8. HgO, added to a cold aqueous solution, produces cyanamide.

PhO acts in like manner. At 100° the product is dicyandiamide (Hofmann, B. 2, 605).-9. Acetoacetic ether reacts forming an ether C₅H₇N₂SO(OEt), which on saponification yields CS NH.CO CH, v. sl. sol. water (Behrend, B. 19, 219).—10. CCl, COCl forms C2H2Cl,N2S2O, crystallising in slender needles, m. sol. alcohol, sl. sol. water, v. sl. sol. ether; converted by Br into CCl₃.SO₂Br (McGowan, C. J. 49, 191; 51, 669; J. pr. [2] 36, 220). CCl, SO, Cl added to an alcoholic solution of thio-urea forms the chloride Cl2CSN2H4, which is ppd. on adding ether, while the filtrate contains the compound $(CSN_2H_4)_2(CCl_3.SO_2)_2$ [125°], which is more sol. alcohol than water, and gives with Br a pp. of CCl_s.SO₂Br.-11. CCl_s.SO₂NH₄ in alcohol forms flat prisms of (CSN₂H₄)CCl₃SO₂H [139°], m. sol. water and ether, v. sol. alcohol, converted by Br into CCl₃.SO₂Br (M'Gowan, C. J. 51, 667; J. pr. [2] 36, 219).—12. Chloro-acetone forms (NH_,CS.NH.CH_,CO.CH_,)HCl [126°] (Paw-lewski, B. 21, 402).—18. ClCO,Et forms (NH_,CS.NH.CO_,Et)HCl [117°].—14. CH_,Cl.OH gives C2H, N2S, a white amorphous powder, insol. water, nearly insol. all usual menstrua, reconverted into thio-urea by HClAq (von Hemmelmayr, M. 12, 89).-15. Di-bromo-succinic acid forms fumaric acid on heating (Nencki a. Sieber, J. pr. [2] 25, 74).—16. A hot cone solution of di-bromo-pyruvic acid forms 'sulphuvinuric acid' C,H, N2SO2, which crystallises with 2aq, reduces Fehling solution to Cu2O in the cold, and to a copper mirror on warming, and yields the salts CaA'₂, MgA'₂, ZnA', HA'HCl, HA'HBr, and HA'HNO₃ aq (N. a. S.).—17. Bromo-pyruvic ether forms amido-thiazole carboxylic ether $S < CH = CCO_2Et \ [173^{\circ}] \ (Steude, A. 261, 30).$

18. Bromo-acetoacetic ether gives μ -amidothiazyl-acetic ether $S \subset CH \xrightarrow{CH} C.CH_2.OO_2Et$

[94°], which yields an acid [c. 130°], split up on fusion into CO₂ and amido-methyl-thiazole (Steude). a - Chloro - acetoacetic ether forms C₂NSMe(NH₂).CO₂Et [175°] (Zürcher, A. 250, 281).—19. Thio-urea heated with an alcoholic solution of chloro-acetic acid and phenyl-hydrazine forms C,H,N,SO [175°] (Probst, J. pr. [2] 45, 416).—20. Fusion with (B)-iodopropionic acid and a little water forms C, H, N, SO, [176°], which crystallises from water in needles (containing 2aq), sl. sol. alcohol, insol. ether (Andreasch, M. 6, 832) .- 21. Di-chloro-acetic acid forms thiohydantoin. a-Bromo-propionic acid forms methyl-thiohydantoin (206° uncor.) (Dixon, C. J. 63, 815).—22. Aldehyde heated in a sealed tube with thio-urea forms CSN₂H₂(CHMe), which is sl. sol. ether and cold alcohol, and is decomposed by hot water (Reynolds, C. N. 24, 87).-23. Succinic anhydride at 140° forms $C_{b}H_{b}N_{2}SO_{s}$ [211°].—24. Citraconic anhydride at 130° forms $C_{b}H_{b}N_{2}SO_{s}$ [223°] (Pike, B. 6, 1104).— 25. Acetyl-acetone and a small quantity of HClAq form CH₂(CMe:N.CS.NH₂)₂HCl [219°]. In presence of excess of HCl the product is CH₂ CMe:N CS [210°] (Evans, *J. pr.* [2] 48, 503).

Crystalline. - B'HCl. Salts .- B'HNO. Formed from the stannous double chloride and

Crystalline.—B'HI. Tables.—B',NH,Cl. [154°]. Felted crystalline mass (Reynolds, O. J. 59, 885).—B',NH,I. [186°].—B',H,PtCl., Yellow prisms, v. sol. water and alcohol. Can be dried at 100°.-B',HPtCl,. Got when saturated solutions of thio-ures and PtCl, are mixed. Dark-red prisms, insol. water, alcohol and ether.— B'_AuCl. Yellow monoclinic crystals, got by adding AuCl, to a saturated solution of thiourea.—B'TISO,.—B'2CuSO,. White silky needles. Got by mixing conc. solutions of thio-ures and CuSO₄.—B',Cu₂(SO₄) 2aq.—B',CuCl. Colourless dimetric crystals, sol. water, but ppd. by HCl and KCl. Alkaline in reaction. Attacked by H₂S with difficulty. Does not absorb CO (Rathke, B. 17, 301).—B'2Cu2Cl2 aq. Ppd. by adding CuCl,Aq to aqueous thio-urea (Rathke, B. 14, 1779; 17, 301). Minute needles, insol. water.—B',CuCl. Flat needles. Formed by mixing the two preceding salts into which it is decomposed by water.—B'4HgO.—B'2(HgO), 3aq. Ppd. by adding neutral Hg(NO₁)₂ to a dilute solution of thio-ures (Emerson Reynolds, A. 150, 235; Kurnakoff, B. 24, 3956). Crystalline. Loses thio-urea on washing with water.— B',HgCl₂: large crystals, v. sol. water.—B',HgCl₂: minute needles, nearly insol. water (Claus, B. 9, 227).—B'HgI₂ (Maly, B. 9, 173).—B'HgCyS. Needles, decomposed by heat into dicyandiamide, Needles, decomposed by heat into dicyandiamide, HCy, and HgS (Nencki, B. 6, 598).—B',PbCl₂: needles (Claus, A. 179, 135).—B'Pb(OyS)₂—B'₂SnCl₂.—B'₂SnCl₂: spherical groups of prisms (from water).—B'₂CdSO₄.—B'₂SiBr₄ (Emerson Reynolds, C. J. 51, 202).—B'₂Ag,O 4aq. Crystalline.—B'AgNO₃ (K.).—B'₂AgNO₃. [141°]. Silky crystals, sol. hot alcohol (Emerson Reynolds, C. J. 61, 249).—B'₂AgCl. [171°]. Needles, m. sol. hot alcohol.—B'₂AgPr. [121°].—B'₂AgL.—B'₂AgCy. [126°].—B'₄Ag₂C₂O₄: needles.

Chloride (CSN₂H₄)₂Cl₂. Formed by passing Cl into a conc. alcoholic solution of thioures (Claus, A. 179, 189). Formed also by the action of CCl₂.SO₂Cl₃ and of ICl, ICl₃, or POCl₄

action of CCl, SO, Cl, and of ICl, ICl, or POCl, on thio-urea (M Gowan, C. J. 49, 191; J. pr. [2] 83, 188). Small needles, v. sol. water, m. sol. alcohol, insol. ether. Decomposed by alkalis, yielding cyanamide. Alcoholic KI removes chlorine, setting thio-urea free. Dilute HNO, forms crystals of (CSN₂H₄)₂(NO₂)₂, which is sl. sol.

water, insol. alcohol and ether.

Bromide (CSN₂H₄)₂Br₂. Formed by adding Br to an alcoholic or cold aqueous solution of thio-urea; in aqueous solution excess of Br produces urea (Claus, A. 179, 138; M'Gowan, C. J. 51, 378; J. pr. [2] 36, 216). Crystals. Its aqueous solution deposits S when heated. Sodium-amalgam reproduces thio-urea.

Iodide (CSN₂H₄)₂I₂. Formed from thioures, I, and alcohol (M'Gowan, J. pr. [2] 83, 192). Prisms.

Methylo-iodide (CSN,H,)MeI. [117°]. Formed slowly by combination of thio-urea with MeI in the cold (Bernthsen a. Klinger, B. 11, 493). Prisms, v. sol. water and alcohol. gives the chloride, which forms the platinochloride (B'MeCl),PtCl, aq

Ethylo-chlorids B'EtCl. Crystals.

Ethylo-bromids (CSN.H.)EtBr. Hexa-gonal plates (Claus a. Siegfried, A. 179, 145). Decomposes at 100°.

Ethylo-iodides (CSN,H,)Etl. Crystal-

line. Gives with moist Ag.O a base. Boiling aqueous alkalis liberate mercaptan.—B', Et, PtCl. (dried at 100°). Tables.—B'zEtI. Needles (from alcohol) (Claus, B. 8, 41).

Bensylo-chloride (CSN,H4)C,H,C!.[168°]. Formed from benzyl chloride and thio-urea. Yields unstable benzyl-thio-urea CSN₂H₄C₇H₇ [72°], which decomposes at 100° into dicyandiamide and benzyl mercaptan (Bernthsen a. Klinger, B. 12, 574).—(CSN₂H₄C₇H₇Cl)₂PtCl₄.

Compound with oxalic etner (CSN,H₄)₂Et,C₂O₄. [150°]. Monoclinic crystals (Nencki, B. 7, 780). Decomposed by hot water.

Compounds with amines

(CSN,H4)4NMeH2Br. [138°]. Formed from methylamine hydrobromide and thio-ures in alcohol (Reynolds, C. J. 59, 392). Felted crystals.—(CSN₂H₄), NEt₂H₂Br. [134°]. Formed by boiling thio-urea together with diethylamine hydro-bromide and absolute alcohol .-(CSN₂H₄),NEt₃HBr.—(CSN₂H₄)₂NEt₄I [185°]. Prisms.—(CSN H4)2NEt4Br [160°]. Rectangular

prisms (Reynolds, C. J. 59, 388).

Acetyl derivative CSN,H,Ac. Mol. w.
118. [165°]. Formed by heating thio-ures with Ac₂O (Nencki, B. 6, 598). Formed also from thio-acetic acid and cyanamide (Prätorius, J. pr. [2] 21, 147). Prisms (from hot water), v. sol. alcohol, m. sol. ether.—BH,PtCl. Crystalline.

Propionyl derivative (Freytag, J. pr.

[2] 20, 881).

Benzoyl derivative CSN2H3Bz. [170°]. Formed by warming thio-urea (2 mols.) with BzCl (1 mol.) (Pike, B. 6, 755). Got also from benzoyl sulphocyanide and NH, Aq (Miguel, Bl. [2] 25, 252). Needles (from alcohol); sl. sol. cold water. Tastes bitter.

o-Oxy-bensoyl derivative NH₂CS.NH.CO.C_eH₄.OH. [182°]. S. (boiling alcohol) 7. Formed from salicyl thiocarbimide NH, CS.NH.CO.C, H, OH. and aqueous NH, (Miguel, A. Ch. [5] 11, 304).

THIO- V-URIC ACID v. THIO-URAMIDO-BARBI-

TURIC ACID.

THIO-VALERIC ALDEHYDE C, H, S. (115°). Formed, together with C₅H₆S₆ [95°], by heating isovaleric aldehyde (23 pts.) with S (9 pts.) in sealed tubes at 250° for eight hours (Barbaglia, G. 11, 95; B. 17, 2654). Oil with penetrating odour like onions, son not combine with NaHSO, combine with NaHSO. [69°]. odour like onions, sol. alcohol and ether. Does

 $V.D. \quad (H=1)$ 50.76. Formed by passing H₂S into a dilute aqueous solution of isovaleric aldehyde. Fibrous

crystals (from ether), insol. water.

THIOXENE v. DI-METHYL-THIOPHENE. THIOXYLENOL v. XYLYL MERCAPTAN.

THIOXYL METHYL KETONE v. DI-METHYL-THIENYL METHYL KETONE.

THORIUM Th. (Thorinum.) At. w. 232. Mol. w. unknown. The m.p. of Th has not been determined; it is certainly very high (Nilson, B. 15, 2541). S.G. 11·1 at 17° (N., l.c., p. 2543). S.H. ·02787 (N., B. 16, 161). Crystallises in regular octahedral and hexagonal forms, isomorphous with Si (Brögger, Z. K. 7, 442; of.

Nilson, B. 15, 2546; 16, 162 note). S.V.S. 20-9. *Historical.*—In 1818 Berzelius announced the discovery of the oxide of a new element in a rare Norwegian mineral; to the metal of this oxide he gave the name thorium (from THORIUM. 711

the Scandinavian god Thor). A few years later Berzelius found that the substance he had supposed to be a new earth was really yttrium phosphate. In 1828, however, Berzelius isolated a new earth from another Norwegian mineral; he named the new earth thoria, and the mineral from which it had been prepared he called thorite (P. 16, 385). In 1851 Bergemann discovered a new metal and called it donarium; but this metal was shown to be identical with the thorium of Berzelius by Damour (C. R. 34, 685), Berlin (P. 87, 608), and Bergemann himself (P. 85, 558). In 1862 Bahr supposed he had found a new metal, which he called wasium, but soon afterwards he recognised that the metal was identical with thorium (P. 119, 572; 132, 227). The metal thorium was isolated by Berzelius in 1828 (P. 16, 385); it was also obtained by Chydenius in 1861 (Bl. [2] 1, 130); and in a state of greater purity by Nilson in 1882 (B. 15, 2537).

Occurrence.—Compounds of Th have been found only in very small quantities in a few Thorite from Lövön in Norway conlocalities. tains c. 59 p.c. ThO2 according to Berzelius; an orange-coloured variety of this mineral, orangite, was found by Bergemann to contain 74 p.c. ThO₂; Nilson (B. 15, 2519) found from 50 to 59 p.c. ThO2 in different specimens of thorite. Small quantities of ThO₂ have been found in specimens of pyrochlore, monazite, samarskite, euxenite, gadolinite, and orthite. Several minerals containing Th have been discovered by Hidden a. Mackintosh in North Carolina and Colorado; one of these is a silicophosphate of Th, another a silicate of Th and Y, another a silicate of Th and U, and another is a uranate of Th, Pb, and Y (Am. S. [3] 36, 461; 38, 474).

Formation. - 1. By reducing ThCl. or 2ThCl, KCl, by heating with K (Berzelius, P. 16, 885). — 2. By reducing fused ThCl., or 2ThCl. KCl, by means of Na (Chydenius, Bl. [2]

1, 130; 6, 433; Nilson, B. 15, 2537).

Preparation.-Powdered thorite is evaporated with HClAq to dryness; the residue is heated to render silica insoluble, and treated with water; the solution is filtered and ppd. by H₂S; the filtrate from PbS is ppd. by NH₂Aq, and the pp. of crude ThO₂xH₂O is thoroughly washed and dissolved in HClAq; Th(C₂O₄)₂ is ppd. from this solution by H₂C₂O₄Aq, and the pp., after being thoroughly washed with boiling water, is decomposed by heating; the crude ThO, thus obtained is converted into Th(SO₄)₂ by treatment in a Pt basin with conc. H₂SO₄ after being thoroughly moistened with water. The impure Th(SO₄), thus obtained is freed from acid and water by heating, and is dissolved in as little ice-cold water as possible (c. 5 parts are required), and the solution is heated to c. 20°, when Th(SO₄), 9H₂O ppts. The Th(SO₄), 9H₂O is dehydrated by heating; the Th(SO₄), is dissolved in the minimum quantity of ice-cold water and the solution is heated to c. 20°; the crystals of Th(SO₄), 9H₂O that separate are de-hydrated, dissolved in ice-cold water, and the hydrated sulphate is ppd. by heating to 20°. This treatment is repeated a few times, when pure Th(SO₄),9H₂O is obtained. This method of purifying crude sulphate of Th, which is given by Nilson (B. 15, 2521), depends on the great differences between the solubilities in water of

Th(SO₄), and Th(SO₄), 9H₂O; the former, when pure, is soluble in 20.6 parts water at 0c, the latter requires 88 parts water at 0°, and much more at 20°, for solution. The sulphates of the metals present in the crude Th(SO,), prepared as described are much more soluble in water at 20° than Th(SO₄), 9H₂O. For other methods of preparing salts of Th from thorite v. Berzelius (l.c.), Chydenius (l.c.), Delafontaine (Ar. Sc. [2]

18, 848), and Cleve (Bl. [2] 21, 115).

To prepare Th from pure Th(SO₄), Nilson (B. 15, 2538) dissolved the salt in water, ppd. by KOHAq and boiled, washed the ThO₂xH₂O thoroughly, dissolved in HOlAq and reppd. by KOHAq (to remove all traces of H.SO.); he thoroughly washed the pp. by decentation, dissolved it in HClAq, added KCl, in the ratio 2ThCl.:KCl, evaporated to dryness in a Pt basin, and dried the residue by warming over a flame, stirring constantly, and then powdering and heating again. The dry double salt of ThOl, and KCl was heated to redness in a stream of dry HCl (prepared from conc. H,SO, and sublimed NH,Cl), HCl was removed by a stream of H, and some of the salt was transferred to a cylinder of wrought iron (fitted with a lid which screwed on), in which a layer of dry NaCl had been placed; pieces of sodium were arranged over the Th-K salt in the cylinder, on these was placed another layer of the Th-K salt, and the cylinder was thus nearly filled with alternate layers of Na and the double salt; the contents of the cylinder were pressed together by a piston which fitted into the cylinder, sufficient dry NaCl was added to fill the cylinder, the lid was screwed on, and the apparatus was heated to full redness for about fifteen minutes; after cooling, the contents of the apparatus were treated with cold water, whereby chlorides were dissolved and the excess of sodium was converted into NaOH and dissolved (the action is apt to be explosive); the thorium was washed with water, then with alcohol, and finally with ether, and dried at 100°.

The yield of thorium by this method is almost equal to that calculated from the equation. For a detailed description of the iron cylinder used for reduction v. Nilson a. Pettersson, W. 4, 554

(on the preparation of Be).

Properties.—Prepared as described, Th forms a greyish-white, glistening powder; the powder consists of small, thin, six-sided, crystalline leaflets; the larger crystals are as white and lustrous as silver, the smaller are grey and less lustrous. The crystals are somewhat brittle. S.G. at 17° 11.1 to 11.23; the latter number referring to the well-crystallised metal (Nilson, B. 15, 2543; 16, 160). S.H. (0° to 100°) ·02787, hence at. heat = 6.4 (N., B. 16, 161). Crystallises in combinations of regular octahedra and hexahedra; isomorphous with Si (N., I.c.; and note, p. 162). S.V.S. 20.9. The emission-spectrum of Th has been mapped by Thalen (Détermin. des Longueurs d'Onde des Raies Métalliques [Upsala, 1868]); the most marked lines are three in the indigo; 4393, 4382, and 4281. Soret (Ar. Sc. [2] 68; 59; [3] 4, 94) found that ThCl, Aq cuts off almost the whole of the ultra-violet part of the spectrum.

Th is unchanged in air up to 100°-120°: when heated to a higher temperature it burns to ThO, with production of much white light. Th 712 THORIUM.

is readily oxidised by heating in O; on this account some of the metal is oxidised in the process of preparation if the lid of the cylinder does not fit tightly. Nilson's experiments (B. 15, 2541) show that Th does not melt at the full heat of a wind-furnace. Th does not react with water; it combines directly with Cl, Br, I and S. The metal is slowly acted on by H₂SO₄ or HNO, rapidly by HClAq or aqua regia; alkali

solutions do not seem to react.

The at. w. of Th has been determined: (1) by analysing Th(SO.), and hydrates of this salt (Berzelius, P. 16, 398 [1829]; Chydenius, P. 119, 43 [1861]; Delafontaine, Ar. Sc. 18, 343 [1863]; Hermann, J. pr. 93, 114 [1864]; Cleve, Bl. [2] 21, 116 [1874]; Nilson, B. 15, 2527 [1882]; Krüss a. Nilson, B. 20, 1665 [1887]); (2) by analyses of the double Th-K sulphate (Berzelius, l.c.); (3) by analyses of the acetate, oxalate, and formate of Th (Ohydenius, l.c.); (4) by determining S.H. of Th (Nilson, B. 16, 153 [1883]); (5) by finding Y.D. of ThCl. (Krüss a. Nilson, B. 20, 1665 The mol. w. of Th is not known.

Th is distinctly metallic in its chemical relations; the only oxide that has been isolated with certainty, ThO₂, is basic, forming salts ThX₂ (X = SO₄, 2NO₅, &c.). There are indications of the existence of a higher oxide (? Th₂O₇). Thorium forms the last member of the even-series family of Group IV. in the periodic classification of the elements. The members of this family are C, Ti, Zr, Ce, —, Th. The four elements Ti, Zr, Ce, and Th closely resemble one another in their chemical properties; they are also similar to the odd-series family of Group IV., namely, Si, Ge, Sn, —, Pb. For a general account of Group IV. (and more especially of C and Si) v. CARBON GROUP OF ELEMENTS (vol. i. p. 682); for a description of the even-series family (Ti, Zr, Ce, Th) v. TITANIUM GROUP OF ELEMENTS (this vol. p. 749); and for the odd-series family (Ge, Sn, Pb; Si being described in vol. i. p. 652) v. Tin GROUP OF ELEMENTS (this vol. p. 735).

The atom of thorium is tetravalent in the

gaseous molecule ThCl4.

Reactions and Combinations (Nilson, B. 15, 2541).-1. Heated in air, oxidation begins at c. 120°, and at much below red heat the metal burns brilliantly to ThO2.—2. Heated in a stream of oxygen, the metal burns to ThO2, with the production of dazzling white light.-3. Th burns to ThX, when heated in chlorine, in bromine vapour, or in vapour of iodine.—
4. When heated with sulphur part of the S volatilises, and when the temperature is above the b.p. of S combination occurs with production of much light, ThS₂ being formed.—5. According to Troost (C. R. 116, 1227), Th combines with carbon, when ThO, is mixed with C and heated in CO, in the electric arc (v. Carbide, infra).— 6. Water does not react with Th at any temperature, so far as experiments have been tried. Nilson (l.c.) gives no details, but says 'Thorium is not changed by water under any conditions.'-7. Sulphuric acid reacts slowly with Th; when dilute, the acid evolves H; when cone. and hot, SO2 is given off.—8. The action of nitric acid is slight; the dilute warm acid acts very gradually; conc. acid is almost without action.—9. Hydrochloric acid rapidly dissolves Th, forming ThOL, and giving off H; the action is most complete with conc. acid.—10. Aqua regia dissolves Th easily .- 11. Alkaline solutions seem not to react with Th.

Detection and Estimation.—Salts of Th are colourless when the acid is colourless. Alkalis, and also (NH₄)₂SAq, ppt. gelatinous, white ThO2.2H2O, insoluble in excess of the precipitant; alkali carbonates form a white pp. soluble in excess; solution in (NH₄)₂CO₂Aq becomes turbid at 60°, but goes clear on cooling; saturation of a solution of a salt of Th with K2SO4 forms white Th(SO₄)₂.K₂SO₄.2H₂O, insoluble in K₂SO₄Aq, but soluble in water; K₄FeCy₆Aq ppts. white ThFeCy,; oxalic acid and alkali oxalates ppt. white amorphous Th(C2O4)2, insoluble in dilute acids, soluble in solutions of alkali oxa-

lates or NH₄.C₂H₄O₂, especially on warming.

The is generally estimated by ppn. as
ThO_{2.2}H₂O or Th(C₂O₄)₂, and subsequent conver-

sion into ThO, by heating.

Thorium, bromide of. The gummy, white mass obtained by Berzelius (P. 16, 385), by evaporating a solution of ThO₂.2H₂O in HBrAq, was probably ThBr. The hydrated salt ThBr. 10aq seems to have been obtained, from ThO₂. 2aq, by Jannasch, Locke, a. Lesinsky (Zeit. f. anorg. Chemie, 5, 283).

Thorium, carbide of. By heating an intimate mixture of ThO₂ and C in the electric arc, in an atmosphere of CO_2 , Troost (C. R. 116, 1227) obtained a hard, brittle solid, containing from 8.2 to 9.5 p.c. C, and therefore approximating to the formula ThC₂ (which requires 9.38 p.c. C). S.G. 10.15 at 15°; scratches glass slightly; decomposes cold water, giving off H and strongly smelling hydrocarbons; gradually decomposes in moist air; burns rapidly and brilliantly when heated to redness.

Thorium, chloride of, ThCl4. Mol. w. 373.5. V.D. 172.2 at 1050°-1270° (Krüss a. Nilson, B. 20, 1671). This compound is formed by heating a mixture of ThO₂ and C in a stream of dry Cl; the ThCl₄ sublimes in white needles. Krüss a. Nilson (B. 20, 1675) prepared pure ThCl₄ by heating Th to incipient redness in a stream of dry, air-free HCl; then raising the temperature to full redness, in order to sublime a little FeCl₂, formed from the iron in the thorium used; and finally subliming the ThCl, at an incipient white heat. (For details of the apparatus used v. Nilson a. Pettersson, J. pr. [2] 33, 1 [on preparation of BeCl. 1).

ThCl, crystallises in lustrous, white needles; moderately hygroscopic, but remains some hours in the air before deliquescing (K. a. N., *l.c.*). Does not volatilise at c. 420° (Chydenius, *P.* 119, 43). The V.D. found by K. a. N. at temperatures from 1050° to 1270° shows that the formula ThCl, is molecular. K. a. N. got the value 142 for V.D. at 1400°, showing that dissociation had begun. Troost (C. R. 101, 360) obtained values for V.D. of ThCl, at c. 950° considerably lower than those of K. a. N. It is likely that his specimen was not pure, or that the apparatus used contained traces of O sufficient to cause decomposition to ThO2 and Cl (K. a. N., l.c., p.

Hydrated thorium chloride ThCl4.8H2O was obtained by Cleve in thin, white prisms, by concentrating a solution of ThO_{2.2}H₂O in HClAq, allowing to crystallise, and drying over H.SO.

(Bl. [2] 21, 116).

Double salts. - ThCl, combines with potaseium chloride to form 2ThCl4, KCl.18H2O; obtained by mixing very cone. solutions of the constituents. Forms small, white crystals; v. sol. water, also sol. alcohol; dehydrated by heating in HCl (Cleve, Bl. [2] 21, 116). By heating a dry mixture of ThOl, and NH,Cl in HCl, Chydenius (Bl. [2] 1, 130; 6, 433) obtained a compound with ammonium chloride, which, when crystallised from water in vacuo, had the composition ThCl.8NH,Cl.8H,O; loses 6H,O at 100°. Cleve (l.c.) prepared a compound with platinic chloride ThCl, PtCl, 12H₂O; and Nilson (l.c.) a compound with platinous chloride 2ThCl, 3PtCl, 24H2O.
Thorium, ferrocyanide of, ThFeCy, 4H2O.

A white powder; prepared by adding K.FeCy, Aq to solution of a salt of Th (Cleve, Bl. [2] 21,

116).

Thorium, fluoride of, ThF4.4H2O. Obtained by adding HFAq to ThCl,Aq; the gelatinous pp. becomes powdery after a time (Chydenius, Bl. [2] 1, 130; 6, 433). Loses H_2O at 100°, and 2H₂O at 140°-200°; heated to redness gives ThO, and HF. Berzelius (P. 16, 385) obtained a heavy white powder by heating ThO_{2.2}H₂O with HFAq. Combines with potassium fluoride; Chydenius (l.c.) obtained two salts, (1) ThF.KF by mixing ThCl.Aq with KHF.Aq, (2) ThF.,2KF.4H.O by boiling ThO.,2H.O with conc. KHF.Aq and HFAq. The isolation of a third salt, 7ThF.,8KF.6H.O, is doubtful.

Thorium, hydride of. By heating ThO., with

excess of Mg powder, in a stream of H, Winkler (B. 24, 885) obtained a grey powder, which gave off H with dilute HClAq, and burnt when heated in O. Analyses of this powder gave 72.86 p.c. Th, 50 p.c. H, 17.57 p.c. Mg, and 9.07 p.c. O. Supposing the ThO₂, Mg, and H interacted in accordance with the equation $ThO_2 + 2Mg + 2H$ =ThH₂+2MgO, then, from the quantities of ThO, and Mg used, the product would give on analysis 73 77 p.c. Th, 63 p.c. H, 15 45 p.c. Mg, and 10 15 p.c. O. From these data Winkler concluded that a hydride, ThH2, was formed in the reaction.

Thorium, hydroxides of, v. Oxides and HY-

DRATED OXIDES (infra).

Thorium, iodide of. The white gummy mass, crystallising after a time, obtained by Chydenius (Bl. [2] 1, 130; 6, 433) by evaporating a solution of $ThO_2.2H_2O$ in HIAq, was probably ThI_4 . The substance goes brown in the light.

Thorium, nitride of. By heating ThO2 or ThCl, in NH, also by heating ThCl, and NH,Cl in HCl, Chydenius (l.c.) obtained a small quantity of what he supposed to be either a nitride of Th or a compound of the metal with N and

Thorium, oxides and hydrated oxides of. Besides the oxide ThO, a peroxide, probably

Th2O7, is said to exist.

THORIUM DIOXIDE ThO2. (Thoria.) Mol. w. not known. Obtained by ppg. a salt of Th by alkali, (NH4)2SAq, or KCNAq, drying, and strongly heating; also by decomposing Th(C2O4), by heat. A fine, white powder. S.G. 10-22 at 17° (Nilson, B. 15, 2536). S.V.S. 25-87. S.H. (0° to 100°) 0548 (N. a. Pettersson, B. 13, 1459). By heating ThO₂ with borax in a porcelain oven, Nordenskijold obtained the oxide in microscopic crystals isomorphous with SnO, and ZrO, (P. 150, 219). Tho, has not been fused; it is not reduced by heating with C, but when mixed with C and heated in Cl the chloride ThCl, is obtained. ThO2 does not react with molten alkalis. It is scarcely acted on by scids; warm conc. H₂SO₄, however, converts it into Th(SO₄)₂.

HYDRATES OF THORIUM DIOXIDE. The dihy $drate \text{ ThO}_2.2\text{H}_2\text{O} (= \text{ThO}_4\text{H}_4, thorium hydroxide})$ is obtained by ppg. a solution of a salt of Th by an alkali, (NH₄)₂SAq, or KCNAq, washing, and drying at 100°. It forms a hard, white solid. ThO_{2.2}H₂O combines with CO₂ in the air; it dissolves in acids, forming salts ThX₂, where X = SO₄, 2NO₄, &c. Another hydrate, 4ThO₂.H₂O₄ (= Th₄O₇(OH)₂, thorium metahydroxide) is obtained, according to Cleve (Bl. [2] 21, 116), by heating ThO2 (obtained by decomposition of Th(C₂O₄)₂) with excess of HNO₂Aq or HClAq at 100° till the acid is driven off, adding water, and then ppg. the opalescent solution so obtained by NH, Aq, washing, and drying at 100°. This hydrate is said to be insoluble in dilute acids.

THORIUM PEROXIDE Th2O,. An oxide with this composition is said to be produced by adding H₂O₂Aq and NH₃Aq to a solution of a salt of Th (Cleve, Bl. [2] 43, 53; Lecoq de Boisbau-

dran, C. R. 100, 605)

Thorium, oxysulphide of. By strongly heating ThO2, obtained from ThO2.2H2O, in CS2 vapour, Chydenius (Bl. [2] 1, 130; 6, 488) obtained a solid substance to which he gave the formula 2ThO. ThS. = Th.O.S. According to Krüss a. Volk (Zeit. f. anorg. Chemie, 5, 75; 6, 49), the compound obtained by heating ThO, in CS, is ThOS, and the same compound is formed by heating 2ThCl₄.KCl in H₂S.

Thorium, phosphide of. By heating Th in vapour of P, Berzelius (P. 16, 385) obtained a dark grey, metal-like, lustrous solid, which was changed to Th phosphate by heating in air.

Thorium, platinocyanide of,

Th(PtCy₁), 16H₂O. Yellow-green, orthorhombic prisms (Cleve, Bl. [2] 21, 116).

Thorium, salts of. All the normal compounds which have been isolated by replacing H of acids by Th belong to the class ThX2, where $X = 2NO_3$, $2IO_3$, SO_3 , SO_4 , $\frac{2}{3}PO_4$, &c.; basic salts also exist. The chief salts of oxyacids are borate, carbonate, chlorate, chromate, iodate and periodate, molybdate, nitrate, oxalate, phosphates, selenate and selenites, silicates, sulphate and sulphite, and tartrate.

Thorium, silicofluoride of. By treating ThO₂2H₂O with H₂SiF₆Aq, Cleve (Bl. [2] 21, 116) obtained a semi-transparent, orystalline solid, probably Th(SiF₄)₂; insol. in excess of H₂SiF₄; over H₂SO, gave off HF and SiF₄.

Thorium, sulphide of, ThS₂. Obtained by heating Th with S; excess of S sublimes, and

when the temperature passes the b.p. of S the elements combine with evolution of heat and light (Nilson, B. 15, 2542; cf. Berzelius, P. 16, Also formed by heating ThO2, obtained 385). from Th(C₂O₄), to white heat in vapour of CS₂ (Chydenius, BL. [2] 1, 130; 6, 433); but, according to Krüss a. Volk (L.c.), the compound thus formed is ThOS. A black powder. Burns when heated in air, giving off SO, and leaving 714 THORIUM.

ThO₂. Not acted on by HClAq or HNO₂Aq; aqua regia dissolves the compound, forming Th(SO), Molten KOH forms K,S and ThO;

heating in Cl produces ThCl.

of. The salt Thorium, sulphocyanide ? Th(SCy), is obtained, as a viscid mass, by eva-porating a solution of ThO₂.2H₂O in HSCNAq. A solution of the compound gives a pp. with HgCy, said to be Th(OH), SCy. HgCy, H,O; and the filtrate is said to give crystals of Th(OH)(SCy), 3HgCy, 12H2O (Cleve, Bl. [2] 21, 116). M. M. P. M.

THUJIGENIN C₁₄H₁₂O, or C₂₂H₂₄O₁₄. Occurs in small quantity in Frondes Thujæ, the green parts of Thujæ occidentalis (Rochleder a. Kawalier, Sits. W. 29, 10). Minute needles, v. sl. sol. water, sol. alcohol. NH₂ colours its alcoholic solution green. AcCl forms resinous C14H11AcO7.

THUJIN C₂₀H₂₂O₁₂. Occurs in the green branches of Thuja occidentalis (Rochleder a. Kawalier, Sits. W. 29, 10). Minute lemon-yellow four-sided tables, sol. alcohol. Tastes bitter. Its alcoholic solution is coloured yellow by alkalis, is turned reddish-brown by air, gives a yellow pp. with lead acetate, and gives a darkgreen colour with FeCl. Reduces ammoniacal AgNO. Boiling dilute H.SO. turns it first green, then yellow, and splits it up into sugar and thujetin. Boiling baryta-water hydrolyses it in like manner.

Thujetin $C_{2n}H_{2n}O_{1n}$. Yellow solid, nearly insol. water, sol. alcohol and ether. Not altered by dilute HCl or H,SO,. Its alcoholic solution is turned a splendid bluish-green colour by NH, Aq. Gives, in alcoholic solution, a red pp. with lead acetate, and a black colour with FeCl. Boiling baryta-water converts it into thujetic acid C₂₈H₂₂O₁₈, which crystallises in minute lemon-yellow needles, sol. alcohol and ppd. by water.

(B)-THUJOL is TANACETONE.

THUJONE $C_{10}H_{18}O$. (212°). S.G. 20 .9265. Occurs, together with the terpene C1.H1s and levorotatory fenchone, in the essential oil of thuja obtained by distilling the ends of the branches and the leaves of *Thuja occidentalis* with water (Schweizer, A. 51, 398; Jahns, Ar. Ph. 221, 748; Wallach, A. 272, 109). Yields an oxim.

Reactions .- 1. When heated with ammonium formate it yields a formyl derivative which on saponification gives rise to 'thujoneamine' C₁₀H_{1,1},NH₂ (199°), which forms the salts B',H,PtCl, and B'HCl. Thujone-amine hydrochloride decomposes on distillation into NH,Cl and thujene $\hat{C}_{l_0}H_{l_0}$ (c. 178°), S.G. 22 ·84.—2. KMnO, forms two 'thujaketonic acids.'—3. Br forms $C_{l_0}H_{l_0}Br_{l_0}O$ [122°] (Wallach, A. 275, 179)

(a)-Thujaketonic acid CieHieOs i.e. CH, CO.C, H₁₂.CO, H. [76°]. S. c. 2·5 at 100°. Transparent plates. Yields an oxim [c. 186°].

 (β)-Thujaketonic acid C,H_{1/2}(OO.CH₂).CO₂H.
 [79°]. S. 1·48 at 100°. Small matted needles.
 On distillation it yields methyl heptyl ketone (186°) S.G. 22° 854, which gives ψ-cumene distillation. hydride C,H₁₄ on heating with ZnCl, at 100° (Wallach, A. 275, 166). Forms an oxim [106°] which crystallises in needles.—AgA': v. al. sol.

cold water.

THULIUM. This name was given by Cleve in 1879 to the metal of a new earth which he announced to be present in specimens of gadolinite (C. R. 89, 478). The name thulium was derived from Thule, the old name for Scandinavia. The presence of the new earth was more or less confirmed by the work of Nilson (B. 13, 1433), and of Soret (C. R. 89, 251). In 1880 Cleve said he had isolated thulia, and described it as a white earth, giving an emission spectrum characterised by two bright lines 5896 and 5306, and forming solutions of salts which showed two absorption bands 6840 and 4645. To this earth Cleve assigned the formula Tm₂O₃, and he determined the maximum value of the atomic weight of the supposed element to be Tm = 170.7. In 1888 Krüss a. Nilson declared that thulium consists of two distinct elements (B. 21, 1681).

The absolute homogeneity of thulia has been by no means established (v. METALS, RARE, vol. iii. p. 245). M. M. P. M.

THYME OIL. The essential oil from garden thyme, Thymus vulgaris, contains lævorotatory thymene $C_{10}H_{16}$ (160°-165°), thymol $C_{10}H_{14}O$ and a little cymene (Lallemand, A. 102, 119; A. Ch. [3] 49, 155). The oil from wild thyme, Thymus serpyllum, contains cymene, thymol, carvacrol, and a phenol coloured violet by FeCl, (Jahns, Ar. Ph. [3] 16, 277; Febre, C. R. 92, 1290; Buri, Ar. Ph. [3] 12, 485).

THYMO-ACRYLIC ACID v. Oxy-GYMYL-

ACRYLIC ACID

THYMOHYDROQUINONE v. HYDROTHYMO-

THYMOL C₁₀H₁₄O i.e. C₂H₃MePr.OH [5:2:1]. Mol. w. 150. V.D. 75·3 (calc. 75) (Eykman, B. 22, 2757). [49·5°] (Reissert, B. 23, 2242). (232°). S.G. § '9941. C.E. (0°-10°) '00085. S. 3. S.V. 188.9 (Pinette, A. 243, 46); 189.3 (Ramsay). $R_{\infty} = 76.56$ (Nasini a. Bernheimer, G. 15, 93). H.F.p. (liquid) 69,250; (solid) -3768. Occurs in essential oil of thyme (Doveri, A. 64, 374; Lallemand, A. Ch. [3] 49, 148; A. 102, 119), of horsemint, Monarda punctata (Arppe, A. 58, 42), of *Ptychotis ajowan* (Stenhouse a. Haines, A. 98, 307; H. Müller, B. 2, 130) of Monarda Didyma, of Ammi Copticum (Buri, Ar. Ph. [3] 12, 485), and in the oil from wild thyme (Jahns, B. 15, 819).

Formation. - By boiling diazo-n-cymene with

water (Widman, B. 15, 170; 19, 245).

Preparation.- By extracting oil of thyme with NaOHAq, ppg. by HCl, and recrystallising from HOAc.

Properties.—Plates (from alcohol), v. sol. alcohol, ether, and HOAc, v. sl. sol. water. Has a slight odour of thyme and a peppery taste. When solid it is slightly heavier than water, when liquid it is lighter. Not ppd. by adding water to its alcoholic solution. Not dissolved by NH,Aq, but absorbs NH, becoming liquid, but re-solidifying after the NH, has escaped. Sol. KOHAq. FeCl, does not colour its aqueous solution. Bromine-water only gives a milkiness. An aqueous solution (1 vol.) warmed first with HOAc (\$ vol.) and then with H.SO. (1 vol.) is coloured reddish-violet, the solution showing two absorption bands at E and D (Wolff, Fr. 22, 96). KOHAq followed by I in KI gives on warming a fugitive red tint (Itallie, Fr. 29, 205).

Estimation: Messinger, B. 23, 2754.

Reactions.-1. Decomposed by distilling with P.O. into propylene and m-cresol.-2. P.S. yields cymene (Fittica, B. 6, 938; A. 172, 305). 8. Oxidised by chromic acid to thymoquinone.-4. PCl. yields chloro-cymene. — 5. Potash-fusion yields several oxy-acids (Barth, B. 11, 567, 1571).—6. Iodine and potash yield a red amorphous pp. [110°] converted by heat or by boiling water into a yellow body [165°] (Messinger a. Vortmann, B. 22, 2316).—7. Boiling POCl₂ (1 mol.) acting on thymol (1 mol.) forms liquid C₁₀H₁₃O.POCl₂ (280°–285°) which is converted by water into liquid C₁₀H₁₃O.PO(OH)₂. which yields BaA" 4sq and is decomposed by heat into thymol and HPO, (Discalzo, G. 15, 279).—8. Boiling POCl_s acting on thymol (2 mois.) forms $(C_{10}H_{12}O)_2$ POCl (330°-340° at 320 mm.), a liquid which is converted by water into (C₁₀H₁₃O), PO.OH, which is insol. water and yields NaA' [74°] (Kreysler, B. 18, 1705) and BaA', 5aq. S. 197 at 21°.—9. PCl₃ (1 mol.) acting on thymol (4 mols.) forms (C₁₀H₁₃O), PO [59°] crystallising from alcohol in prisms (E. a. L.).-10. SiCl₄ forms Si(OC₁₀H₁₅)₄ [48°] (450°) crystallising from CHCl, in prisms (Hertkorn, B. 18, 1693).—11. Cl.CO.NH, reacts forming C₆H₃MePr.O.CO.NH.CO.NH₂ [190°] (Gattermann, A. 244, 44).—12. C₆H₄(NMe₂)(NH₃) [1:4] forms indothymol C₆H₄(NMe₂).N COH₂MePr

[69.5°] crystallising in needles, insol. water and alkalis, forming a blue solution in alcohol, a violet in ether, and a green in HOAc. HClAq decomposes indothymol into thymoquinone, dimethyl-aniline, and NH, (Bayrac, Bl. [3] 7, 97).

13. Chloral forms Cl. H₁, OC, HCl₂O [130°–134°) (Mazzara, G. 13, 272).—14. Chloral, conc. H2SO4, and HOAc yield CCl3.CH(C10H12.OH)2 (Jäger, B. 7, 1197).—15. Dilute H₂SO₄ gradually added to a mixture of benzoic aldehyde (1 mol.) and thymol (2 mols.) forms C.H.CH(C10H12.OH)2 [146°] which is sol. alkalis and yields a diacetyl derivative [126°] (Russanoff, B. 22, 1943).—16. COCl, passed into an aqueous solution of C10H12ONa forms di-thymyl carbonate (C1eH1aO)aCO which crystallises from alcohol in needles [48°] (above 360°) and yields o-oxybenzoic acid when heated with NaOPh at 180° (Richter, J. pr. [2] 27, 505). According to Bender (B. 19, 2268), di-thymyl carbonate, prepared by heating thymyl ethyl carbonate at 300°, melts at 60°. COCl₂ also forms, as a byproduct, C₁₆H₁₂O.COCl, which is converted by ammonia into C₁₆H₁₂O.CO.NH₂ [181°].—17. Cl.OO₂Et acting in the cold upon C₁₆H₁₂O.NA. forms C10H18O.OO.OEt (c. 261°), an oil which yields o-oxy-benzoic acid, PhOEt, and thymol when heated with NaOPh (Richter).—18. Treatment with ClOH, CO, H and NaOHAq produces C₁₀H₁₈O.CH₂.CO₂H [148°] which yields BaA'₂ 2aq, PbA', AgA', oily EtA' (290°), and an amide [97°] (Saarbach, J. pr. [2] 21, 159; Spica, G. 10, 342). 19. a-Chloropropionic acid and KOHAq yield C₁₀H₁₀O.CHMe.CO.H [48°] (Scichilone, G. 12, 50).—20. Thymol (10 g.) mixed with H₂SO₄ (70 g.) containing nitrous acid gives thymolchroin C, H, N,O, a dark-violet amorphous body which dissolves in alcohol, ether, CHCl, and benzene, forming red solutions with pale-green fluorescence. Alkalis turn its alcoholic solutions blue. Sublimes at 140°, giving off violet vapours.

Yields brown amorphous C₄₀H₄₀Ao₄N₂O₄ (Brunner a. Chuit, B. 21, 252; cf. Liebermann, B. 7, 1100).-21. By heating with ammonium-zine chloride it is converted into thymylamine C₁₀H₁₂NH₂ and di-thymylamine (C₁₀H₁₂)₂NH.-22. Cyanuric chloride converts sodium thymol into $(C_{10}H_{13})_2C_3N_3O_3$ [151] a yellow crystalline powder, insol. water, sl. sol. alcohol (Otto, B. 20, 2239).—23. Aqueous KOH and K28,0, form C10H12O.SO2.OK, which crystallises from alcohol in slender silky threads (Heymann a. Königs, B. 19, 3307). It is oxidised by alkaline KMnO. B. 19, 3307). It is oxidised by to oxycuminic acid C₆H₂Pr(OH).CO₂H [4:3:1].

Salts. - C10H18ONa. Crystalline. aqueous solution is ppd. by HgCl₂ and AgNO₃. C₁₀H₁₃O.Hg.HgNO₃.—C₁₀H₁₃O.Hg.HgOAc (Merck, G. P. 48539; Pharm. Zeit. 1889, 625).—(C₁₀H₁₃O)₂Hg₂O (Lallemand).—(C₁₀H₁₃O)₂Al. Decomposed by heat into propylene and (C.H.Me.O) Al, the m-cresylate then splitting up into alumina, m-cresol, di-m-tolyl oxide, and a body C₁₈H₁₄O [200°] S. (alcohol) 17 at 20°; 1 at 78°; S. (benzene) 93 at 21°; S.G. 805; V.D. 208.8, crystallising in pearly plates (Gladstone a. Tribe, C. J. 39, 9; 41, 11).

Acetyl derivative C10H13OAc. (245°). S.G. º 1.009.

Benzoyl derivative C₁₀H₁₈OBz. H.F. 90,480 (Stohmann, J. pr. [2] 36, 9).

Methyl ether C₁₀H₁₃OMe. (216° cor.). S.G. § 9531. C.E. (0°-10°) ·00083. S.V. 214·3 (Pinette, A. 248, 47). H.F.p. 61,429 [C,O, = 94,000; H₂,O = 69,000] (Stohmann, J. pr. [2]

25,260, Formed from thymol, KOH, and MeI (Engelhardt a. Latschinoff, Z. [2] 5, 43).

Ethyl ether C_{1e}H₁₈OEt. (227°) (Pinette; cf. Jungfleisch, Bl. [2] 4, 17). S.G. § 9334. C.E. (0°-10°) 00089. S.V. 240. H.F.p. 68,858 (Stohmann). Yields the ethyl derivative of oxysterophthelic acid on oxidation (Petern et al., 2008). terephthalic acid on oxidation (Paterno a. Canzoneri, G. 9, 460). Decomposes at 3600-400° into thymol and ethylene (Bamberger, B. 19.

Propyl ether C10H13OPr. (243°). S.G. §

9276. C.E. (0°-10°) 90088. S.V. 265·5.

Butyl ether C₁₀H₁₀OC₄H₂. (258°). S.G. §
9230. O.E. (0°-10°) 90085. S.V. 289·2.

Isoamyl ether C₁₀H₁₂OC₅H₁₁. (238°-243°)

(E. a. L.). Oil.

Heptylether C₁₀H₁₃OC,H₁₃. (307°). S.G. § -9097. C.E. (0°-10°) ·00082. S.V. 368·7.

Octylether C₁₀H₁₃OC₈H₁₇. (320°). S.G. § -9026. C.E. (0°-10°) ·00075. S.V. 395·6 (Pinette). Ethylene ether $(C_{10}H_{12}O)_2C_2H_4$. Plates (from ether) (Paterno, G. 5, 13).

Nitroso-thymol v. Oxim of Thymoguinone.

Dithymol v. DI-OXY-DICYMYL.

References .- Amido-, Bromo-Amido-, Bromo-NITRO-, TRI-CHLORO-, IODO-, IODO-AMIDO-, and NITRO-THYMOL.

DI-THYMOL-ETHANE v. DI-OXY-DI-CYMYL-ETHANE

THYMOL DICARBOXYLIC ALDEHYDE C₁₂H₁₄O₂ i.e. C₂HMePr(OH)(CHO)₂. [80°]. Formed as a by-product in the preparation of thymotic acid by the action of chloroform and NaOH on thymol (Kobek, B. 16, 2104). Needles. Gives a red colour with FeCl,

THYMOL (a)-SULPHONIC ACID
C.H.MePr(OH).SO.H. [92°] (Stebbins, Am. 3,
111). Formed from thymol by the action of

 $\rm H_2SO_4$ or $\rm Cl.SO_2.OH$ at 50° (Engelhardt a. Latschinoff, Z. 1869, 44; 1871, 261). Pearly plates (containing aq), v. e. sol. water. FeCl. colours neutral solutions dark-violet. Yields thymoquinone on oxidation.

Salts.—KA'2laq. Efflorescent tables, v. e. sol. water.—Nah'2laq. [167°].—NH,A'2aq. [172°].—CaA'2aq. [157°].—BaA'24aq. Prisms, decomposing at 100°.—PbA'24aq. Bensoyl derivative C₁₂H₁₂(OBz).SO₃H.

-KA' 2aq. -CaA', 4aq. -BaA', 5aq. -PbA', 5aq. Ethyl derivative C₁₀H₁₂(OEt)SO₃H. Formed from the K salt, KOH, and EtI.—KA'. Thin plates.—BaA'2 3aq. Plates, sl. sol. hot Aq.

Isoamyl derivative C_{1e}H₁₂(OC₅H₁₁).SO₈H.
-KA'. Needles, v. sol. water. - BaA'₂3aq.

Thymol (β)-sulphonic acid C₁₀H₁₂(ÕH).SO₃H. Formed in small quantity by the action of H2SO. on thymol.—KA'aq. Plates, sl. sol. cold water. Coloured violet-blue by FeCl,

Thymol (γ)-sulphonic acid $C_{10}H_{12}(OH).SO_3H$. Formed by heating thymol with fuming H2SO. at 100° (E. a. L.).—KA'aq. Granular, v. sol. water.—BaA'23aq. Tufts of needles, more soluble than the (a)-isomeride. Not decomposed at

Benzoyl derivative C₁₀H₁₂(OBz).SO₂H.-KA' 3aq. Flat needles, sl. sol. cold water

Ethyl derivative C₁₀H₁₂(OEt).SO₃H.— KA'. Flat needles.—BaA'₂ Saq. Six-sided tables.

Thymol-sulphonic acid

Ethyl derivative C₆H₂MePr(OEt)SO₃H [1:4:3:6]. Formed by boiling diazo-cymenesulphonic acid with absolute alcohol.—BaA'2 3aq: white plates, sol. hot water, sl. sol. cold (Widman, B. 19, 247)

Thymol sulphonic acid. Methyl derivative C, H, (OMe). SO, H. Formed, together with an isomeride, by sulphonation of the methyl ether of thymol (E. a. L.; Paterno a. Pisati, B. 8, 440).—KA: large tables.—BaA', 3aq. Nodules, v. sol. hot water. S. 3.94 at 26°

Thymol disulphonic acid $C_{10}H_{11}(OH)(SO_3H)_2$. Formed by sulphonating thymol.—K₂A" 1 aq. Efflorescent needles (from alcohol), v. e. sol.

References. - AMIDO-, BROMO-, and IODO-

THYMOL SULPHONIC ACID.

THYMOQUINONE C10H12O2 i.e. C6H2MePrO2 Mol. w. 164. [45.5°]. (232°). [2:5:4:1]. Formed by oxidation of thymol (Lallemand, J. 1854, 592), of the methyl ether of thymol (Paterno, B. 8, 440), of cymenol (Carstanjen, J. pr. [2] 8, 58; 15, 410), of di-oxy-di-cymylethane (Steiner, B. 11, 289), of carvacrol psulphonic acid (Claus, J. pr. [2] 39, 356), and of amido-thymol (Andersen, J. pr. [2] 23, 172; Armstrong, B. 10, 297; Liebermann a. Ilinski, B. 18, 3194). Yellow crystals, with pungent odour, v. sl. sol. water, v. sol. alcohol and ether. Reduced by SO, to hydrothymoquinone at 60° forms crystalline C₁₀H₁₈O₅SK. Phenyl cyanate forms $C_{10}H_{12}O:NO.CO.NHPh$ [182°], crystallising in large yellow needles (Goldschmidt a. Strauss, B. 22, 3106).

Oxim C.H.MePrO(NOH) or C₆H₂MePr(OH)(NO). Nitroso - thymol. [162°]. Formed by the action of nitrous acid on thymol (R. Schiff, B. 8, 1500; Widmann, B. 15, 170; Liebermann, B. 18, 8194). Formed also by the action of hydroxylamine hydrochloride on

thymoquinone (Goldschmidt a. Schmid, B. 17, Slender monoclinic needles, a:b:c =1.987:1:1.8941; β = 94° 57' (Panebianco, G. 10, 78); sl. sol. hot water. Its alkaline solution is red. Oxidised by alkaline K₃FeCy₆ to nitrothymol. Reduced by tin and HCl to amidothymol (Liebermann, B. 10, 77). Does not react with SO₂ (Schmidt, J. pr. [2] 44, 521). Fuming HCl forms di-chloro-thymoguinone and chloroamido-thymol (Sutkowski, B. 19, 2315). KOH and BzCl form C₁₀H₁₂O(NOBz) [110°]. Hydroxylamine (3 mols.) in strongly alkaline solution forms, in 48 hours, a white substance which, if ppd. by HOAc, immediately dissolved in dilute NaOH, treated with an equal weight of a phenol, and exposed to the air, gives a deep-blue colour or pp. turned red by acids (Kehrmann a. Messinger, B. 23, 2818, 3557).

Di-oxim C₁₀H₁₁N₂O₂ i.e. C₁₀H₁₂(NOH)₂. Formed by boiling an alcoholic solution of nitroso-thymol (1 mol.) with hydroxylamine hydrochloride (2 mols.) and partially neutralising with alkali from time to time. Yellowishwhite granules, decomposing at 235° without melting. Insol. water and NH, Aq, sol. KOHAq.

Chlorimide C₈H₂MePr $\stackrel{\vee}{<}_{NCl}$. Formed by adding a solution of bleaching powder to a cold acidified solution of the hydrochloride of p-amido-thymol (Andresen, J. pr. [2] 23, 169). Pungent oil, volatilising even at 15°. Decomposes at 160°-170°. Volatile with steam. Decomposed by conc. HClAq into chloro-amidothymol and mono- and di- chloro-thymoquinone. Conc. HBrAq behaves in a precisely similar way. Alcohol at 140° converts the chlorimide into thymoquinone. Conc. aqueous SO₂ reduces it in a few days to hydro-thymoguinone. Reduced by tin and HCl to p-amido-thymol, considerable quantities of hydrothymoquinone being formed at the same time.

Polymeride [201°]. Formed by the action of daylight on thymoquinone (Armstrong, B. 10, 297; Liebermann, B. 10, 2177; 18, 3193). Silky yellow needles (from alcohol), insol. ether. Changes to thymoquinone on distillation. Not attacked by SO₂ at 180°, but reduced by HI and P, or in alcoholic solution, by Zn and HClAq to hydrothymoquinone.

Phenyl hydrazide [249°]. Insol. benzene. Oxim $(C_{10}H_{13}NO_2)_2$. [263°]. Crystalline, insol. water. Reduced by tin and HCl to amidohydrothymoquinone

 $Dioxim (C_{10}H_{14}N_2O_2)_x$. [c. 290°]. Powder. May be reduced to di-amido-cymene.

References .- DI - AMIDO-, BROMO-, CHLORO-, Iodo-, and Oxy- THYMOQUINONE.

o-THYMOTIC ACID C₁₁H₁₄O₃ i.e. C₆H₂Me(C₃H₇)(OH).CO₂H[6:3:2:1]. Mol. w. 194. [123°]. Formed by the action of CO, upon heated sodium-thymol (Kolbe a. Lautemann, A. 115, 205; Kobek, B. 16, 2101). Silky crystals, v. sol. alcohol, ether, and benzene, almost insol. cold water. Volatile with steam. Gives a deep-blue colour with FeCl₂. Yields thymol when distilled with baryta. PCl₃ acting on the Na salt forms thymotide C₁₁H₁₂O₂ [187°], which

is reconverted into thymotic acid by potash fusion (Naquet, Bl. 4, 92).

p-Thymotic acid C₁₁H₁₄O₂ i.e. C.H.Me(C.H.)(OH).CO.H[6:8:4:1]. [157°] TIN. 717

Formed by heating thymol with aqueous NaOH and CCl, (Kobek). White plates, v. sol. alcohol, ether, and benzene, insol. cold water. coloured by FeCl,

Methyl derivative

 $C_6H_2Me(C_3H_7)(OMe).CO_2H.$ [137°]. Silky needles (from dilute alcohol).-AgA': pp.

p-THYMOTIC ALCOHOL C,1H18O2 i.e. $C_0H_2Me(C_3H_7)(OH).CH_2OH.$ [120°-130°]. Formed by reducing thymotic aldehyde with sodium-amalgam (Kobek, B. 16, 2098). Amorphous

powder, v. sol. alcohol and ether, insol. water.
p-THYMOTIC ALDEHYDE C₁₁H₁₄O₂ i.e. ſ133°]. $C_6H_2Me(C_5H_7)(OH)CHO[6:3:4:1].$ Formed, together with CaHMe(CaH,)(OH)(CHO), [6:3:4:5:1] [80°], by heating thymol with chloroform and NaOHAq (Kobek, B. 16, 2096). Silky needles, v. sol. alcohol, sl. sol. hot water. On heating with aniline it yields the compound C₆H₂Me(C₃H₇)(OH).CH:NPh [142°] orystallising in yellow needles.

Methyl derivative

C₆H₂Me(C₈H₂)(OMe).CHO. (278°). Formed by methylation of the aldehyde. Oil, yielding an anilide C₆H₂Me(C₈H₇)(OMe).CH:NPh [80°] crystallising in transparent tables.

THYMYLAMINE C₁₀H₁₈NH₂. (230°). Formed, together with di-thymylamine, by heating thymol with ammoniacal ZnBr, or ZnCl, and NH,Br or NH,Cl at 350°-360°, the yield being 25 p.c. (Lloyd, B. 20, 1260). Colourless oil.— B'2H,PtCl: yellow needles.

Acetylderivative C₁₀H₁₈NAc. [112°].

Isomeride v. Carvacrylamine.

Di-thymyl-amine $(C_{10}H_{19})_2NH$. (340°-345°). Formed as above, the yield being 25 p.c. Oil. Its solution in P₂SO₄ is coloured blue by nitrites or nitrates.—B'₂H₂PtCl₆.

Acetyl derivative (C₁₀H₁₃)₂NAc. [78°].

THYMYL CYANURIC ACID v. CYANIC ACID.

THYMYL MERCAPTAN C, H13SH. Thiothymol. (231°). S.G. 989. Formed by heating thymol with P₂S₅ (Fittica, A. 172, 325; Bechler, J. pr. [2] 8, 167). Liquid with pungent odour. Oxidised by HNO₄ to sulpho-toluic acid. $-Hg(SO_{10}H_{18})_2$. Greenish rhombohedra [78°]. -PbA'2: golden needles (from alcohol).

THYMYL PHOSPHATE (C,oH,s),PO. [59° Formed by heating thymol with POCl, the yield being 75 p.c. of the theoretical amount (Kreysler,

B. 18, 1705). Colourless needles.

Formed by heating thymol with THYMYL SILICATE (C10H13)4SiO4. (c. 450°). SiCl., the yield being 70 p.c. of the theoretical amount (Hertkorn, B. 18, 1692).

TIGLIC ACID C.H.O. i.e. CH. CH: CMe.CO.H. Methyl-crotonic acid. Mol. w. 100. [65°]. (199° i.V.) (Kopp, A.195, 84). Occurs as a glyceryl ether in croton oil (Geuther a. Fröhlich, Z. 1870, 549; Schmidt a. Berendes, A. 191, 94; B. 10, 835; Ar. Ph. [8] 13, 213), and as isoamyl ether in Roman oil of chamomile (Köbig, A. 195, 101).

Formation .- 1. By the action of heat or of conc. H2SO, on angelic acid (Demarcay, B. 9, 1933) .- 2. By reducing the dibromide of angelic acid with sodium-amalgam (Schmidt, A. 208, 253).—3. From CEtMe(OH).CO₂Et (derived from oxalic ether, EtI, MeI, and Zn) by treatment with PCl₂ and saponification of the product

(Frankland a. Duppa, A. 136, 9).-4. By distilling CH₃.CH(OH).CHMe.CO₂H (Rohrbeck, A. 188, 235), or by heating it with HIAq (Rücker, A. 201, 61).-5. By reducing methyl-acetoacetic ether with sodium-amalgam and heating the resulting CH₃.CH(OH).CHMe.CO₂H at 200° (Wislicenus, A. 250, 243).-6. By heating veratrine with alcoholic potash (Wright a. Luff, C. J. 33, 347).—7. By heating veratrine with conc. HClAq (Ahrens, B. 23, 2704)

Properties.—Triclinic plates, v. sol. hot water, alcohol, and ether. Smells like benzoic acid. Volatile with steam. Not attacked by sodium-amalgam. Forms with isovaleric acid the double salts CaA'(C,H,O,) 4 aq and

AgA'C,H,O2.

Reactions .- 1. Yields acetic aldehyde and acid on oxidation with KMnO, (Beilstein a. Wiegand, B. 17, 2261; Kondakoff, J. R. 20, 523).—2. Potash-fusion gives propionic and acetic acids.—3. Fuming HI forms an iodovaleric acid [87°] (Schmidt, B. 12, 252).—4. HI and P yield CEtMeH.CO.H.—5. Bromine forms C.H.Br.O. [88°] (Pückert, A. 250, 240; Wislicenus, A. 272, 21; 274, 99).—6. HOCl forms a mixture of CH, CH(OH). CClMe. CO, H [112°] and CH3.CHCl.CMe(OH).CO2H [75°] (Melikoff, Bl. [2] 47, 166).

Salts.-KA'.-CaA', 3aq. S. (of CaA', 6.4 at 17°.—BaA'2 4aq. S. (of BaA'2) 18.5 at 16°.—

AgA'

Ethyl ether Eth'. (155°). S.G. $^{\circ}$.942. Isoamyl ether $C_{s}H_{11}A'$. (205°). Reference.—Chloro-tiglic acid.

TIGLIC ALDEHYDE C.H.O. Guaiol. (118°). V.D. 2.92. Formed by distillation of gum guaiacum (Deville, C. R. 17, 1143; 19, 134; Völckel, A. 89, 346; Herzig, M. 3, 118). Formed also by heating acetic aldehyde (1 mol.) with propionic aldehyde (1 mol.) and aqueous (28 p.c.) NaOAc for 30 hours at 100° (Lieben a. Zeisel, M. 7, 53). Pungent oil, miscible with alcohol and ether. Oxidised by air to tiglic acid. Alkaline sodium nitroprusside gives a violet-red colour destroyed by HOAc (Von Bitto, A. 267, 376). Br forms di bromo-valeric aldehyde. Iron and dilute (50 p.c.) acetic acid reduce it to valeric aldehyde (91°), an inactive amyl alcohol CHMeEt.CH_OH (129°) and tiglic alcohol $C_0H_{10}O$ (c. 130°). Reacts with phenyl-hydrazine, and combines with NaHSO. CrO, forms acetic acid (Gilm, A. 106, 379). Aqueous SO₂ at 0° forms oxy-pentane disulphonic acid (Haymann, M. 9, 1055).

TIGLYL ALCOHOL v. PENTENYL ALCOHOL.

TIN Sn. At. w. 118'8 (v. infra). Mol. w. not known with certainty (v. infra). Melts at 231'68° (Callendar a. Griffiths, C. N. 63, 1); for other data, giving m.p. from 220° to 235°, v. Carnelley's Melting and Boiling-point Tables (2, 12). Boils between the m.p.s of Ni and Fe, i.e. between 1450° and 1600° (Carnelley a. Williams, C. J. 35, 566). S.G. c. 7.3 (determinations vary from c. 7.0 to c. 7.5) (v. Clarke's Table of Specific Gravities [2nd ed.] 5). The so-called allotropic variety of tin seems to have S.G. from 5.8 to 6.1 (v. Schertel, J. pr. [2] 19, 322). S.H. (0°-100°) 0559; so-called allotropic tin has S.H. 0545 (0°-100°) (Bunsen, P. 141, 1); S.H. liquid tin (250°-350°) = 0687 (Person, A. Ch. [3] 24, 129). O.E. (linear, 0°-100°) .00002296 (Matthiessen,

Pr. 15, 220; v. also Kopp, A. 81, 1; Fizeau, C. R. 68, 1125). T.C. 15.2 (Ag = 100) (Wiedemann, P. M. [4] 19, 243). E.C. (Hg at 0° - 1) at 15° = 8.823 (Kirchoff a. Hausemann, W. 13, 406); at $0^{\circ} = 8.726$, at $100^{\circ} = 6.091$ (Lorenz, W. 13, 422). Heat of fusion (for 1 kilo. tin) = 14,252 (Person, A. Ch. [3] 24, 129). For spectrum of tin v. Thalén (A. Ch. [4] 18, 237); Salet (C. J. [2] 9, 1147); also Hartley a. Adeney (T. 1884 [i.] 118).

Historical.—Tin was known long before the Christian era. Pliny distinguished plumbum album or candidum from plumbum nigrum; in the fourth century A.D. the former of these substances began to be called stannum, from which

word the symbol Sn is derived.

Occurrence.—Tin has been found in Siberia and in South Australia. Tinstone or cassiterite, consisting essentially of SnO₂, is found in considerable quantities, both in veins and in alluvial deposits, in many parts of the world. Tin pyrites, containing SnS, with Cu2S, FeS, and ZnS, is found in small quantities in various tin veins. Small quantities of SnO₂ occur in various *lithia-micas* (Sandberger, J. M. 1878. 291, 657). According to Lockyer (Pr. 27, 279), there are indications of the occurrence of tin in the sun.

Formation.—1. By ppn. from solution of SnCl, in HClAq by tin; or by tin and iron simultaneously, with a little Fe₂O₃ to neutralise excess of acid (Schultze, B. 28,974).—2. By electrolysis of SnCl, in dilute HClAq.—3. By reduction of

SnO, by heating with C.

Preparation.—Tinstone is cleansed by agitation with water; it is then sorted, stamped or rolled to a certain fineness, and washed. The washed ore is calcined in reverberatory calciners to remove sulphur from the sulphides of Fe that are present in most tin ores; the calcined ore is exposed to the air for some days to insure the oxidation of any sulphides that remain in the ore to sulphates, and it is then washed in water, whereby sulphates of Fe, Cu, &c., dissolve and the tin oxide sinks to the bottom of the vessels. The tin oxide is then separated into lighter and heavier parts by agitation with water; the heavier parts are mixed with charcoal or anthracite, some lime or fluorspar is added as a flux, and the mixture is smelted in a reverberatory furnace. The product of the smelting is purified by melting, when the tin runs off, leaving the less fusible foreign metals behind; this purified tin is again melted, the molten mass is stirred with billets of green wood for some hours, and is then allowed to partially solidify by cooling, when the metal separates into different strata, which are ladled into moulds: the upper stratum is the purest and the undermost is the least pure. Sometimes the smelting is conducted in a blast furnace. For details v. Tin in Dictionary of Applied CHEMISTRY, vol. iii. p. 834.

Commercial tin generally contains small quantities of As, Ou, Fe, and Pb, and occasionally traces of Sb, Bi, Mn, Mo, W, and Zn. To prepare pure tin, the purest commercial metal is granulated and dissolved in conc. HClAq; a little water is very slowly poured into the liquid so as to form two layers, and a plate of tin is placed in the liquid so that part of it is in the upper (less cone.) and part in the under (more cone.) layer. A slight electric current is thus produced, and crystals of pure tin are deposited on the plate (Hiller, A. 85, 258).

Pure tin was prepared by Bongartz a. Classen (B. 21, 2903) by heating the purest commercial tin (Banca tin) in dry Cl and so forming SnCl. distilling fractionally, and collecting that which passed over at 120°, adding to the SnCl, about four times its volume of water, and then crystallised Na S until the pp. of SnS that formed was dissolved, then adding NaOHAq about equal to half the quantity of Na,S used, allowing to stand for some days, drawing off the clear, almost perfectly colourless liquid into a Ptbasin, and passing an electric current through this alkaline liquid, washing the ppd. tin with water, then with absolute alcohol, and drying at 100°. The current was arranged so that it produced c. 2 to 3 c.c. detonating gas per minute from water; c. 2 to 3 g. tin were obtained in 24 hours.

Properties .- Pure tin is a silver-white metal; the commercial metal generally has a slightly yellowish tinge. It is unchanged in pure air; but it tarnishes in the air of large towns from formation of a film of sulphide. Tin is one of the least tenacious metals; a wire 1.6 mm. diameter breaks with a weight of c. 14 kilos. Tin may be hammered into plates c. 0254 mm. thick; the malleability varies much with temperature, at 200° the metal is so brittle that it crumbles when hammered. Tin readily crystallises; by fusing, allowing to cool partially, and pouring out the still fluid metal, fair-sized crystals are obtained. Crystals are formed by depositing tin by electrolysis. Stolba (J. 1873. 282) covers the outside of a Pt basin, all except a small portion, with wax, places the unwaxed surface of the basin on a plate of amalgamated zinc in a larger porcelain basin, fills the Pt basin with a dilute and not very acid solution of SnCl, and the porcelain basin with water containing c. $\frac{1}{20}$ part of HClAq; crystals of tin deposit on the bottom of the Pt basin. The forms of the crystals of tin are probably rhombic and quadratic (von Foullon, J. M. 1885 (11) Ref. 266). According to Frankenheim (P. 40, 456), the crystals belong to the regular system; Miller (B. J. 24, 133) obtained quadratic prisms by separating tin by electrolysis. If a piece of tin is brushed over with warm, dilute aqua regia, or with a mixture of warm dilute H₂SO₄Aq and HNO₂Aq, the surface becomes covered with a fretwork of crystals, from the facets of which light is unequally reflected, and the surface appears like watered silk. When a bar of tin is bent a crackling sound may be heard due to the crystals in the inner parts of the bar breaking against one another. When warmed, tin has a characteristic smell.

By exposure for some time to a very low temperature (c. -39°) tin crumbles to a grey powder (v. Fritzsche, B. 2, 112, 540); S.G. c. 5.8 to 6.1 (v. Schertel, J. pr. [2] 19, 322). Tin which had been kept for 300 to 400 years was found by Schertel (l.c.) to have become reddishgrey, and so brittle that it was crushed by pressure between the finger-nails; by immersion in boiling water the metal became more coherent and lighter in colour, and the S.G. increased to 7.8 (cf. Oudemanns, P. M. [5] 4, 470).

Tin remains lustrous in pure air at the ordinary temperature, and also in water; a

greyish-white film of SnO₂ forms on the surface of tin kept molten in the air; when heated to whiteness in air the metal burns brightly to SnO₂. Tin dissolves in HOlAq, forming SnCl₂ solution; dilute H₂SO₄Aq has only a slight action, conc. hot H₂SO₄ produces SnSO₄ and SO₅; dilute HNO₃Aq dissolves tin, with formation of Sn(NO₂)₂Aq and NH₄NO₂Aq, conc. HNO₂ transforms the metal into SnO₂. Tin dissolves in hot conc. NaOHAq, forming Na₂SnO₂ solution and giving off H.

When haloid compounds of tin are volatilised in a stream of H and the gas is ignited, the flame shows two cones; the inner cone is blue with SnCl₂, green with SnBr₂, and yellow with SnL₃, and gives a continuous spectrum; the outer cone is carmine-red, and gives a spectrum with two characteristic lines (Salet, O. J. [2] 9,

1147).

Tin is metallic physically, and in most of its chemical relations. The oxide SnO₂ interacts towards several acids as a basic oxide, forming salts SnX₂ (X = SO₄, &c.); but with conc. KOHAq or NaOHAq this oxide reacts as an acidic oxide, forming salts M₂SnO₃. Tin is the third member of the odd-series family of Group IV. in the periodic classification of the elements; it is closely allied to Ge and Pb, and less closely to Si; it shows very marked analogies with Ti, Zr, Ce, and Th, which (with C) form the even-series family of Group IV. For details of the chemical relations of tin v. Tin group of elements, this vol. p. 735.

The atom of tin is divalent in the gaseous molecule SnCl₂, and tetravalent in the gaseous

molecule SnCl.

Supposed allotropic form of tin. The grey brittle powder formed by the action of great cold on tin, and also found in some very old specimens of the metal, is sometimes regarded as an allotropic variety of tin (v. supra). the specimens of tin, however, which have been observed to undergo this change have contained small quantities of impurities; until the change has been effected with pure tin, and has been proved to occur without any change of mass, the existence of an allotropic variety of tin cannot be said to be established. According to von Foullon (J. M. 1885 [1] Ref. 266) there are three varieties of tin: (1) rhombic, S.G. 6.52 to 6.56; (2) quadratic, S.G. 7-196; (3) grey brittle tin, B.G. 5.781 to 5.809.

Atomic weight of tin. The at. w. has been determined (1) by oxidising Sn to SnO₂ by HNO₃ (Gay-Lussec, A. Ch. [2] 80, 163 [1812]; Berzelius, P. 8, 184 [1812]; Mulder, A. 72, 212 [1849]; Vlaanderen, B. J. 1858. 183 [1858]; Dumas, A. Ch. [3] 55, 134 [1859]; van der Plaats, C. R. 100, 52 [1885]; Bongartz a. Classen, B. 21, 2900 [1888]); (2) by ppg. Cl from SnCl₂ by AgNO₂Aq (Dumas, A. Ch. [3] 55, 156 [1859]); (3) by electrolysing SnCl₂.2NH₂Cl and SnCl₂.2KCl (B. a. C., l.c.); (5) by reducing SnO₂ in H. (v. d. P., l.c. [1885]); (6) by determining S.H. of Sn (Bunsen, P. 141, 1 [1870]); (7) by determining V.D. of, and analysing, SnCl₂ and SnCl₄. The older determinations gave values from 115.9 to 117.8. The results of the determinations of B. a. C. are summarised by them (l.c., p. 2909) as follows:—

Number of experi- ments	Method	At. weight	Diff. be- tween min. and max. found		
11	oxidation of Sn to SnO.	118.7606	•459		
16	electrolysis of SnCl2NH,Cl	118-8093	•228		
10	electrolysis of SnCl2KCl	118-7975	•163		
10	electrolysis of SnBr.	118-7809	-144		

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B. a. C. regard the second and third methods as the most trustworthy; in the oxidation of Sn to SnO₂ the Pt vessel was also slightly acted on, and there was the possibility of a very small loss in dissolving SnBr₄ in the last method. The mean of the results obtained by the electrolysis of the two double chlorides is 118.8 (O = 15.96).

Molecular weight of tin. As the V.D. of tin has not been determined the molecular weight is not known. Ramsay determined the depression of the vapour pressure of Hg caused by dissolving tin therein; assuming that equal volumes of dilute solutions contain equal numbers of molecules of the dissolved bodies, and that the mol. w. of liquid Hg is 200, the results obtained by B. gave the mol. w. of tin as 1174 in one experiment and 149.5° in another.

Reactions and Combinations.-1. When tin is heated in air or oxygen, a film is formed on the surface, containing both SnO and SnO; when heated to whiteness tin burns brightly, forming SnO₂ (v. Emich, M. 14, 845).—2. Heated in *chlorine*, bromine, or iodine, the compounds SnX2 and SnX, are formed, according to whether there is excess of tin or halogen. - 3. Tin combines with sulphur, and with selenion, to form SnS and SnSe, when heated with these elements.—4. Tin probably combines with phosphorus when heated in vapour of that element .- 5. Tin forms alloys with many metals (v. Tin, alloys of, p. 720).— 6. Water is decomposed by tin at a full red heat with formation of SnO and H.-7. Sulphydric acid has a slight action on tin, in moist air, forming a film of SnS .- 8. Hydrochloric acid dissolves the metal, forming solution of SnCl, and giving off H; the reaction proceeds more rapidly with warm conc. HClAq.—9. Pure conc. nitric acid, S.G. c. 1.55, has no action on tin. When the acid is diluted the products of the reaction vary with the concentration and the temperature; according to Walker (C. J. 63, 845), both Sn(NO₂), and Sn(NO₂), are formed, the quantity of Sn(NO₂), increasing as concentration increases at a fixed temperature; with very dilute scid an increase of temperature causes only a slight increase in the quantity of the stannic salt, but the effect of temperature is very marked with more conc. acid; the proportion of stannous to stannic salt formed is only slightly affected by changing the relative masses of tin and acid; when a fairly conc. solution of tin in HNO, Aq is heated, hydrated stannic nitrates, varying in composition, are deposited (v. also Montemartini, G. 22, 884, 897, 426; abstract in C. J. 62, 1402; v. also Hay, C. N. 23, 298; Scott, C. N. 28, 322). Pure

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HNO, diluted sufficiently to start the reaction produces chiefly SnO₂.xH₂O (v. Hydrated stannic oxide, under Tin oxides and Hydrated oxides, p. 725). When tin dissolves in HNO,Aq, NH,NO, is formed; the gases evolved are NO, N.O. and N (v. Ackworth a. Armstrong, C. J. 32, 84).—10. A mixture of nitric and sulphuric acids, or nitric and hydrochloric acids, produces NH₂OH along with NH₃ (Divers, C. J. 43, 443; D. a. Shimidzu, C. J. 47, 597; cf. Hydroxyl-amine, vol. ii. p. 734). With certain proportions of HNO, Aq and H2SO, Aq, Bassett (C. N. 53, 172) obtained Sn(NO₃)₂ in solution, and almost pure N₂O.—11. Dilute sulphuric acid reacts slowly when warmed with tin, forming SnSO, if the tin is kept in excess, and chiefly Sn(SO₄)₂ if the acid is in excess, and giving off Conc. sulphuric acid reacts to form SnSO., or $Sn(SO_4)_2$, according to the proportion of tin to acid, and evolves H_2S or SO_2 , or both, according to the concentration of the acid and the temperature; with pure hot H_2SO_4 , SO_2 is given off, and S separates (v. Calvert a. Johnson, C. J. [2] 4, 435; also Pattison Muir a. Robbs, C. N. 45, 69).—12. Aqua regia dissolves tin, forming SnCl,-13. Tin reacts with mercuric ohlorids and bromide, when heated with these compounds, forming SnCl₂, or SnBr₂, and Hg.-14. With sulphurous chloride, SnCl₂ is formed, with production of much heat (Wöhler, A. 73, 374).-15. Tin dissolves in warm cone. caustic potash, or soda, solution, forming M2SnO3 and giving off H.—16. Fusion with nitre forms K,SnO.

Detection and Estimation.—H., S ppts. dark brown SnS from slightly acidified solutions of stannous salts; the pp. is soluble in warm yellow ammonium sulphide, forming (NH₄)₂SnS₂Aq, from which acids ppt. yellow SnS₂. AuCl₄Aq produces a purple pp., a compound of Sn, Au, and O (v. Purple of Cassius, under Tin OXIDES AND HYDRATED OXIDES, p. 727). H₂S ppts. yellow SnS₂ from acidified solutions of stannic salts; the ppt. dissolves in warm yellow ammonium sulphide, and acids re-ppt. SnS₂ from this solution. Tin is generally estimated as SnO₂.

Tin, alloys of. Tin forms alloys with many metals. For a general account of the properties of different classes of tin alloys, v. Dictionary of Applied Chemistry, vol. iii. p. 837. For alloys of tin with antimony, v. Karmarsch (D. P. J. 123, 267); Kestner (Kast. Archiv, 19, 424); Johnson (Chem. Gasette, 1855. 180); Chaudet (A. Ch. [2] 3, 376); with bismuth and lead, and with bismuth, cadmium and lead, v. vol. i. p. 511; with cadmium, v. vol. i. p. 655; with cadmium and gold, v. Heycock a. Neville (C. J. 59, 936); with copper, v. vol. ii. p. 254 (also Dictionary of Applied Chemistry, vol. iii. p. 838); with copper and lead, v. vol. iii. p. 125 (also French, S. C. I. 8, 36); with iridium, platinum, and rhodium, v. Deville p. Debray (C. R. 81, 839); with iron, v. vol. iii. p. 53 (also Headden, Am. S. [3] 44, 464); with gold, v. Laurie (P. M. [5] 33,94); with lead, v. vol. iii. p. 125; with lead and sinc, v. vol. iii. p. 125 (also Wright a. Thompson, Pr. 45, 461; 48, 25); with mercury, v. infra; with platinum, v. vol. iv. p. 288; with silver, v. W. a. T. (Pr. 48, 25); with sodium, v. Bailey (C. N. 65, 18). Tin alloys with sinc in all proportions; the alloys are harder than tin, but softer than sinc; they are less

malleable than tin. Many varieties of bronse are alloys of tin with sinc and copper (v. Rudberg, P. 18, 240). Amalgams of tin are readily formed by immersing tinin Hg; the action is more rapid if the Hg is warmed (v. Böttger, J. pr. 1, 305; Joule, Chem. Gasette, 1850. 339, also C. J. 16, 384; Kupffer, A. Ch. [2] 40, 293).

16, 384; Kupffer, A. Ch. [2] 40, 293).

Tin, bromides of. Tin combines with Br to form two compounds, SnBr, and SnBr. The V.D. of the latter has been determined, and the formula SnBr, is molecular; from the analogy of the chlorides it is probable that the formula SnBr, is molecular also. Watts a. Bell (C. J. 33, 442) obtained both SnBr, and SnBr, by heating SnO, in Br vapour mixed with CO.

STANNOUS BROMIDE SnBr₂. (Dibromide of tin.) Obtained by Balard (A. Ch. [2] 32, 337); more fully examined by Rayman a. Preis (4. 223, 323). Prepared by heating tin in HBr gas, allowing the oily liquid so obtained to cool, dissolving the crystals that form in dilute HBrAq, and purifying the solid that separates by distilling it in a stream of N. A crystalline, slightly yellow, transparent solid; melts at 215.5° to a transparent liquid, which solidifies at 215°. S.G. 5.117 at 17°. Carnelley a. Williams (C. J. 35, 563) say that SnBr₂ boils between 617° and 634°. Partly decomposed by water, with separation of an oxybromide of tin (R. a. P., I.c.). By crystallising a solution of tin in warm conc. HBrAq, R. a. P. (l.c.) obtained the hydrate $SnBr_2.xH_2O$ (x=1 or 2), in colourless needles. By cooling a mixture of NH₄Cl and solution of tin in warm conc. HBrAq, the compound SnBr₂.2NH₄Cl.H₂O was obtained in large rhombic,

porcelain-like crystals (R. a. P., l.c.).

STANNIC BROMIDE SnBr. (Tetrabromide of tin.) Mol. w. 437.8. Melts at 30° (Carnelley a. O'Shea, C. J. 83, 55), at 33° (Rayman a. Preis, A. 223, 323). Boils at c. 201° (C. a. O'S., l.c.), at 203.3° (R. a. P., l.c.). S.G. 3.349 (R. a. P., l.c.). V.D. at 228°-260° = 227-230 (C. a. O'S., l.c.). Prepared by passing vapour of Br over melted tin in a tube shaped thus

the tin being at b and the Br at a. The tin burns in the Br vapour; the product is distilled backwards and forwards from b to c until the distillate is quite colourless. On cooling the liquid solidifies to colourless crystals of SnBr, (C. a. O'S., l.c.). R. a. P. (l.c.) prepared SnBr, by heating tin with Br, and crystallising from SnCl₄ or SnBr₂Cl. SnBr₄ forms white crystals, which are slowly decomposed in the air; according to R. a. P. (l.c.) the crystals deliquesce to a clear liquid, which when placed over H.SO, deposits crystals of the hydrate SnBr, 4H.O. SnBr, is soluble in cold water; after some hours SnO, xH,O is deposited; boiling hastens this change; addition of HNO,Aq to hot SnBr, Aq ppts. all the tin as SnO, xH,O, with evolution of a little Br (C. a. O'S., l.c.). R. a. P. (l.c.) found that SnBr, absorbs NH; on heating some NH, was given off, and a yellow sublimate of SnBr. 2NH, was obtained. Accord-ing to Nickles (C. R. 52, 869), SnBr. combines with ether to form SnBr. Et.O.

Stannibromhydric acid H.SnBr. xH.O. (Bromostannic acid.) Hydrobromostannic acid.) This compound was obtained by Seubert (B. 20, 194) by adding 74:1 parts HBrAq (50 p.c. HBr)

to 100 parts SnBr, (i.e. in the ratio 2HBr:SnBr,). A yellow liquid was formed which solidified after a time to a mass of yellow, needle-shaped crystals; the small quantity of mother-liquor was removed by melting and allowing to solidify partially. It is doubtful whether the acid crystallises with 7 or 9H₂O. The acid forms amberyellow needles, probably triclinic; it is very deliquescent; fumes in air, giving off HBr. By adding Na₂OO₂ to H₂SnBr₆Aq, and allowing to evaporate, Seubert (l.c.) obtained sodium stannibromide, Na₂SnBr₆GH₂O, in yellow needles. Several other stannibromides are described by Rayman a. Preis (A. 223, 323); their composition is MSnBr₆:xH₂O, where M = (NH₄)₂, Ba, Ca, Fe, Mg, Mn, Ni, Na₂, Sr, and x is 6, 8, or 10.

Tin, bromochlorides of. Rayman a. Preis

111, bromochlorides of. Rayman a. Freis (A. 223, 828) obtained a compound to which they gave the formula SnBr, Cl by heating tin in Br that contained some Cl, and repeatedly distilling. The compound is a liquid boiling between 181° and 190°. It dissolves SnBr, which crystallises out unchanged. By adding Br to excess of SnCl, Ladenburg (A. Supplbd. 8, 60) obtained indications of the formation of bromochlorides,

probably SnBr, Cl and SnBrCl,

Tin, chlorides of. Tin and Cl combine to form two compounds, SnCl₂ and SnCl₄. Both formulæ are molecular.

STANNOUS CHLORIDE SnCl₂. (Dichloride of tin. Tin-salt.) Mol. w. 189·54; perhaps 379·08 (= Sn₂Cl₂) in the liquid state (v. infra). Melts at 249·3° (Carnelley a. Williams, C. J. 35, 63). Boils between 617° and 628° (C. a. W., l.c.); at 606° (Biltz a. Meyer, Z. P. C. 2, 184). V.D. v. infra. H.F. (Sn,Cl²] = 80,790; [Sn,Cl²,Aq] = 81,140; [Sn,Cl²,2H²O] = 86,520 (Th. 3, 327).

Preparation.—1. By heating a mixture of equal parts tin filings and HgCl₂; Hg sublimes and leaves SnCl₂.—2. By heating tin in HGCl gas.—3. By dissolving tin in warm, fairly conc. HClAq, evaporating to the crystallising point, drying the crystals of SnCl₂.2H₂O that separate, heating them in a capacious crucible so long as gases are given off, pouring the liquid into a small crucible, pounding when cold, and distilling from a retort; the first portions of the distillate are said to be pure SnCl₂ (Capitaine, J. Ph. 25, 552).

Properties.—A semi-transparent, white solid; melts at c. 250° and boils at incipient redness, (v. supra). Heated to bright redness, out of air, some SnCl₄ is given off, then SnCl₂. Heated in air gives SnCl₄ and SnO₂. SnCl₂ dissolves in a little water; the solution is decomposed by much dilution, with separation of a hydrated oxychloride SnCl₂SnO.2H₂O. According to Michael a. Kraft (A. Ch. [3] 41, 471), 1,000 c.c. saturated SnCl₂Aq contains, at 15°, 1,333 g. SnCl₂ and 494g. water, and the S.C. is 1.827. Gerlach (D. P. J. 186, 131) gives the following data for a solution of the hydrate SnCl₂.2H₂O:—

P.c.	S.G.	P.o.	8.G.	
8nCi2H.O	of solution	SnCl.2H,O	of solution	
5	1.0331	45	1.3850	
10	1.0684	50	1.4451	
15	1.1050	55	1.5106	
20	1.1442	60	1.5823	
25	1.1855	65	1.6598	
30	1.2300	70	1.7452	
85	1.2779	75	1.8399	
40	1.8298			
Vor IV	7			

An aqueous solution of SnCl₂ generally becomes turbid from ppn. of Sn₂OCl₂; digestion with tin and a little HClAq dissolves the pp.; addition of tartaric acid, or NH₄Cl, stops the solution from becoming turbid, the former by dissolving the oxychloride, the latter by forming a double salt SnCl₂·2NH₄Cl, which is very soluble in water.

Mallet (C. J. 35, 524) obtained a semi-transparent, jelly-like solid from an aqueous solution of SnCl₂ which had stood in a loosely closed bottle for a year or two; the solid dried to a substance like gum arabic; it reddened litmus; the composition of the substance was SnO₂.HCl. Mallet writes the formula as SnO.OH.Cl, and suggests the name chlor-stannic acid.

Molecular weight of stannous chloride. The V.D. of the gas obtained by heating SnCl₂ has been determined repeatedly. The following

table presents the results :-

Calculated for SnCl. 94.7	Sn	ated for .Cl. 9·4	Temp., and Observers
96·5 90·1	103·7 104·5 104·5 123·7 123·7 122·9 119·5 111·4 102·7	?temp. 185·9 189·2	Rieth, B. 3, 668 [1870]. at 619° V.a. C. Meyer, 697° [1879] 800° [1879] 800° [1879] 880° lin, B. 13, 970° lin, B. 13, 639° 675° Biltz a Meyer, Z 760° 760° 790° 1113°

From their determinations, Biltz a. Meyer concluded that molecules of the composition Sn₂Cl₄ do not exist in the gaseous state, and that the mol. w. of the gas comes to a constant value, corresponding with the formula SnCl,, only at a temperature very much above the b.p. of the liquid (SnCl, boils at 606°). In some of the earlier determinations at c. 800° partial decomposition occurred, with production of Cl. The results, taken as a whole, seem to be explained by assuming that the molecules of the liquid, near to the b.p., have the composition Sn2Cl4, that some of these are dissociated to 2SnCl₂ at c. 30° above the b.p., and that this process of dissociation continues slowly, and is not completed at even 500° above the b.p.

Reactions .- 1. Heated strongly in air, SnCl. is given off, and SnO₂ remains. Heated to its b.p. in a retort, SnOl₂ and SnOl₄ distil, and the oxychloride Sn₂OCl₂ remains (Capitaine, J. Ph. 25, 552).—2. When sulphur is added to molten SnCl₂ the products are SnCl₄ and SnS; with selenion the products are SnCl, and SnSe (Schneider, R. 127, 624).—3. Many salts are reduced by SnCl2Aq, either to lower salts or to metals. Au, Hg and Ag are ppd. from solutions of their salts; ferric and manganic salts are reduced to ferrous and manganous salts; CuCl2Aq to Cu2Cl2-4. The higher oxides of Bi, Cr, Pb, Mo, W, &c., are reduced to the lower oxides of these metals.—5. Arsenious oxide is reduced to As by a considerable excess of SnCl₂Aq.—6. Ac. cording to Böttger (Polytech. Notizbl. 35, 96), a mixture of 2 pts. SnCl, rubbed with 1 pt.

potassium chlorate gets hot, and produces an oxychloride of tin, KClO₄, oxide of chlorine, and H₂O. For reactions with water v. supra, Properties; also infra, Combinations, No. 1.

Combinations.—1. With water to form the hydrate SnCl, 2H₂O. This compound is prepared by dissolving tin in warm conc. HClAq and evaporating to the crystallising point, in contact with tin to prevent formation of SnCl. SnCl22H2O, commonly known by the name of tin-salt, forms large white, monoclinic crystals (Marignac, J. 1856, 394); melts at c. 37.7°; S.G. 2.71 at 15.5° solid), 2.588 at 37.7° (liquid) (Penny, C. J. 4, 239). H.F. [Sn,Cl²,2H²O] = 86,520 (Th. 3, 327). When gradually heated to 100°, most of the water is removed; when rapidly heated, H2O and HCl are given off and Sn₂OCl₂ remains, which gives off SnCl₂ when more strongly heated. SnCl₂·2H₂O decomposes slowly on exposure to the air, forming Sn₂OCl₂ and SnCl₄.--2. With ammonia, forming SnCl₂.NH₃.-3. According to Engel (C. R. 106, 1398), a compound of SnCl₂ with hydrogen chloride is formed by passing HCl gas over SnCl2.2H2O at 0°; the liquid thus formed solidifies at 27°, forming crystals of SnCl₂.HCl.3H₂O, which melt at 40°.—4. With chlorides of the alkali and alkaline earth metals, to form $MCl_2.SnCl_2.xH_2O$, where $M = (NH_4)_2$, K_2 , Ba, or Sr(v. Marignac, C. R. 55, 650; Rammelsberg, Krystall. Chemie, 211; Richardson, Am. 14, 89). Poggiale (C. R. 20, 1180) described a salt SnCl₂.4NH₄Cl.3H₂O. Richardson (l.c.) obtained SnCl₂.KCl.H₂O as well as SnCl₂.2KCl.2H₂O. These double salts are perhaps best regarded as stannochlorides, derived from the acid HSnCl_s (v. Combinations, No. 3) and the hypothetical acid H2SnCl4.

STANNIC CHLORIDE SnCl₄. (Tetrachloride of tin.) Mol. w. 260·28. Boils at 113·9° at 760 mm. (Thorpe, C. J. 37, 372); at 114·1° at 760 mm. (Young, C. J. 59, 912). Solidifies at -33° (Besson, C. R. 109, 940). S.G. 6° 2·27875; 1·97813 at b.p. (Thorpe, l.c.). V.D. 133·1 at 124° (Dumas, A. Ch. [2] 33, 385). S.H. ·1402 (10°-15°) (Regnault, A. Ch. [3] 9, 332). Heat of vaporisation = 30,530 (gram-units for 1 kilo.) (Andrews, C. J. 1, 27). H.F. [Sn,Cl·¹] = 127,250; [Sn,Cl·⁴,Aq] = 157,170; [SnCl²,Cl²] = 46,460; [Sn,Cl⁴,Aq,Cl²] = 76,030 (Th. 3, 327). C.E. (cubical, 0°-100°) ·00130244 (Thorpe, l.c.). S.V. 131·07 (Thorpe, l.c.; for S.V. from 0° to 280° v. Young, C. J. 59, 938, 935). For vapour pressures from = 10° to 319° v. Young (l.c., p. 927). Critical temperature = 318·7°; critical pressure = 28080 mm. (Y., l.c., p. 928).

Formation.—1. By heating tin in excess of Cl.—2. By heating 1 pt. tin filings intimately mixed with 4 or 5 pts. HgCl₂.—3. By heating a mixture of Sn(SO₄)₄ and NaCl.—4. By passing vapour of CCl₄ over heated SnO₂ (Watts a. Bell, C. J. 83, 442).—5. By adding fuming sulphuric acid to tin, and then passing in HCl gas and distilling (Heumann a. Köchlin, B. 15, 416).

Preparation.—Dry Cl is led over tin filings heated in a retort connected with a dry flask, which is kept cold; the distillate is distilled from tin filings, and then repeatedly redistilled in a current of dry CO₂. A solution of SnCl₂ is obtained by passing Cl into SnCl₂Aq until no pp. is produced with HgCl₂Aq, or by dis-

solving tin in dilute HClAq containing a little $HNO_{\bullet \bullet}$

Properties.—A thin, mobile, colourless, fuming, very corrosive liquid. Solidifies at -83° to small white crystals (Besson, C. R. 109, 940). SnCl₄ dissolves crystalline S, also P, I, Br, CS₂, &c. (Girardin, C. R. 51, 1057). SnCl₄ does not conduct electricity, even at its b.p.; addition of absolute alcohol forms crystals, probably (SnCl₄.5EtOH), which conduct when dissolved; ether behaves similarly to alcohol (Coldridge, P. M. [5] 29, 383, 480). SnCl₄ withdraws moisture from the air, probably forming SnCl₄.3H₂O (v. infra, Hydrates of stannic chloride). SnCl₄ dissolves in water, with production of heat [SnCl⁴.Aq] = 29,920 (Th. 3, 327), and considerable contraction of volume (v. also Reactions, No. 1). Gerlach (D. P. J. 178, 49) gives the following table:—

Pontge. SnCl.	S.G. SnOl ₄ Aq	Vol. of 100 pts. by wt. of solu- tion; vol. of 100 pts. water=100	Vol. of mixture of SnCl4 and water; vol of sum of consti- tuents=100
0	1.000	100.00	₽ 100·00
10	1.082	92.42	97.82
20	1.174	85.18	95.76
30	1.279	78.19	93.72
40	1.404	71.12	91.42
50	1.556	64.26	88.78
60	1.743	57.37	85.81
70	1.973	50.68	82.63
100	2.234	44.76	100.00

Hydrates of stannic chloride. – Various hydrates have been isolated: (1) SnCl₄, 3H₂O, by exposing SnCl₄ to air (Casselmann, A. 83, 272); also by adding 18 pts. water to 260 pts. SnCl₄, when § of the SnCl₄ remains unchanged, and § forms the hydrate, which sinks in the excess of SnCl₄ (Gerlach, D. P. J. 178, 49); also by dissolving 260 pts. SnCl₄ in 54 pts. water, and letting the solution cool to 60° (G., l.c.); (2) SnCl₄.2H₂O, by keeping the trihydrate in vacuo (Scheurer-Kestner, C. R. 50, 50); also by drying the pentahydrate over H₂SO₄ (Lewy, A. Ch. [3] 16, 303); (3) SnCl₄.5H₂O, by dissolving SnCl₄ completely in water and evaporating (G., l.c.; L., l.c.); (4) SnCl₄.8H₂O, by strongly cooling dilute SnCl₄Aq (G., l.c.); (5) SnCl₄.9H₂O, obtained by Nöllner (Z. [2] 1, 445) by treating SnCl₂Aq with aqua regia, and exposing the solution to a winter temperature. Gerlach (l.c.) gives a table (v. next page), showing the S.G. and percentage composition of solutions of the pentahydrate.

Reactions.—1. With water: SnCl₄ dissolves in water (v. supra) with production of much heat. Vignon (C. R. 108, 1049; 109, 372) found that H₂SnO₅ ppd. from freshly prepared SnCl₄Aq, from SnCl₄Aq after keeping, and from SnCl₄Aq after heating, showed markedly different heats of neutralisation by potash. He concluded that SnCl₄Aq contains HClAq and H₂SnO₂Aq, and that the H₂SnO₃ in solution gradually polymerises. Dilute SnCl₄Aq gradually decomposes, giving HClAq and SnO₂.xH₂O (Casselmann, A. 83, 272). Heating in a sealed tube with a little water is said to give SnO₂.—2. With conc. nitric acid SnCl₄ forms SnO₂.xH₂O.—3. Dry hydrogen sulphide forms white crystals of SnCl₄.5H₃S, which gives off H₂S and HCl when heated, and

Pontge. SnCl ₄ .5H ₂ O	S.G. of solu- tion at 15°	Pontge. SnCl ₄ .5H ₂ O	S.G. of solu- tion at 15°
0	1.000	48	1.347
ĭ	1.006	49	1.357
2	1.012	50	1.3661
3	1.018	51	1.376
4	1.024	52	1.386
5	1.0298	53	1.396
6	1.036	54	1.406
7	1.042	55	1.4154
8	1.048	56	1.426
9	1.053	57	1.437
10 10	1.0593	58	1.447
11	1.066	59	1.458
12	1.072	60	1.4684
13	1.078	61	1.480
14	1.084	62	1.491
15	1.0905	63	1.503
16	1.097	64	1.514
17	1.104	65	1.5255
18	1.110	66	1.538
19	1.117	67	1.550
20	1.1236	68	1.563
21	1.130	69	1.575
22	1.137	70	1.5873
23	1.144	71	1.601
24	1.151	72	1.614
25	1.1581	73	1.627
26	1.165	74	1.641
27	1.173	75	1.6543
28	1.180	76	1.669
29	1.187	77	1.683
30	1.1947	78	1.698
31	1.202	79	1.712
32	1.210	80	1.7271
33	1.218	81	1.743
34	1.226	82	1.759
35	1.2338	83	1.775
36	1.242	84	1.791
37	1.250	85	1.8067
38	1.259	86	1.824
39	1.267	87	1.842
40	1.2755	88	1.859
41	1.284	89	1.876
42	1.293	90	1.8939
43	1.302	91	1.913
44	1.310	92	1.932
45	1.3193	93	1.950
46	1.329	94	1.969
47	1.338	95	1.9881

leaves SnS₂ (Coldridge, P. M. [5] 29, 383, 480).— 4. SnCl₄ dissolves stannous oxide, forming SnCl₂ and a solution of SnO₂·H₂O in excess of SnCl₄ (Scheurer-Kestner, C. R. 50, 50).

Combinations.—1. With hydrogen chloride. By passing dry HCl over SnCl.,5H₂O, then saturating the liquid so formed with dry HCl at 28°, and cooling to 0°, Engel (C. R. 103, 213) obtained white crystalline leaflets of the compound SnCl.,2HCl.6H₂O, melting at c. 20°, and giving off HCl when more strongly heated. The same compound was prepared by Seubert (B. 20, 793) by adding to SnCl., such a quantity of conc. HClAq that the ratio of water in the acid was to the SnCl., as 6H₂O:SnCl., (100 pts. SnCl., require 62·15 pts. of 33 p.c. HClAq), and after a little passing in c. 8 pts. dry HCl; the whole then solidified to a crystalline mass, melting at 19·2°. This

compound is best called stannichlorhydric acid; it has also been called chlorostannic acid (Mallet gave the name chlor-stannic acid to SnO.OH.Cl, v. Stannous chloride, Properties, p. 721), and hydrochlorostannic acid. Chassevant (A. Ch. [6] 30, 5) has described a compound SnCl. HCl. 3H2O. -2. With several metallic chlorides. The double salts of SnCl, and alkali and alkaline earth chlorides have the composition $MSnCl_a.xH_2O$, where $M = (NH_J)_{33}$, K_{23} , Na_{23} , Ba, Ca, Mg, or Sr; these salts are best named stannichlorides (for details, v. Bolley, A. 39, 101; Lewy, J. pr. 37, 479; Rammelsberg, Krystall. chemie; Wittstein, R. P. 64, 7; Topsoš, W. A. B. 69 [2] 261; Morel, C. C. 1891 [1] 492; Chassevant, A. Ch. [6] 30, 5). Cleve (Bl. [2] 31, 195) obtained double salts of the form xSnCl, 2MCl, xH, Q, where M = Ce, Di, La, or Y, x = 2 and 5, and s had large values (from 18 to 45).—8. With certain nonmetallic chlorides; the compound SnCl, 2SCl, is said to be obtained by the interaction of Cl and SnS (H. Rose, P. 44, 320; Casselmann, P. 42, 517); the compound SnCl. PCl, was obtained by Casselmann (l.c.) by heating a mixture of SnCl. 2SCl. and PCl. in a stream of HCl.-4. With certain nonmetallic oxychlorides: (1) SnCl. POCl, formed by the reaction of POCl, and SnCl, (Casselmann, A. 91, 248; 98, 213); (2) SnCl₄ (Casselmann, A. 91, 248; 98, 213); (2) SnCl₄.2SeOCl₂, by combining the constituent compounds (Weber, B. B. 1865. 154); (3) SnCl₄2NOCl, by passing the dry vapour from aqua regia over SnCl₄ (Hampe, A. 126, 43), also by subliming SnCl₄.N₂O₃ formed by the action of N₂O and NO₂ on SnCl₄ (Weber a. Hampe, P. 118, 471).—5. With hydrogen sulphide to form SnCl₄.6H₂S; decomposed by heating to SnS₂, H₂S, and HCl (Coldridge, P. M. [5] 29, 383).—6. With hydrogen cyanide, to form crystals that With hydrogen cyanide, to form crystals that are decomposed in moist air (Klein, A. 74, 85). 7. With ammonia, to form SnCl. 2NH, according to H. Rose (P. 24, 163), to form SnCl.4NH, according to Grouvelle a. Persoz (A. Ch. [2] 44, 322).—8. With phosphorus trihydride, to form SSnCl, 2PH₃ (H. Rose, P. 24, 159); heated to 100° in CO2 there are formed HCl and Sn2P2Cl2, according to Mahn (J. Z. 5, 160).—9. With certain nonmetallic oxides: (1) with SO, to form a solid [? composition] (H. Rose, P. 44, 320); (2) with NO a compound is formed, according to Kuhlmann(A.39,319), but no action occurs according to Hampe (A. 126, 43); (3) with N_2O_3 to form SnCl., N.O., produced by passing NO2 and N2O over SnCl. (Hampe, l.c.).—10. With ethylic alcohol, to form SnCl. 5EtOH (Coldridge, P. M. [5] 29, 383, 480).—11. With amylic alcohol, to form SnCl. 2C, H, (OH) (Bauer a. Klein, Z. [2] 4, 870) 12. With ether, to form SnCl. 2Et,O (Coldridge, l.c.).—13. With various nitriles, forming crystallisable compounds (Lewy, C. R. 21, 371).

Tin, chloropromides of, v. TIN BROMOCHLORIDES, p. 721.

Tin, chloro-iodide of; v. Tin iodochloride, p. 724.

Tin, chlorosulphide of; v. Tin thiochloride, p. 733).

Tin, ferricyanides of; v. vol. ii. p. 340.

Tin, ferrocyanides of; v. vol. ii. p. 337.

Tin, fluorides of. Only one fluoride, SnF₂ has been isolated; double salts of stannic fluoride (SnF₄) are known.

8 A 2

STARROUS FLUORIDE SnF2. (Difluoride of tin.) Formed, in small opaque, lustrous, prisms, by dissolving SnO.xH2O in HFAq, and evaporating; heated in air it forms the oxyfluoride $SnOF_2(=SnF_a.SnO_2)$ (Marignac, Ann. M. [5] 15, 221; Fremy, A. Ch. [3] 47, 37). By dissolving freshly ppd. SnO.xH.O in solutions of alkali fluorides acidified by HFAq, Wagner (B. 19, 896) obtained stannofluorides of the form xSnF₂.2MF.yH₂O, where x = 1 and 3, y = 1 and 2, and M was NH4, K, and Na.

STANNIC FLUORIDE. This compound has not been isolated; a solution of SnO₂.xH₂O in HFAq coagulates when heated, but does not yield any definite compound; when evaporated in air HF is given off, and an oxyfluoride, SnOF2, is depo-

sited (Marignac, l.c.).

Stannifluorides. A series of these salts, MSnF₈xH₂O, where M = one atom of a divalent metal or two atoms of a monovalent metal, has been obtained by Marignac (Ann. M. [5] 15, The salts are isomorphous with the corresponding silicofluorides and titanifluorides; they are generally obtained by saturating the stannates with HFAq and evaporating, some are formed by double decomposition from the Pb or Ag salt. The following salts have been obtained: $MSnF_{a}.xH_{a}O$; $M = NH_{a}$, Ba, Cd x = 6, Ca x = 2, Cu x = 4, Pb x = 3, Li x = 2, Mg x = 6, Mn x = 6, Ni x = 6, K x = 1, Ag x = 4, Na, Sr x = 2, Zn x = 6.

Tin, haloid compounds of. Tin and the halogens combine to form two series of compounds, SnX₂ and SnX₄ (SnF₄ has not been isolated). The V.D.s of SnCl2, SnCl4, and SnBr4 have been determined; it is probable that the simplest formula in every case is molecular. One or two compounds of the forms SnXX' and SnXX3' are known, where X and X' are different halogens. A few oxyhaloid compounds are known, and probably also one or two thiohaloid compounds. The compounds SnBr, and SnCl, combine with HBr and HCl respectively, forming acids H2SnBr, and H2SnCl, from which series of salts, stannibromides and stannichlorides, are derived; stannisluorides are An acid HSnCl, has also been also known. isolated, and a few stannochlorides-MISnCl, and M1,SnCl,-are known, as also some stannoiodides.

Tin, hydrosulphide of. The compound SnS₂H₂, which has probably been isolated, may be called hydrosulphide of tin (v. TIN, THIO-

ACIDS AND SALTS OF, p. 733).

Tin, hydroxides of, v. Tin oxides and HY-DRATED OXIDES, p. 725; also Tin oxyacids and вацтв, р. 727.

Tin, hydroxyl chloride of, SnO.OH.Cl, v.

Chlor-stannic acid, under Stannous Chloride, Properties, p. 721. Tin, iodides of. Two compounds are formed by combining tin and I; they correspond in composition with the two bromides and the two chlorides. The V.D.s of the iodides have not been determined, but the simplest formulæ are probably molecular.

STANNOUS IODIDE SnI2. (Di-iodide of tin.)

Formula probably molecular.

Preparation.-1. Tin and I are heated together in the ratio Sn:2I (1 pt. tin to 2.14 pts. I); the brown crystalline solid so formed (a mixture of SnI, and SnI,) is mixed with tin

filings and heated, when orange-red SnI, sub limes, and SnI, remains as a red crystalline solid mixed with particles of tin, which are easily separated (Henry, T. 1845. 363).-2. Conc. KIAq is added to warm conc. SnCl,Aq; the yellow crystalline pp. is dried and melted, out of contact with air, and a red crystalline mass of SnI₂ is formed on cooling (Boullay, A. Ch. [2] 34, 372).—3. Tin foil is heated for some hours with fairly conc. HIAq in a sealed tube at 120°-150° (Wöhler a. Dünhaupt, A. 86, 374); SnI separates, on cooling, in shining yellow-red prisms.

Properties and Reactions.—A red, crystalline solid. Melts at 316° (Carnelley a. Williams, C. J. 35, 564). Slightly soluble in cold, more soluble in hot, water, without decomposition (Boullay, l.c.). According to Personne (C. R. 54, 216), SnI₂ is decomposed by a large quantity of water, forming HIAq and several oxyiodides SnI₂ is soluble in SnCl₂Aq. heated out of contact with air, SnI, melts without decomposition, but when heated in the air it is decomposed to an oxylodide, which remains

in the vessel, and SnI, which subfimes.

Combinations.—1. With stannous chloride to form SnI₂.SnCl₂ (v. Iodochloride, infra).—2. With stannous oxide to form several oxyiodides (q. v.). — 3. With ammonia, probably forming SnL₂NH₃ (Rammelsberg, P. 48, 169).—4. With alkali iodides and with iodides of the alkaline earth metals to form stanno-iodides; these salts have the composition M2SnI4xH2O and MSnI₃.xH₂O, corresponding with the stannochlorides (v. Stannous Chloride, Combinations, p. 722); $M = NH_4$, K, Na, also $\frac{1}{2}Ba$ and $\frac{1}{2}Sr$. The stanno-iodides are formed by mixing solutions of the constitutent salts, or by adding SnCl,Aq to excess of the alkaline iodide in solution; the salts must be crystallised from alcohol, as they are decomposed by water, forming stannous oxyiodides (v. Boullay, l.c.; Personne, l.c.).

STANNIC IODIDE SnI. (Tetra-iodide of tin.)

Formula probably molecular. This compound is probably formed by heating tin with 4.2 pts. I, and subliming from the product (Henry, T. 1845. 363). Schneider (P. 127, 624) recommends to add 6 pts. CS, to 1 pt. tin filings, and then to add gradually, cooling frequently, 4 pts. I, and to allow the yellow liquid to evaporate.

SnI, orystallises in orange-red octahedra; melts at 146°, sublimes at 180°, and boils at 295° (Personne, C. R. 54, 216). S.G. 4.696 at 11° (Bödeker, Die Beziehungen zwischen Dichten u. Zusammensetzung bei festen u. liquiden Stoffen [Leipzig, 1860]). Solubility in CS2 at 15°=145 (Schneider, *l.c.*); it is also soluble in CHCl₂, EtOH, Et₂O, and C₂H₃. Decomposed by water to SnO₂.xH₂O and HIAq. Combines with ammonia to form SnI_4 where x = 3, 4, and 6 (Personne, l.c.), also = 8 (Rammelsberg, P.48, 169). SnI, is said not to combine with alkali iodides.

Tin, iodochloride of, SnICl (=SnI2.SnCl2). According to Henry (T. 1845, 363) the addition of I to conc. SnCl_Aq causes ppn. of SnI, and on evaporating the mother-liquor (which contains SnCl₂, SnCl₃, and SnL₂) straw-yellow crystals are deposited that have the composition SnI₂SnCl₂. The crystals are decomposed by water, with separation of SnI₂.

Tin, iodosulphide of, v. Tin Thio-lodide, p. 1 733

Tin, oxides and hydrated oxides of. oxides have been isolated, SnO and SnO2; various compounds of these oxides with H₂O seem to exist, but their composition readily undergoes change with variation of temperature. Both oxides interact with acids to form corresponding salts, SnX_2 and SnX_4 , where $X = NO_3$, SO₄, PO₄, &c. Some of the hydrates of SnO₂ also react as weak acids, especially SnO2.H2O (=H₂SnO₃), from which is derived a series of stannates $MSnO_3$, $M = Na_2$, Ca, Pb, &c.; and $xSnO_2xH_2O$ (= xH_2SnO_3), x probably = 5, from which a series of metastannates, xMSnO, is derived. Stannic and metastannic acids and salts derived from them are described under the heading Tin oxyacids and salts and derivatives THEREOF (p. 727). The oxides SnO and SnO, are described in this section of the article Tin, and a brief account is also given of the experiments on the hydrates of these oxides other than stannic and metastannic acids.

When tin is strongly heated in air or oxygen a film forms on the surface consisting of SnO and SnO2; at a full red heat, or incipient white heat, tin burns to SnO₂. According to Vignon (C. R. 107, 734), tin ppd. by zinc from neutral SnCl,Aq or SnCl,Aq oxidises easily in air; after being exposed to the air for some days it contains from 20 to 33 p.c. SnO. Emich (M. 14, 345) found that the surface of tin that was kept molten in the air became covered with crystalline nodules of SnO₂.

STANNOUS OXIDE SnO. (Protoxide of tin.)

Mol. w. not known

Formation.—1. By heating finely-divided tin in the air: the metal becomes covered with a film of SnO (Vignon, C. R. 108, 96). Also by exposing tin ppd. by zinc from SnCl₂Aq or SnCl₄Aq to the air at ordinary temperatures (V., C. R. 107, 734).—2. By dehydrating SnO.xH₂O, ppd. from SnCl,Aq by alkali carbonates, either by heating in a stream of CO₂, or by boiling with water containing a little KOH .- 3. By dissolving SnO.xH₂O in cold KOHAq, and allowing the solution to stand in the air (Ditte, A. Ch. [5] 27, 145).—4. By heating SnC₂O₄ in a tube of hard glass without free access of air (Liebig, A. 95, 116).—5. By ppg. a stannous salt by KCNAq, and boiling the pp. for some days with KCNAq (Varenne, C. R. 89, 360).

Preparation.—SnCl,Aq is ppd. by Na,CO,Aq, the white pp. of SnO.H.O is thoroughly washed with cold water, and is then boiled with water containing a little KOHAq or NaOHAq (less than sufficient to dissolve the pp.), when the SnO.xH₂O is gradually dehydrated, and small black, lustrous crystals of SnO are obtained (Fremy, A. Ch. [3] 12, 460).—2. Tin is dissolved in warm HClAq, the solution is evaporated, in contact with tin, until it solidifies to SnCl2 on cooling; 7 parts, or rather more, Na₂CO₂.10H₂O are then added for each part of SnCl2 in the basin, and the liquid thus formed is heated with constant stirring until it becomes black, and is then allowed to cool; the brownish-black powder, SnO, thus obtained is thoroughly washed with cold water, and dried at 100° (Sandall, J. pr. 14, 254).—8. A solution of SnCl, is ppd. by a slight excess of KOHAq in the cold, the pp. is thoroughly washed and dissolved in cold KOHAq (c. 1 part KOH in 10 parts H2O); the solution, which should be saturated with SnO.xH.O, is allowed to stand in the air, when SnO is gradually ppd. as small, blue-black, shining crystals (Ditte, A. Ch. [5] 27, 145).

Properties and Reactions.—Prepared by any of the methods described above, SnO forms small, black, lustrous, regular crystals (for crystalline form v. Nordenskjöld, P. 114, 612). S.G. 6.1 (N., l.c.); 6.6 at 0° (Berzelius; Ditte, A. Ch. [5] 27, 145).

According to Fremy (A. Ch. [3] 12, 460), another modification of SnO is obtained by heating the black crystals (prepared by the first process given above) to 258°; the crystals are said to swell up and change into soft, olive-green

laminæ, without any change in weight.

By evaporating very dilute NH, ClAq, holding SnO.xH2O in suspension, until NH2Cl began to crystallise, Fremy (A. Ch. [3] 12, 460) obtained a cinnabar-red powder, which he took to be a third form of SnO. Roth (A. 60, 214) obtained red, crystalline SnO by digesting SnO.xH2O, at 56°, with a solution of SnO.xH₂O in acetic acid; the solution containing a little free acetic acid, and having S.G. 1.06.

For the S.G. and appearance of SnO prepared

in various ways v. also Ditte (l.c.).

SnO is unchanged in air at ordinary temperatures. According to Ditte (A. Ch. [5] 27, 145), SnO that separates from an alkaline solution is unchanged at 300°-310°; when heated to redness it is partly decomposed to Sn, and SnO2 which combines with unchanged SnO to give 2SnO.SnO₂. SnO is readily converted into SnO₂ by heating with oxidising agents; the change is effected by heating to 500° in NO (Sabatier a. Senderens, C. R. 114, 1429). SnO dissolves in acids to form stannous salts, SnX2, X = NO3, ½SO₄, ½PO₄, &c.; it is not acted on by NH₂Aq; boiled with fairly conc. KOHAq or NaOHAq it gives a solution of an alkali stannate (M2SnO2) and tin; heated in Cl forms SnCl, and SnO2; mixed with S and strongly heated, SnS2 and SO2 are produced; SnO is reduced to tin by heating to redness in H or with C.

HYDRATED STANNOUS OXIDE. The white pp. formed by adding an alkali carbonate to solution of a stannous salt, washing with air-free cold water, and drying at a temperature not above 80°, is said to have the composition 2SnO.H.O=Sn.O(OH). According to Ditte (A. Ch. [5] 27, 145) the pp. formed by adding KOHAq or NaOHAq to SnCl2Aq, washing thoroughly, and drying in vacuo at 14° is SnO.2H₂O(=SnO₂H₂H₂O). Stannous hydroxide is a yellowish white amorphous powder. It is dehydrated, to SnO, by heating in CO2; also by the action of boiling water containing a little KOH, or a trace of HCl, or acetic acid-HNO,Aq and H2SO4Aq form stannous salts; also by boiling with NH ClAq (Ditte, l.c.). The hydroxide is gradually oxidised by exposure to air to SnO₂xH₂O. Boiled with conc. KOHAq it gives K,SnO,Aq and tin. Many metallic salts are reduced, to lower salts or to the metals, by the action of SnO.xH2O (for details of the interaction with CuO in presence of alkali v. Lenssen. J. pr. 89, 90).

STANNIC OXIDE SnO. (Dioxide of tin.) Mol.

w. unknown; probably at least $Sn_{10}O_{20}$ (v. Carnelley a. Walker, C. J. 53, 92).

Occurrence.—Tinstone is more or less pure SnO₂; the percentage of the oxide varies from c. 85 to c. 99, the other constituents are generally SiO₂, and oxides of Al, Fe, and Mn. Tinstone crystallises in quadratic forms. Crystalline SnO₂ has been found in the fused slag from a furnace used for casting gun-metal (Abel, C. J. 10, 119).

Formation.—1. By exposing molten tin to the action of the air (Emich, M. 14, 345).—2. By heating to c. 600° SnO₂,xH₂O formed by the interaction of tin and HNO₃Aq, or of alkalis and stannic salts, or of alkali stannates and dilute acids.—3. By strongly heating SnO or SnO.xH₂O.—4. By passing a mixture of vapour of SnCl₄ and steam through a red-hot tube.—5. By heating SnC₂O₄, in small quantities, in contact with the air (Vogel, C. C. 1855. 413).—6. By passing a little CO₂ into a dilute solution of an alkali stannate (Ditte, A. Ch. [6] 30, 282).

Preparation.—A. Crystalline stannic oxide.—1. Molten tin is kept in contact with the air until the surface becomes covered with crystalline nodules of SnO₂ (Emich, M. 14, 345).

2. A stream of CO₂ is passed through SnCl₄, and then through a red-hot porcelain tube through which a current of steam is passed at the same time (Daubrée, C. R. 29, 227).—3. Amorphous SnO₂ is strongly heated in a rapid current of dry HCl (Deville, C. R. 53, 161).—4. SnC₂O₄ is strongly heated, in small quantities at a time, in an open porcelain or silver dish (Vogel, C. C. 1855. 413).

B. Amorphous stannic oxide.—5. The hydrated stannic oxide obtained by ppg. a stannic salt by an alkali, decomposing an alkali stannate solution by dilute acid, or by treating tin with HNO,Aq, is thoroughly washed and dried, and then heated to c. 630°, at which temperature dehydration is complete (Carnelley a. Walker, C. J. 53, 83).

Properties .- Crystalline stannic oxide is a hard, lustrous, white solid. It is dimorphous. Prepared by heating the amorphous oxide in HCl gas, SnO₂ crystallises in quadratic forms isomorphous with tinstone and anatase (TiO₂) (Daubrée, C. R. 29, 227); prepared by decomposing vapour of SnCl, by steam, it crystallises in trimetric prisms isomorphous with brookite (TiO₂). S.G. of crystalline $SnO_2 = 6.7$ to 6.85 (Playfair a. Joule, C. J. 1, 187; Mallet, J. 3, 705; Bergemann, J. 10, 661; Daubrée, C. R. 29, 227). The crystals obtained by long continued heating molten tin in air had S.G. 7.0096 at 4° (Emick, M. 14, 345). Crystalline SnO₂ is hard enough to scratch glass. The amorphous oxide has S.G. 6.6 to 6.9 (P. a. J., *l.c.*; Herapath, P. M. 64, 321; Boullay, A. Ch. [2] 43, 266). It is a hard, yellowish-white powder. SnO₂ has not been fused. It is not acted on by acids (but v. Reactions, No. 6). Fusion with KOH or NaOH forms M2SnO3 which dissolves in water. The product of fusion with KHSO, dissolves in water, but addition of more water ppts. all the tin as SnO_2xH_2O .

Mallet (C. J. 35, 524) obtained a compound SnO₂HCl, to which he gave the formula SnO₂OH.Cl, and the name *chlor-stannic acid*, by keeping SnCl₂ for a year or two in a loosely alosed bottle.

Reactions.—1. Fusion with sulphur produces SnS₂ and SO₂.—2. When strongly heated in chlorine SnCl₄ is formed.—3. SnO₂ is reduced to tin by heating to a high temperature in hydrogen, or with potassium, sodium, or carbon, or in carbonic oxide.—4. Fusion with caustic potash or caustic soda forms alkali stannate (M₂SnO₃), which dissolves in water.—5. Heating with phosphorus trichloride to 160° forms SnCl₂, SnCl₄, and P₂O₅ (Michaelis, J. Z. 6, 239; 7, 110).—6. According to Ditte (C. R. 104, 172), SnO₂ dissolves very slowly in hot sulphuric acid (1 acid to 8 water), and on concentrating till not more than 3 or 4 vols. water are present to 1 vol. H₂SO₄ crystals of SnO₂:2H₂SO₄ separate.

HYDRATED STANNIC OXIDE. Carnelley a. Walker (C. J. 53, 60, 68, 83) examined the dehydrating action of heat on SnO₂.xH₂O, obtained by decomposing Na2SnO2Aq by cold dilute HClAq and drying the pp. in the air for five months. The pp. lost less and less weight for each increase of c. 10° from 50° to c. 110°, at which temperature rather less water was present than corresponded with the formula SnO2.H2O, the loss of weight for each rise of 10° was then approximately constant up to c. 360°, at which temperature the composition was nearly that required by the formula 3SnO2.H2O; at a little above 360° the solid changed colour from brown to pale yellow, and at the same time lost weight at a rate nearly three times as great as during the previous rise of 100°; after changing colour the solid had the composition 7SnO₂.2H₂O; the loss of weight, per 105 increase, then decreased very much for the next 30° or 40°, and after that dehydration continued irregularly until at 630°-635° the oxide SnO₂ remained. From these results, considered with results obtained by the same method for other hydrated oxides, C. a. W. concluded that probably a large number of hydrates of SnO2 exists, but that none of these is stable through more than a very small range of temperature. The results of J. van Bemmelen (B. 13, 1466) on the dehydration by heat of stannic hydrates, obtained by oxidising tin by HNO, Aq, and by decomposing SnCl, Aq by CaCOs, and on the rehydration of the products obtained by heat, by placing them in air more or less saturated with moisture, at different temperatures, show that the quantity of water of hydration varies with variations in temperature, in the molecular states of the solids, and in the moistness of the surrounding air. Not only does the quantity of water of hydration vary with variations in the molecular state of the hydrates, but the firmness or looseness wherewith the water is held also varies much as the molecular condition varies. According to J. van B. the loosely held water is given up in dry air; and when the product is placed in moist air water is taken up until a state of equilibrium is attained wherein as many molecules of water are taken up as are lost in the unit of time.

The following compositions have been given to different hydrates of SnO₂.

I. Hydrates obtained by decomposing soluble stannates by dilute acids: (1) 3\$\text{SnO}_x7\text{H}_2\text{O}, by drying in a stream of dry air (Fremy, A. Ch. [8] 12, 463); (2) \$\text{SnO}_x2\text{H}_2\text{O}, by drying in the air at the ordinary temperature (Weber, P. 122, 858); (8) \$\text{SnO}_x\text{H}_2\text{O}, by drying in vacuo (Fremy,

(4) \$SnO₂.2H₂O, by drying at 140° (Fremy,
 (5) to these should probably be added (5)
 7SnO₂.2H₂O, by drying at c. 365° (Carnelley a.

Walker, l.c.).

II. Hydrates obtained by oxidising tin by HNO₂Aq: (6) 5SnO₂.10H₂O, by drying at the ordinary temperature (Fremy, l.c.); (7) 5SnO₂.5H₂O, by drying at the ordinary temperature (Thomson, Ann. Phil. 1817. 149), by drying over H_2O_4 (Weber, l.c.), by drying in vacuo, or at 100° (Fremy, l.c.); (8) $5\mathrm{SnO}_2.4\mathrm{H}_4\mathrm{O}$, by drying at 130° (Fremy, l.c.); (9) $5\mathrm{SnO}_2.3\mathrm{H}_2\mathrm{O}$, by drying at 160° (Fremy, l.c.); (10) $2\mathrm{SnO}_2.\mathrm{H}_2\mathrm{O}$, by drying at 55° (Thomson, l.c.).

The hydrates obtained by decomposing stannates by diluteacids, or by ppg. stannic salts by CaCO₃ or BaCO₃, differ in properties from the hydrates obtained by oxidising tin by HNO, Aq; the former are generally distinguished as stannic hydrates, and the latter as metastannic hydrates. The stannic hydrates, dried in air, form hard, semi-transparent lumps, like gum arabic; soluble in the stronger acids, forming stannic salts SnX₄, where X=NO₃, ½SO₄, &c.; soluble in alkali solutions, forming stannates M¹₂SnO₃ (v. Stannates, under TIN OXYACIDS, AND SALTS AND DERIVATIVES THEREOF, infra). Metastannic hydrates are white solids (? crystalline) that do not dissolve in HNO, Aq, or H2SO, Aq (v. infra). These hydrates interact with HClAq, and the product is soluble in water but insoluble in conc. HClAq; according to Weber (P. 122, 358), the solution gives SnCl. 3SnO. 5H.O, when evaporated over H2SO4; by prolonged boiling, with fresh additions of water, all the tin in the solution is ppd. as metastannic hydrate (Fremy, A. Ch. [3] 12, 463; 23, 393). Metastannic hydrates dissolve in KOHAq and NaOHAq, forming metastannates M2H8Sn5O15 (v. Metastannates, under Tin oxyacids, and salts and derivatives тневеог, р. 730).

According to Ditte (C.R. 104, 172), both stannic and metastannic hydrates dissolve in warm H₂SO₄Aq (1 part acid and 8 parts water), and on concentrating till not more than 3 to 4 vols. water are present to 1 vol. H2SO, the solution deposits white crystals of SnO₂.2H₂SO₄: these crystals are very deliquescent; they are decomposed by water; if so much water is added that not more than 43 g. H₂SO₄ are present in 1,000 c.c. of the liquid, a pp. of SnO₂.xH₂O is formed. By dissolving stannic or metastannic hydrate in warm H₂SeO₃Aq, and concentrating, D. (l.c.) ob-

tained crystals of SnO₂.2H₂SeO₄.

For further details of reactions of stannic and metastannic hydrates v. Tin oxyacids, and SALTS AND DERIVATIVES THEREOF (infra).

Oxides of tin other than stannous and STANNIC OXIDES. Oxides which are most simply

regarded as xSnO.ySnO2 seem to exist.

According to Fuchs (J. pr. 5, 318), a greyishwhite, slimy solid, having the composition $SnO.SnO_{g}(=Sn_{2}O_{s})$, is obtained by diffusing freshly ppd. Fe₂O_s $xH_{2}O$ in SnCl₂Aq free from acid, and boiling

 $(2\operatorname{SnCl}_2\operatorname{Aq} + \operatorname{Fe}_2\operatorname{O}_3 = \operatorname{SnO}.\operatorname{SnO}_2 + 2\operatorname{FeCl}_2\operatorname{Aq}).$ This oxide is said to dissolve completely in

NH,Aq, and also in HClAq.

By digesting SnO₂.xH₂O, ppd. by alkali from SnCl, Aq, with SnCl, Aq, Schiff (A. 120, 58) obtained an orange-yellow solid, to which he gave !

the composition SnO.6SnO.5H.O; by digesting SnO₂.xH₂O, formed by oxidising tin with HNO₂Aq, with SnOl₂Aq Schiff (l.c.) obtained a yellow powder, said by him to be SnO.6SnO.9H2O. or when dried at 85° to be SnO.6SnO.4H.O (v. also Tschermak, C. C. 1862. 305).

PURPLE OF CASSIUS. A purple-coloured solid is obtained by adding solution of a stannous salt to dilute AuCl, Aq; this solid has been examined by many chemists, who have generally assigned to it formulæ which represent it as a compound of SnO2, sometimes of SnO.SnO2, and AuO. According to the most recent investigations the composition varies according to the conditions of preparation, and all that can be said with certainty is that the purple solid is a compound of tin, gold, and oxygen (v. Buisson, J. Ph. 16, 629; Bohlen, Ar. Ph. 57, 277; Capaun, J. pr. 22, 153: Fuchs, J. pr. 5, 318; Wächter, A. 68, 116; Figuier, Ph. C. 1844, 724; Debray, C. R. 100, 1035; Müller, J. pr. [2] 30. 252; and, for a bibliography (to 1866), Fischer,

D. P. J. 182, 39). Tin oxyacids, and salts and derivatives thereof. Some of the hydrates of SnO2 interact with alkalis to produce salts wherein tin forms part of the acidic radicle; compounds are also known the acidic radicles whereof contain tin in combination with other negative elements besides oxygen, or tin in combination with both

metallic and nonmetallic elements.

The reactions of the hydrates of SnO2 ppd. from solutions of stannates by dilute cold acids, or ppd. from stannic salts by alkalis, are distinctly different from the reactions of the hydrates of SnO₂ obtained by oxidising tin by HNO_sAq, or by ppn. by dilute acids from solutions of another class of salts which have the same composition as stannates, except that they always contain H and O in the ratio 2H:O in addition to the constituents of stannates. is therefore usual to distinguish the acidic hydrates of SnO₂ as stannic and metastannic acids; to the former is given the composition $H_2SnO_3(=SnO_2H_2O)$, and to the latter the composition $H_{10}Sn_5O_{15}(=5SnO_2.5H_2O)$. Neither of these formulæ is to be regarded as more than an attempt to connect the differences in the chemical behaviour of two compounds having identical percentage compositions (if the H₂O in one class is omitted) with differences in the complexities of their interacting atomic aggregates; the molecular weight of neither compound is known. It is, indeed, doubtful whether a compound with the composition xH,SnO, can exist apart from other hydrates of the composition ySnO₂.zH₂O; it is certain that the composition of the hydrates of SnO₂ varies with very small variations of temperature, by whatever methods these hydrates are prepared (v. p. 726,

Hydrated Stannic Oxide).

That the hydrates of SnO₂, obtained respectively by ppg_solution of stannates by dilute acids and by oxidising tin by HNO, Aq, differed in their reactions, was noticed by Berzelius in 1811 (v. Lehrbuch [5th ed.] 2, 596). Berzelius regarded these compounds as hydrates of two different modifications of stannic oxide; he called the oxide supposed to be a constituent of the hydrates obtained by ppg. solutions of stannates stannic oxide, and designated it as aSnO₃; the other oxide he called metastannic oxide, and designated it as $b\mathrm{SnO}_2$. Gmelin called the hydrate from stannates ordinary stannic acid, and the other hydrate abnormal stannic acid.

STANNIC ACID H2SnO.

Preparation.—By adding cold dilute HClAq to an aqueous solution of K₂SnO₂, prepared by fusing SnO2 with KOH, or by heating tin with KOH and KNO, (v. Potassium stannate, p. 730), a gelatinous pp. is obtained which, when washed and dried in vacuo, has the composition $SnO_xH_2O=H_xSnO_x$ (Fremy, A. Ch. [3] 12, 463). The same hydrate is formed, according to F. (l.c.), by adding CaCO, or BaCO, to excess of SnCl, Aq, washing, and drying in vacuo. (For details regarding the results obtained by different chemists who have examined the compositions of the stannic hydrates, v. Hydrated STANNIC OXIDE, p. 726). Graham (T. 1861. 213) obtained colloidal stannic acid (no analyses are given) by dissolving SnO₂xH₂O ppd. from a stannic salt (? by alkali from SnCl₄Aq) in SnCl₂Aq, and dialysing until all the Cl had passed into the dialysate. Colloidal forms of stannic acid are also obtained (1) by dialysing a solution of the ppd. acid in HClAq or in an alkali; (2) by the action of CO2, or of air, on an alkaline solution of the ppd. acid (J. van Bemmelen, R. T. C. 7, 87; abstract in C. J. 54, 1160 [1888]; the results of experiments on the dehydration of the colloidal acid are given).

Properties and Reactions .- A gelatinous, amorphous solid, drying (in air or in vacuo) to hard, semi-transparent lumps, like gum-arabic. Reddens litmus paper. Vignon (C. R. 108, 1049) found that the heat of neutralisation, by KOHAq, of stannic acid formed by ppg. SnCl₄Aq by KOHAq, decreased when the ppd. acid was kept in contact with water for some days, and decreased more rapidly when the acid was heated with water in a sealed tube for some hours. J. van Bemmelen (J. pr. [2] 23, 324) found that when stannic acid is shaken with solutions of HCl, or H₂SO₄, or with solutions of certain salts, such as KCl or K2SO4, a definite number of molecules of the acid or salt is taken up by the stannic acid so as to form a loose combination therewith; and that the number of molecules thus absorbed varies with the proportion between the quantities of stannic acid and the acid or salt in the solution used, and also with the concentration of the solution used. The loose compounds thus formed are called absorptioncompounds by J. van B. Dissolves in the stronger acids (HNO₃, H_2 SO₄, HCl, &c.). According to Ditte (C.R. 104, 172), when a solution of stannic acid in warm H₂SO₄Aq (1 part acid to 8 parts water) is evaporated until not more than 8 to 4 vols. water are present to 1 vol. H2SO4, and allowed to cool, it yields white deliquescent crystals of SnO2.2H2SO4; and a solution of strnnic acid in warm H₂SeO₄Aq yields a similar crystalline compound SnO₂·2H₂SeO₄. Dissolves in excess of caustic alkali solutions, forming stannates (q. v. p. 780) M₂SnO₃. According to J. van. B. (i.c.) colloidal stannic acid does not form any compound when shaken with dilute cold KOHAq, but only complex molecular aggregates, the composition of which varies considerably with temperature and concentration. Stannic acid, dried at the ordinary temperature by pressure, is gradually changed to metastannic acid by heat; the change begins at c. 55° (J. van B., l.c.; v. also infra, Change of stannic to metastannic acid, and vice versa). For other reactions of stannic acid v. infra, Distinctions between stannic and metastannic acids.

METASTANNIC ACID H10 Sn5O15.

Preparation.—Tin, in thin pieces or granulated, is heated with HNO, Aq, S.G. c. 1.35, until the metal is converted into a greyish-white powder, which is washed with dilute HNO3Aq, and then with water, and dried. The solid is said to have the composition $H_{10}Sn_{5}O_{15}$ (= $5SnO_{2}.5H_{2}O$) when dried *in vacuo* (Fremy, A. Ch. [3] 12, 463), when dried over H₂SO₄ (Weber, P. 122, 358), or when dried at the ordinary temperature (Thomson, Ann. Phil. 1817. 149). For details of the results obtained by different chemists who have examined the compositions of the metastannic hydrates v. Hydrated stannic oxide (p. 726); v. also J. van Bemmelen, R. T. C. 7, 87 (abstract in C. J. 54, 1160 [1888]). It should be noted that the empirical formula assigned to metastannic acid dried in vacuo (Fremy), or dried over H₂SO₄ (Weber), is the same as that a signed to stannic acid dried in vacuo (Fremy); this formula is SnO2.H2O. The different reactions of the two compounds show that if both have the same empirical formula one must be an isomeride or a polymeride of the other. Metastannic acid is also obtained by decomposing an aqueous solution of the sodium salt (v. p. 730, Metastannates) by boiling. Graham (T. 1861. 213) obtained colloidal metastannic acid by adding a little HClAq to the acid obtained by the action of HNO, Aq on tin, dissolving the solid so formed in water, and dialysing. Vignon (C. R. 108, 1049) found that the heat of neutralisation of metastannic acid by KOHAq decreased when the acid was dried at 110°, and decreased very considerably when the acid was heated with water, for some hours, at 250° in a sealed tube.

Properties and Reactions.-A white powder. Insol. in nitric acid. Contact with conc. HClAq for a short time produces a compound (according to J. van Bemmelen, R. T. O. 7, 87; abstract in C. J. 54, 1160 [1888], no definite compound is formed, but only complex molecular aggregates of metastannic acid and HCl) which dissolves in water, but is insol. in HClAq. According to Barfoed (J. pr. 101, 368), the compound with HCl is quite insol. in HClAq S.G. 1.1, and may be purified by washing with acid of that concentration. The solution of this substance in water gives off HCl and H2O, with a very little SnCl4, when distilled; and metastannic acid separates out (Fremy, A. Ch. [3] 12, 463; 23, 893; H. Rose, A. 68, 272). Evaporation of the aqueous solution over H₂SO, is said to give an oxychloride 3SnO₂SnCl₄.5H₂O (Weber, P. 123, 358). Metastannic hydrate dissolves in conc. HClAq after prolonged boiling, forming SnCl, solution (Löwenthal, J. pr. 77, 321). Barfoed (l.c.) says that conc. HClAq gradually converts recently ppd. and moist metastannic acid into stannic acid, and that the quantity of metastannic acid thus changed increases with the quantity of conc. HClAq, the time of contact, and the tempersture. J. van Bemmelen (J. pr. [2] 23, 324) found that metastannic acid forms absorption compounds with HCl, H₂SO₄, KCl, K₂SO₄, &c. (cf. STANNIC ACID, Properties, p. 728). For other reactions of the solution in water of the product of the action of HClAq on metastannic acid v. infra, Distinctions between stannic and metastannic acids. According to Ditte (C. R. 104, 172), metastannic acid dissolves in warm H₂SO₄Aq (1 pt. acid to 8 pts. water), forming a solution which gives crystals of SnO, 2H, SO, when evaporated until not more than 3 to 4 vols. water are present for one vol. H₂SO₄; the crystals are decomposed by water, with ppn. of SnO2xH2O (? metastannic hydrate), when sufficient water is added to insure that not more than 43 g. H₂SO₄ are present in 1,000 c.c. of the liquid. A similar compound was obtained with selenic acid, viz. SnO₂·2H₂SeO₄ (D., l.c.). Allen (C. J. 25, 274) found that metastannic acid dissolved in boiling conc. H₂SO₄, forming Sn(SO₄)₂ solution; by pouring this solution into water some of the tin was ppd., after a time, as stannic acid, but on boiling the whole of the tin came down as metastannic acid. Metastannic acid is said to dissolve slowly in fairly dilute cold KOHAq or NaOHAq; addition of conc. KOHAq or NaOHAq is said to eppt. K or Na metastannate; when the solution in KOHAq or NaOHAq stands in the air metastannic acid gradually deposits. According to J. van Bemmelen (l.c.), colloidal metastannic acid does not form any definite compound with KOH when shaken with KOHAq of different concentrations. By fusing metastannic acid with KHSO, or NaHSO, Allen (C. J. 25, 274) obtained a product which dissolved partially in water; the aqueous solution gave a pp. of metastannic acid on boiling.

Distinctions between stannic and metastannic acids (v. Fremy, A. Ch. [3] 12, 462; 23, 393; H. Rose, P. 75, 1; 105, 561; Löwenthal, J. pr. 77, 321; Barfoed, J. pr. 101, 368; J. van permelen, R. T. C. 7, 87; abstract in C. J. 54, 1160 [1888]).—Stannic acid is soluble in dilute HNO₂Aq, H₂SO₄Aq, or HClAq; metastannic acid is insol. in these acids. Cold conc. HClAq dissolves stannic acid, and the solution gives the reactions of SnCl₄; cold conc. HClAq forms a compound (? molecular aggregates) with metastannic acid, which is insol. HClAq of S.G. 1·1, or greater S.G., but dissolves in cold water.

The following reactions apply to solutions of stannic acid in HClAq on the one hand, and to solutions in water of the substance formed by the action of HClAq on metastannic acid on the other hand. Stannic acid solution, if conc., does not become turbid on boiling; long continued boiling, with additions of water, ppts. all the acid. Metastannic acid solution, even when conc., goes turbid on boiling; long-continued boiling, accompanied by dilution, ppts. all the acid. Stannic acid solution, when distilled, gives off H₂O, with a little HCl and SnCl₄; and a little stannic acid separates. According to Barfoed (J. pr. 101, 368), when a solution of stannic acid in HClAq S.G. 1.1 is distilled, the whole of the tin passes over as SnCl. Metastannic acid solution, when distilled, gives off H2O, with some HCl and a very little SnCl, and the meta-acid separates in the retort. Addition of tartaric acid to the stannic solution, followed by gradual addition of NH,Aq, excess of NH,Aq being avoided, produces no pp. Treatment of the metastannic solution with tartaric acid and NH, Aq (not in

excess) ppts. the meta-acid. SnCl,Aq gives no pp. with the stannic solution; SnCl₂Aq gives a yellowish pp. (? SnO.6SnO₂.9H₂O) with the metastannic solution. Addition of dilute H.SO, Aq to the stannic solution produces no pp., unless the stannic solution is very much diluted. Dilute H₂SO₄Aq produces a pp. even in very dilute metastannic solution; by washing this pp. with hot water, metastannic acid remains (cf. Ditte's experiments are described under both STANNIC ACID and METASTANNIC ACID, pp. 728, 729). stannic solution is not ppd. by conc. HClAq. The metastannic solution, if nearly neutral, is ppd. by conc. HClAq. Addition of K,FeCy,Aq ppd. by conc. HCIAq. Addition of Agency Sn to the stannic solution, in the ratio K FeCy. Sn To present, ppts. all the tin as SnFeCy. To ppt. all the tin from the metastannic solution, the K.FeCy.Aq must be added in c. the ratio 19K FeCy. Sn present; the pp. is not SnFeCy, it is said by some observers to be metastannic acid; J. van B. (l.c.) calls it an absorption compound of metastannic acid and K.FeCy. stannic solution gives no pp. with excess of fairly conc. NaOHAq. The metastannic solution is completely ppd., as sodium metastannate, by excess of fairly conc. NaOHAq.

Change of stannic to metastannic acid and vice versa. A solution of stannic acid in HClAq gradually changes to metastannic acid; the change is the more rapid and complete the more dilute the solution (H. Rose, l.c.; Barfoed, l.c.); by boiling for some time with repeated additions of water the whole of the stannic acid may be converted into metastannic acid. A very conc. solution of stannic acid in HClAq remains unchanged (cf. reaction of meta- acid with conc. HClAq, infra). According to Löwenthal (l.c.), the change from stannic to metastannic acid is stopped by addition of tartaric acid, even when the solution of stannic acid in HClAq is dilute; the amount of change to the meta- acid may be determined (according to L., l.c.) by finding the quantity of K.FeCy, Aq required to completely ppt. the tin in solution. Barfoed (l.c.) says the best way to measure the amount of change is to add excess of fairly conc. NaOHAq, which ppts. all the meta-acid (as a sodium salt) but none of the stannic acid.

Stannic acid is converted into the meta-acid by heat (H. Rose, *l.c.*). J. van Bemmelen (*l.c.*) found that stannic acid dried by pressure at the ordinary temperature is completely soluble in conc. HClAq, but that by heating to c. 55°, or by heating with water to below 100°, some metastannic acid is formed which is insoluble in conc. HClAq.

A solution of stannic acid in excess of KOHAq gradually deposits metastannate of K on exposure to air, with formation of K₂CO₂Aq.

Metastannic acid is gradually changed to stannic acid, according to Barfoed (l.c.), by contact with conc. HClAq; the quantity of stannic acid produced increases with the quantity of HClAq used, the time of contact, and the temperature.

Stannic acid is said to be obtained from the meta-acid by fusing the latter with a large excess of KOH, dissolving in water, and ppg. by cold dilute HClAq (H. Rose, L.c.; Fremy, L.c.).

dilute HClAq (H. Robe, l.c.; Fremy, l.c.).

Oxyacids of Tin other than Stannic and
Metastannic. (1) H₂Sn₂O₇. Spring (Bl. [8] 1,

180) obtained a colloidal solid, drying at 100° to a white mass with the composition H2Sn2O2, by adding excess of BaO2.xH2O to a solution of SnCl, in HClAq, and dialysing the turbid liquid until BaCl, ceased to pass into the dialysate. regarded the new compound as an acid of the hypothetical perstannic oxide SnO₂, analogous to CeO₂, TiO₃, and (?) ZnO₃. (2) H₄Sn₂O₄ and H.Sn.O. The empirical formula of both these hydrates of stannic oxide is SnO2.H2O. According to Musculus (C. R. 65, 961), the hydrates differ somewhat in properties both from stannic and metastannic acids; they are said to be formed by keeping stannic acid under water, and to form K salts, KH₃Sn₂O₆ and KH₅Sn₂O₉.

STANNATES, MI2SnO, and MIISnO. alkali stannates are obtained by dissolving stan-nic acid in MOHAq (M=NH4, K, or Na), and evaporating over H₂SO₄; the stannates of the alkaline earth metals may be obtained by the interactions of solutions of the alkali stannates

with MO_2H_2 or MCO_3 (M = Ba, Ca, Sr). Ammonium stannate. A salt, said to have

the composition $(NH_4)_2SnO_3.SnO_xxH_2O$, is obtained, as a yellowish jelly, by evaporating a solution of stannic acid in NH₅Aq over H₂SO₄ (Moberg, J. pr. 28, 230). When K₂SnO₃Aq is added to NH ClAq, stannic acid is ppd. according to Ditte (C. R. 96, 701).

normalBaSnO₃.6H₂O is said to be obtained by adding BaCl2Aq to K2SnO2Aq; it is described as a heavy, white powder (Moberg, l.c.). By adding K₂SnO₃Aq to excess of BaCl₂Aq, Ditte (l.c.) obtained a basic

salt BaSnO₂.BaO.10H₂O as lustrous leaflets.
Calcium stannate. Ditte (l.c.) obtained the
normal salt CaSnO₃.5H₂O, in white crystals, by adding K.SnO.Aq to excess of CaCl.Aq, washing, and drying at 100°. By heating for some hours a mixture of stannic acid and CaCl2Aq, with a little CaO, D. obtained CaSnO, in regular crys-

tals.

Potassium stannate K₂SnO₂.xH₂O. pared by dissolving stannic acid in KOHAq, or by fusing SnO₂ or metastannic acid with excess of KOH for some time and dissolving in water, and evaporating over H₂SO₄; transparent, rhombic prisms; crystallising with 4H₂O (Fremy, A. Ch. [3] 23, 393), with 3H₂O (Moberg, l.c.). Marignac (Ann. M. [3] 15, 277) obtained the salt, in rhombohedral crystals, by gradually heating 3 pts. metastannic acid with 8 pts. KOH till the mixture boiled, allowing to cool, dissolving in water, filtering, and evaporating. Ordway (Am. S. [2] 40, 173) prepared the pure salt by adding an equal volume of absolute alcohol to a solution of the commercial salt (v. infra), repeatedly treating the syrupy layer that separated with alcohol, drying the pasty mass so obtained by pressure, dissolving in water, evaporating in vacuo, and washing the crystals with alcohol. Kastannate dissolves easily in water; O. (l.c.) gives S. at 10° = 106.6, and at 20° = 110.5; solution reacts alkaline. Insoluble in alcohol. By adding a dilute acid solution sufficient to neutralise 3 of the alkali, a white flocculent pp. of K metastannate is said to be produced (O., l.c.); excess of cold dilute acid of cold dilute acid ppts. stannic acid. K₂SnO₃.xH₂O is dehydrated at a red heat.

Commercial stannate of potash solution is prepared (1) by fusing tinstone with K.S. KOH, or | vol. p. 285.

KNO, and dissolving in water; (2) by boiling tinstone with KOHAq; (3) by fusing tin with KNO, and K₂CO₃, or boiling the metal with KOHAq containing KNO, and KCl (v. Haeffely, D. P. J. 144, 66; Vaughan, B. 5, 396; and DICTIONARY OF APPLIED CHEMISTRY, vol. iii. pp.

843-4)

Sodium stannate Na₂SnO₃.xH₂O. Prepared similarly to the K salt. Crystallises in hexagonal plates with 3H2O (M., l.c.; O., l.c.). Marignac (l.c.) obtained rhombohedral crystals. Jonas (C. C. 1865. 607) obtained rhombic crystals of Na₂SnO₂.9H₂O by recrystallising the commercial salt. Haeffely (D. P. J. 144, 66) obtained crystals of Na₂SnO₃.3H₂O by heating an aqueous solution of S.G. 1.3; on allowing to cool again the crystals dissolved, the S.G. became 1.35, and after a time crystals of Na,SnO,8H,O were deposited. Scheurer-Kestner (Bl. [2] 8,839) obtained crystals of the composition Na₂SnO₃.10H₂O by evaporating a dilute solution of the ordinary salt, free from excess of NaOH, at a low temperature. Na₂SnO₂.3H₂O is more soluble in cold than in hot water; O. (*l.c.*) gives S. at $0^{\circ} = 67.4$, and at $20^{\circ} = 61.3$. Insoluble in alcohol. According to H. (l.c.), an aqueous solution of the octohydrate gives a crystalline pp. of Na metastannate when heated, or when left at the ordinary temperature for some weeks. Commercial stannate of soda solution is prepared similarly to the solution of the potash salt (v. supra).

Strontium stannate. A basic salt 2SrSnO₂.SrO.10H₂O was obtained by Ditte (C. R. 96, 701), in lustrous octahedra, by adding

K₂SnO₂Aq to excess of SrCl₂Aq.

METASTANNATES. Only alkali metastannates have been prepared. The empirical formula of these salts is M₂O.5SnO₂.4H₂O; as they are decomposed by removing water, the formula is generally written as M₂H₈Sn₃O₁₅ (cf. Metastannic acid, p. 728). Metastannic acid dissolves slowly in KOHAq or NaOHAq; on adding conc. MOHAq the salts are ppd. The salts are better obtained by adding conc. MOHAq the salts are ppd. obtained by adding conc. MOHAq to a solution in water of the substance obtained by the interaction of metastannic acid and conc. HClAq (Barfoed, J. pr. 101, 368). Dilute acid solutions ppt. metastannic acid from solutions of metastannates. Metastannates heated with excess of MOH give stannates. For more details regarding $M_aH_aSn_aO_{1a}$ (M = K or Na) v. Fremy (A. Ch. [3] 23, 393); Weber (P. 122, 358); Haeffely (D. P. J. 144, 66).

S. A compound was obtained by ARSENIO-STANNATES. $Na_2O.2As_2O_5.6SnO_2.5H_2O$ Haeffely (D. P. J. 140, 290) by adding excess of HNO, Aq to a boiling solution of Na, SnO, and Na₃AsO₄, and treating the gelatinous pp. with excess of NaOHAq.

OXALO-STANNATES. The compound 2KHC₂O₄.SnO₅.5H₂O was obtained, in lustrous leasiets, by Pechard (C. R. 116, 1513) by dissolving stannic acid in hot KHC, O, Aq and allowing to cool.

PLATINO-STANNATES. Compounds of SnO and SnO₂ with PtO₂, derived from the acids H₂Sn₂Pt₂O₆ and H₂Sn₂Pt₂O₄ (v. Platino-stan-nates, this vol. p. 285). PLATINO-SELENO-STANNATES v. this

SILICO-STANNATES. Bourgeois (Bl. [2] 47, 897) obtained the salt CaO.SiO2.SnO2 in monoclinic, lustrous crystals, by fusing SiO, and SnO2 with excess of CaCl2, and extracting with water.

In connection with stannic and metastannic acids and their derivatives, v. Chlor-stannic acid, under Stannous chloride (p. 721); Tin, oxides and hydrated oxides of (p. 725); and

TIN, THIO-ACIDS AND SALTS OF (p. 735).

Tin, oxybromides of. By adding pieces of tin to the mother-liquor from BaSnBr_s (v. Stannibromides, under Stannio Bromide, p. 720). Rayman a. Preis (A. 223, 323) obtained colourless, prismatic crystals to which they gave the formula Sn₂OBr₂.12H₂O; and from the motherliquor from this compound fine, colourless needles of Sn_sO₂Br_s.10H₂O separated after a time

Tin, oxychlorides of. According to Ditte (A. Ch. [5] 27, 145), when $SnO.xH_2O$, obtained by ppg. a stannous salt by NH, Aq, is boiled with water containing a trace of SnCl2, a gelatinous oxychloride is obtained having the composition Sn₂OCl₂6H₂O (=SnO.SnCl₂.6H₂O); and when this compound is boiled with a little more dilute SnCl₂Aq another oxychloride, Sn₅O₃Cl₄.6H₂O (=3\$nO.28nOl2.6H2O) is formed in small, white, pearly tablets.

The oxychloride SnOCl2 is obtained, according to Scheurer-Kestner (A. Ch. [3] 47, 1), by evaporating SnCl₂Aq at 50° to 60°; also by heating 100 pts. SnCl₂2H₂O, dissolved in 50 pts. water, with HNO₃Aq containing 16 pts N₂O₅.

An oxychloride is also formed by treating metastannic acid with conc. HClAq, pouring off the excess of acid, dissolving in water, and evaporating over H₂SO₄ and CaO; to this oxychloride, which is an amorphous, white solid, Weber (P. 122, 358) gave the formula $Sn_4O_6Cl_4.5H_2O$ (= 3SnO_.SnCl_4.5H_2O). Tschermak (W. A. B. 44, 2) obtained a crystalline oxychloride $Sn_8O_8Cl_{14}$ (= $4SnO_2.3SnCl_4.SnCl_2$) by dissolving moist metastannic acid in hot conc. SnCl.Aq containing HCl.

Mallet (C. J. 35, 524) obtained a semi-transparent, jelly-like solid from SnCl₂Aq which had been kept in a loosely-stoppered bottle for a year or two; to this solid he gave the formula

 SnO_2 .HCl (= SnO.OH.Cl).

Tin, oxyfluoride of. When SnF, is heated in air it is said to form the oxyfluoride SnOF, (=SnO_xSnF₄) (Fremy, A. Ch. [3] 47, 37; Marignac, Ann. M. [5] 15, 221). Tin, oxyiodides of. According to Personne

(C. R. 54, 216), various oxylodides are formed by decomposing SnI₂ by much water; P. analysed four compounds, to which he assigned the compositions $Sn_3O_2I_2$ (= $2SnO.SnI_2$), Sn_2OI_2 $= SnO.SnI_2$), Sn_3OI_4 ($= SnO.2SnI_2$), and Sn_4OI_6 $(=SnO.3SnI_2).$

divided tin in vapour of P, Schrötter (W. A. B. 1849. 301) obtained a silver-white, brittle solid to which he gave the formula SnP; S.G. 6-56; easily acted on by HClAq, but not by HNOsAq. Other crystalline solids have been obtained by heating tin and P together, but their compositions are not determined with certainty (v. Pelletier, S. 55, 106; Berthier, A. Ch. [2] 33, 180; H. Rose, P. 24, 326; Lüpke, C. C. 1890 [ii.] 643). Natanson a. Vortmann (B. 9, 1459) obtained a phosphide of tin by heating glacial phosphoric acid with C and tin; HClAq gave off PH, and left SnP.

Tin, salts of. Tin forms two classes of salts by replacing the H of acids; the stannous salts SnX₂, and the stannic salts SnX₄ (X = ClO₃, NO₂, $\frac{1}{3}$ CO₂, $\frac{1}{2}$ SO₄, $\frac{1}{3}$ P₄O, &c.). The chief salts of oxyacids are the following: (1) Stannous salts: antimonate, arsenate, borate, carbonate, chlorate, chromate, iodate, nitrate, phosphates, phosphite, sulphates, and sulphite. (2) Stannic salts: antimonate, arsenate, bromate, chlorate, iodate, molybdate, nitrate, phosphates, phosphite, selenate, selenite, sulphates, and tungstates (v. NITRATES, SULPHATES, &c.).

Tin, selenides of. Tin and Se form two compounds, SnSe and SnSe₂. The former can be prepared by the direct union of the elements; the latter does not seem to be obtained by this method. Both selenides dissolve in alkali solutions, and in solutions of alkali sulphides.

STANNOUS SELENIDE SnSe. (Tin mono-selenide or protoselenide.) Mol. w. unknown. Obtained by melting together Se and tin (Berzelius; Welsmann, A. 116, 122). With excess of Se the disulphide SnSe, was said to be formed (Little, A. 112, 213), but this was probably a mistake (Schneider, P. 127, 624). Also formed by adding powdered Se to molten SnCl, heating till SnCl, is volatilised, and allowing to cool, when SnSe crystallises; excess of SnCl, is removed by washing with dilute HClAq (Schneider, l.c.). Steel-grey, lustrous prisms; probably isomorphous with SnS; S.G. 5.24 at 15° (S., l.c.). Also obtained, as a black powder, by passing H₂Se into SnCl₂Aq, washing, and drying at 100°. SnSe prepared by ppn. is soluble in alkali solution; the crystals prepared by heating Se with SnCl, are insoluble, even in boiling alkali solutions. Both crystalline and amorphous SnSe are soluble in solutions of alkali sulphides or selenides. SnSe is insoluble in HClAq; oxidised by HNO₂Aq to SnO₂.xH₂O, H₂SeO₃, and H₂SeO₄. Heated with I, forms SnI, and SnSe2; or, with an excess of I, SnI, and Se; reacts similarly with Br. SnSe is not reduced by heating in H. Heated in air it is gradually oxidised to SnO2 and SeO₂ (S., I.c.). According to Ditte (C. R. 96, 1790), SnSe can be sublimed in a stream of H at a red heat; D. says S.G. is 6.179 at 0°.

STANNIC SELENIDE SnSe. (Tin disclenide.) Mol. w. unknown. Not formed by heating together tin and Se, as excess of Se over that required to form SnSe sublimes : Little's statement that SnSe₂ is formed by directly combining tin and Se by heat (A. 112, 213) is probably wrong (Schneider, P. 127, 624). Prepared by rubbing 5 pts. I and 8 to 10 pts. crystallised SnI₂ (the SnI₄) is used to enable the I to be thoroughly powdered), adding 4 pts. powdered SnSe, then sufficient CS₂ to form a pasty mass (stirring constantly), and then ædding more CS₂ to dissolve SnL, and washing with CS2, when SnSe2 remains as a dark, red-brown, indistinctly crystalline powder, which becomes darker when dried at 100° (Schneider, l.c.). S.G. 1.85. According to Welsmann (A. 116, 122), SnSe, is obtained, as a dark reddish-brown powder, by passing H.Se into SnCl,Aq; heated in H, this powder gives off Se and leaves SnSe.

SnSe, is not acted on by water, or by dilute acids; it is slowly attacked by conc. boiling HClAq; aqua regia, or conc. HNO, Aq, oxidises it to SnO2, H2SeO3 and H2SeO4; it dissolves in hot conc. H,SO, and when the olive-green solu-tion is poured into water Se separates and Sn(SO₄)₂ remains dissolved. SnSe₂ is easily soluble in caustic alkali solutions, including NH,Aq, forming blood-red liquids. with I, in the ratio SnSe2:4I, SnI, and Se are formed; Br reacts similarly to I (Schneider, l.c.).

Tin, silicides of. No definite compound has been isolated. Substances which seem to be No definite compound of the nature of alloys are formed by heating together tin and Si; treatment with HClAq dissolves tin from these bodies, and separates Si along with SiO₂ (v. Winkler, J. pr. 91, 193).

Tin, silicofluoride. According to Berzelius (Lehrbuch [5th ed.] 3, 767), the salt SnSiF₆ separates (? from solution of SnO.xH₂O in H₂SiF₆Aq) in prismatic crystals, which are

readily crystallised from water.

Tin, sulphides of. Two sulphides of tin are known, SnS and SnS2: the former is readily obtained by heating tin and S; the latter by heating tin with an excess of S and some substance (e.g. NH,Cl) which readily volatilises, and so removes heat, which would else drive off the S above that required to form SnS. Both sulphides are also formed by ppn. by H2S from corresponding salts in solution. Both sulphides disselve in solutions of alkali polysulphides, forming thiostannates, M., SnS,.

STANNOUS SULPHIDE SnS. (Tin monosulphide

or protosulphide.) Mol. w. unknown.

Preparation.—1. Finely divided tin is heated with c. equal parts of S; the product is powdered, and repeatedly heated with S in a closed vessel. Ditte (C. R. 96, 1790) recommends to sublime the SnS thus prepared, by heating to redness in a porcelain tube in a stream of H. 2. H₂S is passed into a solution of a stannous salt until the liquid smells strongly of HoS; the brownish black pp. of amorphous SnS is washed and dried, and is then added, little by little, to molten SnCl₂ as long as it is dissolved thereby; after cooling, SnCl₂ is dissolved out by dilute HClAq, and any dark-brown powder that is present is washed away from the heavier, greyish, lustrous, crystalline particles of SnS (Schneider, P. 95, 167).

Properties.—A dark, lead-grey, crystalline solid; S.G. 4.85° (Karsten, S. 65, 394), 5.27 (Boullay, A. Ch. [2] 43, 266). Ditte (l.c.) describes SnS, after sublimation in H, as crystallising in thin, lustrous squares with an angle of c. 90°; with a grey-blue, metal-like lustre; soft and friable; S.G. 5.0802 at 0°. Crystalline SnS melts at a red heat; it expands considerably on cooling (Ditte, l.c.). Prepared by ppn. SnS is a

brownish-black, amorphous solid.

Reactions.—1. SnS, prepared by heating together tin and S, may be sublimed in hydrogen at a red heat (Ditte, C. R. 96, 1790); but continued heating in H is said to reduce it to tin, H₂S being given off.—2. Heated in air SnS is gradually converted into SnO2 and SO2-3. Reacts with chlorine, even at the ordinary temperature, to form SnCl, and SnCl, 2SCl, = (SnS, Cl, 2) (H. Rose, P. 47, 517).—4. Fusion with potassium cyanide produces tin and KONS.—5. SnS dissolves gradually in boiling hydrochloric acid, forming SnCl₂ solution and giving off H₂S (for details of interaction with HClAq of different concentration, and with HCl gas, v. Ditte, C. R. 97,42).—6. Gradually oxidised to SnO₂ by heating with *nitric acid.*—7. Dissolves in solutions of alkali polysulphides, forming alkali thiostannates M₂SnS₃.—8. SnS is generally said to be nearly insoluble in solutions of alkali monosulphides (M₂S). According to Ditte (C. R. 94, 1419, 1470), SnS is not acted on by K2SAq at the ordinary temperature if the concentration of the solution does not exceed 20K₂S:100H₂O; but a more conc. solution of K2SAq, out of contact with air, gradually converts the SnS into a grey, spongy mass of tin; and still more conc. K2SAq dissolves this, forming K2SnS2Aq and giving off H: if air is admitted the reactions are more complex.

STANNIC BULPHIDE SnS₂. (Tin disulphide.)

Mol. w. not known.

Preparation.-1. By saturating SnCl, Aq containing a little HClAq with H2S, warming, again saturating with H.S, warming gently for some hours, collecting the pp., washing with dilute H2SAq, drying, and heating to above 1000 out of contact with air. Pure SnS2 can scarcely be prepared in this way; there seems to be always some SnO2.xH2O present in the pp. (Kühn, A. 84, 110; Barfoed, J. pr. 101, 368).—2. By passing the mixed vapours of SnCl, and H_2S through a red-hot porcelain tube, or by passing H2S into SnCl, and heating the white crystals of SnCl₄.5H₂S so obtained (Coldridge, P. M. [5] 29, 383).—3. A mixture of finely-divided tin and S, or of SnS and S, with some substance that gradually volatilises and so removes heat, is slowly heated to redness in a glass retort, or a loosely covered flask, imbedded in sand; the volatile substance gradually passes off, then the excess of S is volatilised and the SnS2 remains, partly on the sides and partly on the bottom of the vessel. If tin is heated with S only, the heat produced in the reaction is so great that the SnS₂ formed is resolved into SnS and S. Various mixtures have been employed by different chemists; the following give good results: 1) equal parts sifted tin-filings, S, and NH Cl (Pelletier); (2) 4 parts tin-filings, 3 parts S, 2 parts NH₂Cl (Woulfe); (3) a pulverised amalgam of 12 parts tin and 6 parts Hg, with 7 parts S, and 6 parts NH,Cl (W.); (4) 5 parts SnS and 8 parts HgCl, (W.).

References.—Pelletier (Crell's Chem. Ann. 1797 [1] 46); Woulffe (ibid. 1, 149); Bullion (ibid. 1793 [1] 89); Proust (Gehlen's Journ. f. Chem.

und Phys. 1, 250).

According to Gmelin (Handbuch [5th ed.] 3,75) if NH,Cl is heated with tin and S there is formed a compound of NH₄Cl and SnCl₂, which then interacts with the S, forming SnS₂ (?2Sn + 8NH₄Cl = $2(2NH_4Cl.SnCl.) + 2H_2 + 4NH_3$; $2(2NH_1Cl.SnCl_2) + 2S$

 $= SnS_2 + 2NH_4Cl.SnCl_4 + 2NH_4Cl).$

Properties .- Prepared by sublimation, SnS, is a soft golden-yellow, lustrous, crystalline solid; S.G. 4.6 (Karsten, S. 65, 894), 4.42 (Boullay, A. Ch. [2] 43, 266). Crystalline SnS₂ is known BS mosaic gold (v. DICTIONARY OF APPLIED CHEMISTRY, vol. iii. p. 845).

Reactions and Combinations.-1. Heated in a closed vessel gives SnS and S, and a sublimate

of SnS2-2. Heated in air SnO2 and SO2 are formed.—3. Chlorine liquefies SnS2, on cooling yellow crystals of SnCl₄.2SCl₄ are formed (H. Rose, P. 42, 517). — 4. The compound SnS₂I₄(=SnSI₂.SI₂) is said to be formed by heating SnS₂ with *iodine* in a stream of CO₂ (Schneider, J. pr. 79, 419; v. also Tin, thio-IODIDE, infra). According to Schneider (l.c.) a boiling alcoholic solution of I does not act on SnS₂ prepared by sublimation, but with amorphous SnS2 prepared by ppn. it gives SnS2I4-5. Amorphous, but not crystalline, SnS₂ is slowly decomposed by boiling conc. hydrochloric acid, H2S being given off and SnCl, solution formed.—6. The amorphous sulphide is slowly oxidised by hot nitric acid; aqua regia oxidises both amorphous and crystalline SnS, to SnO, and H2SO4Aq. -7. Fusion with lead monoxide produces a mixture of sulphides and oxides of tin and lead; with excess of PbO, SO2 is given off and lead remains. - 8. SnS₂ was said by Dumas (S. 66, 409) to combine with stannic chloride, forming SnS₂.2SnCl₄; the compound being produced by the interaction of SnCl, and H2S. According to Coldridge (P. M. [5] 29, 383) this compound does not exist, the product of the reaction being SnCl_{4.5}H₂S.—9. SnS₂ dissolves in alkali sulphide solutions, forming thiostannates (q. v., infra), M₂SnS₃, — 10. In alkali solutions SnS₂ dissolves, probably forming stannates and thiostannates (?3SnS2+6KOHAq

= K₂SnO₂Aq + 2K₂SnS₂Aq + 3H₂O).
Tin Sesquisulphide. Berzelius (*Lehrbuch* [5th ed.] 2, 600) gave the formula Sn₂S₂ to a greyish-yellow, lustrous solid obtained by heating a mixture of 3 parts SnS and 1 part S in a retort; the substance was almost certainly a

mixture of SnS and SnS2.

Tin, sulphochlorides of; v. Tin thiochlorides, infra.

Tin, sulpho-iodide of; v. Tin thio-iodide, infra.

Ta.
Tin, sulphocyanide of; v. vol. ii. p. 352.

Tin, telluride of, SnTe. A grey, lustrous, metal-like, crystalline solid; S.G. 6.478 at 0°; obtained by heating together tin and Te, and slowly subliming in a stream of H (Ditte, C. R. 96, 1700)

Tin, thio-acids and salts of. It is doubtful whether a thio-acid of tin has been isolated with certainty; a few salts derived from the acid

H2SnS, have been obtained.

THIOSTANNIC ACID. (Sulphostannic acid.) Kühn (A. 84, 110) obtained an olive-brown pp. by adding HClAq or H.C₂H₂O₂Aq to a conc. solution of Na₂SnS₂.Na₂S.12H₂O (v. infra); after washing, and drying at 100°, the pp. was a leadengrey, lustrous solid having the composition Storch (M. 10, 255), repeating Kühn's experiments, always obtained brown pps., which did not contain more S than required by the ratio Sn:S=1:2.19. S. concluded that the pps. were mixtures of H2SnS, and SnS2. By adding dilute oxalic acid solution to solution of Na, SnS, (prepared by saturating Na, SnO, Aq with H,S) in quantity just sufficient to combine with the Na present, S. (l.c.) obtained a deepyellow liquid which remained clear for hours; after removing the H2S by a current of air S. found that the liquid contained tin and S in the ratio Sn:38; he concluded that H.SnS, was present in the liquid. The liquid was decolourised by NH₂Aq, KOHAq, Na₂CO₂Aq, (NH₃)₂CO₂Aq, and NH₄ClAq; strong acids gave brownish pps. containing rather more S than required by the formula SnS₂.

THOSTANNATES (Kühn, l.c.; Höring, Hirsel's Zeit. für Pharm. 1851. No. 8). The alkali salts are obtained by dissolving SnS, in alkali sulphide solutions, M,SAq; the alkaline earth salts are formed by double decomposition from the alkali salts.

Potassium thiostannate K₂SnS₂.10H₂O. Obtained by dissolving SnS₂ in K₂SAq, and adding alcohol, when the salt separates as a dark brown heavy oil; all H₂O is given off at 100°.

Sodium thiostannates. The normal salt Na₂SnS₂.2H₂O is obtained, in yellow, glassy, regular crystals, by adding tin, little by little, to molten Na₂S_s, treating the fused mass with water, and evaporating at a low temperature. A solution of SnS and S in Na SAq deposits colourless, monoclinic crystals of the basic salt, which, when dried over H₂SO₄, has the composition Na₂SnS₅.Na₂S.12H₂O. The stronger acids (HCl, H₂SO₄, CHCl₂CO₂H) ppt. SnS2 at once from Na2SnS, Aq; but weak acids (H₂PO₄, H₂C₂O₄, CH₂Cl.CO₂H, &c.) form brownyellow solutions from which red-brown pps. separate more or less slowly (Storch, M. 10, 255). Kühn (l.c.) gives the reactions of the two Na thiostannates with solutions of several metallic salts; many of the pps. were doubtless thiostannates of the metals employed.

PLATINO-THIOSTANNATES. Schneider (Z. [2] 5, 629; 6, 270, 613) obtained salts to which he gave the composition M₂Pt₂SnS₄ (=M₂S.3PtS.SnS₂), where M=K or Na, by fusing SnS₂, Pt, M₂CO₃, and S, and lixiviating

with water.

Tin, thiochlorides of. A compound SnS₂Cl₁₂ (= SnCl₄.2SCl₄) is said to be formed, along with SnCl₄, by the interaction of Cl and SnS or SnS₄ (H.Rose, P.42, 517). Dumas (S. 66, 409) described a compound Sn₂S₂Cl₄(= SnS₂.SnCl₄) as obtained by passing H₂S into SnCl₄; but according to Coldridge (P. M. [5] 29, 383) this compound is not formed. C. (L.c.) says that the passage of H₂S into SnCl₄ produces white crystals of SnCl₄.5H₂S, which are decomposed by heat, giving off HCl and H₂S, and leaving SnS₂.

Tin, thio-iodide of. By melting together crystalline SnS₂ and I, in the ratio SnS₂:4I, allowing the liquid to cool, and either heating in a stream of CO₂, or dissolving in CS₂ and crystallising, Schneider (J. pr. 79, 419) obtained brown, lustrous crystals of SnS₂I₄ (= SnSI₂:SI₂). This thio-iodide is dissolved unchanged by CS₂ or CHCl₃; alcohol separates S; water, or a solution of a caustic alkali, produces SnS₂, S, and HIAq (or MIAq); HClAq or HNO₂Aq decomposes it, with separation of S. The compound SnS₂I₄ is also produced by the interaction of a solution of I in CS₂ and ppd., dried SnS₂(S., I.c.).

M. M. P. M.

TIN ORGANIC COMPOUNDS.

Stannie methide SnMe. (78°). S.G. 2 1·313. V.D. 6·00 (calc. 6·15). Got by heating an alloy of tin and sodium (14 p.c.) with MeI (½ pt.) at 100°-120° (Ladenburg, A. Suppl. 8, 74; cf. Cahours, A. 111, 256; 114, 372). Oil, with ethereal odour. Reduces alcoholic AgNO.

Stannic tri-methylo-iodide SnMe, I. (170°). S.G. 2 2.143. Formed by the action of I on SnMe, Oil. Yields crystalline SnMe, OH and the salts (SnMe₃)₂SO₄, SnMe₃.O.CO.H, and SnMe₃.OAc. Forms the compounds SnMe₃OEt (Ladenburg, B. 3, 358) and SnMe₃I2NH₃ (Ca-

hours, A. 122, 56)

Stannic di-methylo-di-iodide SnMe₂I₂. [30°]. (228°). S.G. 22 2.872. Formed by heating tinfoil with MeI at 150° (C.). Monoclinic crystals (from ether-alcohol). Converted by ammonia into amorphous SnMe2O, which is insol. water, but dissolves in acids forming the following crystalline salts: SnMe₂Cl₂ [90°] (189°), forming trimetric crystals; a:b:c = 834:1: 941.— SnMe₂PtCl_e 7aq, forming trimetric crystals; a:b:c = .888:1:.977. — SnMe₂Br₂. (209°). — SnMe₂SO₄, forming monoclinic crystals (Hjortdahl, C. R. 88, 584).

Stannic ethide SnEt. Mol. w. 234. (181°). S.G. ²³ 1·187. Formed from SnEt, L and EtI (Buckton, A. 109, 218; 112, 223; Frankland, A. 111, 44). Formed also, together with tin and ZnEtCl, by adding fused SnCl₂ to cooled ZnEt₂ and then distilling (Frankland a. Lawrance, C. J. 35, 184). Prepared by heating powdered tin with Etl and the zinc-copper couple; and also by heating ZnEtI with powdered tin at 160° (Letts a. Collie, C. J. Proc. 2, 166). Oil. Has no action at 180° on Al, Na, or Mg. Slowly absorbs sulphur dioxide, forming crystalline SO₂(OSnEt₈)₂, insol. ether and an oil SnEt, SO, Et, sol. ether. Hot conc. HClAq forms ethane and SnEt, Cl. Iodine forms SnEt, I and

Stannous ethide SnEt₂. S.G. 1.558. Formed by adding zinc to a warm solution of SnEt₂Cl₂ (Frankland). Oil. Decomposed at 150° into SnEt, and tin. Br forms SnEt, Br.,

Distannic hexa-ethide Sn_Et_s. (205°-270°). S.G. 2 1·412. V.D. 14·8 (calc. 14·7). Got by distilling SnEt₃I with sodium (Ladenburg, A. Suppl. 8, 66; B. 3, 647). Pungent oil. I forms SnEt, I. Conc. HClAq forms SnEt, Cl., ethane,

and hydrogen.

Stannic tri-ethylo-iodide SnEt_zI. (231°). S.G. ²² 1.833. Formed by the action of EtI on an alloy of tin and sodium (Ladenburg, A. Suppl. 8, 60; B. 3, 353, 647; cf. Löwig, A. 84, 308). Liquid. Combines with NH_s (2 mols.). On distilling with KOHAq it yields SnEt,OH crystallising from ether in prisms [43°] (271°) converted by heat into oily (SnEt₂)₂O, which recombines with water forming the hydroxide. NaOEt converts the iodide into SnEt, OEt (191°) S.G. 2 1.263, which is at once changed by The hydroxide water into the hydroxide. SnEt,OH is sol. water, strongly alkaline, absorbs CO₂ from the air, and is converted by acids into the salts:—SnEt,Cl. [c. 0°]. (209°). Br. (228°) (Cahours). — SnEt, I tCl, .— Br. (228°) (Cahours). — SnEt, Cy. — S.G. SnEt.Br. SnEt, Cy().—SnEt, NO, .—(SnEt,), SO, . Hexagonal prisms terminated by pyramids (Hjortdahl).—(SnEt₃)₂S.—SnEt₃SH.—(SnEt₃)₄PO₄.

SnEt₃OAc.—SnEt₅OBz.—(SnEt₃)₂CO₅.—(SnEt₃)₂CO₅.

(SnEt₃)₂CO₄. aq.—Formate (SnEt₃)CHO₅.— Sutyrate (SnEt₃)C₄H₂O₅ aq. Tartrates (SnEt₃)₂C₄H₂O₆ aq. (SnEt₃)HC₄H₂O₆ aq. Stannic di-ethylo-di-iodide SnEt₄I. [44·5°]. (245°). Formed from tin and Eti by exposure

to sunlight or by heating with EtI at 150° (Frankland, A. 85, 829; Löwig, A. 84, 308; Cahours a. Riche, A. 84, 333). Needles (from alcohol), sl. sol. cold water. May be sublimed. Alcoholic NH, forms SnEt, O, which is an amorphous powder, insol. water, converted by acids into the following salts: SnEt_Cl. [85°]. (220°), forming trimetric crystals; a:b:c=859:1:943.—SnEt_Cl.OH: plates.—SnEt_Br₂ (233°). V.D. 11.6 (calc. 11.7).—SnEt₂(NO₃)₂.—SnEt₂ICy.—
SnEt₂(SCy)₂.— SnEt₂S.— SnEt₂SO₄.— Crystals. SnEt₂(OAc)₂.— SnEt₂C₂O₄.— Formate

SnEt₂(CHO₂)₂.
Staunic ethylo-trimethide SnEtMe₃. 128°). Formed from SnMe, I and ZnEt, (Cahours, A. 122, 59). Oil; converted by I into EtI and SnMe,I.

Stannic di-ethylo-di-methide ShEt₂Me₂. (145°). S.G. º 1.260. V.D. 6.84 (calc. 7.14). Formed from SnEt₂I₂ and ZnMe₂ (Frankland, A. 111, 50; Morgunoff, A. 144, 157). Oil.

Stannic methylo-tri-ethide SnMeEt,. (163°). V.D. 6.72 (calc. 6.65). Formed from SnEt, I and ZnMe, in the cold (C.). Heavy oil.

Stannic tetrapropylide SnPr. (224). S.G. 1.179. Formed from SnPr. and ZnPr. 14 1.179. (Cahours, C. R. 76, 133). Pungent oil.

Stannic tripropylo-iodide SnPr.I. (261°). S.G. 14 1.692. Formed from PrI and an alloy of Na and tin (Cahours a. Demarçay, C. R. 88, 725, 1112). Oil, converted by moist Ag₂O into

crystalline SnEt, OH.

Stannic di-propylo-di-iodide SnPr.I. (272°). Formed from tin and PrI. Oil. Converted by alkalis into amorphous SnPr₂O, insol. water, which yields SnPr₂Cl₂ [81°].

Stannic tri-isopropylo-iodide SnPr.I. (257°).

Got in like manner.

Stannic di-isopropylo-di-iodide

(267°). Yields SnPr₂Cl₂ [57°].

Stannic tri-isobutylo-iodide Sn(C,H,),I. (285°). S.G. 15 1.540. Formed from CH2PrI and a 9 p.c. alloy of Na in tin (Cahours a. Demarcay, C. R. 89, 68). Pungent oil, converted by KOH into Sn(C,H,),OH (313°).

Stannic di-isobutylo-di-iodide Sn(C,H₂)₂I₂. (290°-295°). Formed by heating tinfoil with isobutyl iodide at 125°. Liquid, converted by alkalis into amorphous Sn(C, H,)2O, and by HCl

into Sn(C₄H₉)₂Cl₂ [c. 6°] (261°).

Stannic tetra-isoamylide Sn(C,H11). Formed from isoamyl iodide and an alloy of Na (1 pt.) and tin (6 pts.) (Grimm, A. 92, 383). Liquid. Stannic tri-isoamylo-iodide $\operatorname{Sn}(C_bH_1)_3I$.

 $\operatorname{Sn}(C_{\mathfrak{b}}H_{11})_{\mathfrak{p}}I.$ (304°). Accompanies the preceding compound (C. a. D.). Converted by potash into oily

 $\dot{\mathbf{S}}\mathbf{n}(\mathbf{C}_{\mathbf{s}}\mathbf{H}_{11})_{\mathbf{s}}\dot{\mathbf{O}}\mathbf{H}$ (337°).

Stannic tetraphenylide SnPh. [226°]. (over 420°). Formed by heating bromo-benzene with a tin-sodium alloy and acetic ether (Polis, B. 22, 2915). Colourless prisms, insol. ligroin, v. sl. sol. alcohol, v. sol. boiling benzene.

Stannic tri-phenylo-chloride SnPh.Cl. [105°]. Formed by the action of NaNO₂ on a solution of SnPh₂Cl₂ in HOAc, the yield being 85 p.c. of that represented by the equation 8SnPh.Cl. -2SnPh₂Cl+SnCl₂ (Aronheim, B. 12, 509).

Converted by potash into SnPh₂OH 1 aq [118°].

Stannic di-phenylo-dichleride SnPh₂Cl₂.

Stannic di-phenylo-dichloride [42°]. (335°). Prepared by boiling HgPh, with SnCl, and ligroin (Aronheim, B. 10, 2229;

Triclinic prisms; 11, 2285; A. 194, 145). a:b:c=588:1:1:067; $a=62^{\circ}$ 47'; $\beta=76^{\circ}$ 48' $\gamma = 94^{\circ}$ 3'. Sol. alcohol and ligroïn, miscible with ether. Converted by means of water into SnPh2Cl.OH, an amorphous insoluble powder [187°]. Potash forms SnPh₂O. Ether, alcohol, and sodium-amalgam form SnPh2(OEt)2 [124°].

Stannic di-phenylo-di-bromide Snlh,Br2. [c. 38°]. (230° at 42 mm.). Formed from SnPh, and Br (Polis), and from the oxide and

HBr (A.).

Stannic di-phenylo-chloro-bromide

SnPh,BrCl [39°]. Crystalline. Formed from SnPh2Cl(OH) and HBr.

Stannic di-phenylo-chloro-iodide SnPh,ClI. [69°]. Yellow monoclinic crystals (from ether), decomposed by water.

TIN GROUP OF ELEMENTS. Group IV. in the periodic classification of the elements contains the following members:-

Even series 10 C Ti \mathbf{Th} Zr Ce Odd series 3 5 7 11 9 Si Ge Sn Ph

Subdividing this group in accordance with the chemical similarities of the elements we get the following families:-

Carbon family: C and Si.

Titanium family: Ti, Zr, Ce, -, Th.

Tin family: Ge, Sn, -, Pb.

The carbon family is considered in the article CARBON GROUP OF ELEMENTS (vol. i. p. 682); and that article also gives a sketch of the chemical relations of all the members of the group. The elements of the titanium family are considered under the heading Titanium Group of ELEMENTS (this vol., p. 749). The tin family is considered in the present article.

Ge occurs in small quantities, as sulphide, in a very few rare minerals. Tin and lead are found native in not very large quantities. The chief ore of tin is tinstone, which contains SnO2; SnS, is also found in tin pyrites; and various micas contain more or less SnO2. Galena, containing PbS, is the chief ore of lead; sulphate, carbonate, phosphate, &c., are also found in considerable quantities. Ge is obtained by heating its ore with Na₂CO₃ and S, dissolving in water, decomposing Na₂S by H₂SO₄Aq, ppg. Ag (Ag₂S is a constituent of the ore) by HClAq, then ppg. GeS, by H₂S, oxidising the GeS, to GeO₂ by HNO, and reducing the GeO₂ by heating in H or with C. Tin is obtained by calcining tin-stone, washing, and reducing the SnO₂ by heat-ing with C. To obtain lead, galena is roasted in such a quantity of air that it is partially oxidised, both PbO and PbSO, being formed; by then heating out of contact with much air the PbS, PbO, and PbSO₄ interact to form SO₂ and lead. Tin and lead have been known from very early times. Ge was isolated in 1885.

The table on page 736 presents some of the principal physical and chemical properties of the tin elements.

General formulæ and characters of compounds. Oxides:-MO, MO2; also Pb3O4 and Pb.O.: various hydrates of most of these oxides are known. Sulphides:—MS, MS₂ (?PbS₂). Haloid compounds: — MX, and MX.

Acids: — H,MX, where M = Ge or Sn, and
X = either Br, Cl, F, or I; also H,PbI4; H,SnO2 | H,H,Sn,O12, are to be classed with the weak

and $H_2H_8Sn_5O_{15}$; $?H_2SnS_3$. S alts: $-MX_2$ and MX_4 , where M=Sn or Pb and $X=NO_3$, $\frac{1}{2}SO_4$, PO4, &c. Salts containing M in the acidio radiole: — R¹2MX₈, where M = Ge, Sn, (?Pb), and X is a halogen; R₂MO₈, where M = Sn, or Pb; also $R_2^r PbO_2$; also

R¹₂·H₈Sn₅O₁₅; also R¹₂SnS₅.

The oxides MO are basic; they react with acids to form salts MX_2 , $X = NO_3$, $\frac{1}{2}SO_4$, $\frac{1}{3}PO_4$, &c. (no salts of GeO have yet been isolated). PbO also dissolves in molten KOH, forming K2PbO2. The oxides MO2, where M=Sn or Pb, are both basic and acidic; they react with some acids to form salts MX,; several of these salts have been isolated when M = Sn, very few when M = Pb: they also react with molten KOH or NaOH $(SnO_2xH_2O$ also with MOHAq) to form salts $K_2(orNa_2)MO_3$; when M = Sn two classes of salts are known, stannates M2SnO3, and metastannates, probably M2.H8Sn5O15. GeO2 dissolves in acids, also in molten KOH or K2CO3, but the products of these reactions have not yet been examined. Pb,O4 reacts with dilute strong acids to form PbO₂, and PbX₂ which dissolves; with glacial acetic acid Pb₃O₄ forms Pb(C₂H₃O₂)₂ and Pb(C2H3O2)4. Pb2O3 probably reacts with dilute strong acids to give PbO, and a salt of PbO. The hydrates of MO and MO2 react similarly to the oxides. Some of the hydrates SnO2.xH2O are acidic; one class of these is probably polymeric with the other. There are indications of the existence of an oxide MO₃ in the case of tin; this oxide seems to be acidic.

The sulphides of Ge have not been much investigated; GeS_2 dissolves in alkali sulphide solutions, and is probably acidic. GeS has been gasified and V.D. determined. The sulphides of tin are both basic and acidic: with hot conc. HClAq they give SnCl, and SnCl, respectively; both dissolve in alkali sulphide solutions to form thiostannates M.SnS3. PbS is basic; with acids it gives salts PbX2, and H.S; there are indications of the existence of a higher sulphide, perhaps PbS₂, which will probably form thioplumbates

with the sulphides of strongly positive metals.

The haloid compounds MX, (none isolated yet when M = Ge) may be formed by the direct union of the elements, and by the other usual methods. SnCl2 and PbCl2 have been gasified, and the formulæ are molecular; molecules of the composition Sn₂Cl, probably exist, but are decomposed at c. the boiling-point of the compound. Several salts are known of the form M_2SnX_4 , where X = Br, Cl, F, or I; and an acid HSnCl₂xH₂O and some salts MSnI₃ are also said to exist. H₂PbI, and salts M,PbI, have been isolated; and also several salts coming under the general formula $xPbX_2yNH_4Cl$, where X = Bror Cl. The haloid compounds MX, have been gasified, and V.D.s determined, when M = Ge and X = Cl or \P , and when M = Sn and X = Br or Cl. PbCl, has not been isolated with certainty, but salts of the form M.PbCl, are known, and also some salts xPbCl, yNH,Cl. The acids H.GeF, H.SnBr, and H.SnCl, have been obtained, and also salts derived from these acids; salts M2SnF and M2PbCl6 are also known. GeHCl2 has been isolated. Many oxyhaloid compounds x.MO.yMX.

	GERMANIUM.	TIN.	LEAD.
Atomíc weights.	72.8	118-8	206·4
Melting-	also been determined di	f each element have been go rectly. There is some evid les of tin and lead, in solution 282°	asified. Specific heats have lence in favour of the con- on in Hg, are monatomic.
points. (approx.). Boiling- points.		1,450°_1,600°	1,450°_1,600°
(approx.). Spec. gravs. (approx.).	5.5	7∙3	11-4
Spec. heats. Atom. wts. Spec. gravs.	·0737 13·2	•0559 16·3	*0315 18·1
(approx.).	Double sulphide of Ge and Ag occurs as a rare mineral. GeO ₂ also found in some specimens of euxenite, with oxides of Nb, Ta, and Y.	Metal found in small quantities. SnO ₂ occurs in tolerable quantities widely distributed.	Small quantities of lead found. Chief ore con- tains PbS; widely dis- tributed in considerable quantities.
	by H or C.	Prepared by reducing SnO ₂ by C.	Prepared by interaction of PbS, PbO, and PbSO ₄ ; also by reducing PbO by C.
Physical properties.	Greyish white, lustrous, very brittle. Crystallises in regular octahedra. Melts at c. 900°; slightly volatilised at 1,850° in N or H.	Silver-white, lustrous, malleable, but malleability varies with temperature; tenacity very small. Crystallises very easily in rhombic and quadratic forms. Melts easily (at 232°), and boils between 1,450° and 1,600°.	White, with tinge of blue; lustrous; very malleable and ductile; tenacity very small; very soft. Crystallises easily in regular octahedra. Melts easily (at 330°), but at higher temperature than tin, and boils between 1,450° and 1,600°.
Chemical properties.	Unchanged in air at ordinary temperature; oxidised to GeO, when heated in fine division. Combines readily with Br, Cl, and I when heated, forming GeX,. Heated in HClgas forms GeHCl. Dissolves in H ₂ SO, Aq. No salts have yet been isolated by replacing H of oxyacids by Ge. GeO, dissolves in fused KOH or K ₂ CO ₂ . H ₂ GeF, Aq and salts of this acid are known.	Unchanged in pure air at ordinary temperature. Heated in air gives SnO and SnO ₂ . Heated in Br, Cl, or I forms SnX ₂ and SnX ₄ . Heated with S forms SnS and SnS ₂ ; also forms SnS and SnSe ₂ . Reacts with acids forming two series of salts, SnX ₂ and SnX ₄ , X = NO ₃ , \$SO ₄ , \$PO ₄ , &c. Conc. HNO ₂ produces SnO ₂ xH ₂ O. SnO is basic; SnO ₂ xH ₂ O. SnO ₂ xH ₂	Tarnishes in air, film of oxide (? Pb ₂ O) forming on surface. Melted in air forms PbO and Pb ₂ O ₂ . Combines with halogens to form PbX ₂ ; PbCl ₄ also probably isolated. Combines with S and Se, forming PbS and PbSe; PbS.xS also probably exists. Reacts with acids forming salts PbX ₂ ; X = NO ₃ , ½SO ₄ , ½PO ₄ , &c. PbO is basic; PbO ₂ feebly basic and feebly acidic; Pb(C ₂ H ₂ O ₂), has been isolated, and probably PbCl ₄ ; with molten KOH, PbO ₂ forms K ₂ PbO ₃ . No thioplumbates isolated. No acid H ₂ PbF ₆ , or salts thereof, isolated; but 3KF.HF.PbF ₄ probably exists. Mol. w. perhaps same as at. w. in solution in Hg. Atom is divalent in gaseous molecule PbCl ₂ , and tetravalent in gaseous

acids; the one is probably a polymeride of the other. The salts of these acids show very distinct differences: the acids themselves interact with strong acids to form salts containing tin in the positive radicles. Of acids containing tin and halogens the following have (probably) been isolated: HSnCl₂, H₂SnBr₂, and H₂SnCl₃. H₄GeF₆ is known, and H₂Pbl₄ is said to exist. It is doubtful whether thiostannic acid H₂SnS₂ has been obtained pure, but salts of this acid have been isolated.

The salts of oxyacids containing these metals in the negative radicles are M₂SnO₈ and M₂H₆Sn₃O₁₈, M₂PbO₂ and M₂PbO₈. These salts are generally easily decomposed. Oxyacids and salts of oxyacids of Ge have yet to be searched for.

Salts derived from oxyacids by replacing H by M are known when M=Sn or Pb; no Ge salts of oxyacids have yet been isolated. The tin salts belong to two classes, SnX₂ and SnX₄, where X = NO₂, ½SO₄, ½PO₄, &c.; both classes of salts readily undergo decomposition and show a general tendency to separate off SnO or SnO₂. Almost all the lead salts belong to the class PbX₂; they are more definite and less ready to split off oxide or decompose to basic salts than the corresponding salts of tin. Pb(C₂H₂O₂)₄ is the only lead salt of an oxyacid of the class PbX₄ that has been certainly isolated; it is decomposed by the least trace of moisture, giving a pp. of PbO₂.

The elements of the tin family are distinctly

The elements of the tin family are distinctly metallic in their physical properties, and in most of their chemical properties also; they exhibit, however, non-metallic characters in their highest salt-forming oxides, MO₂, which are acidic towards strong bases, while they also react as basic oxides towards several acids. The tin elements are less chemically metallic, on the whole, than the members of the even-series, or titanium family, of Group IV.; the two distinctly non-metallic elements of the group are the first member of the even series—carbon, and the first member

of the odd series-silicon.

In considering the chemical analogies of the tin elements, the position of the members of the family in their respective series should be looked to (v. table in vol. iii. p. 811). Ge comes in series 5; it is preceded by Ga, Zn, and Cu, and succeeded by As, Se, and Br: tin comes in series 7; it is preceded by In, Cd, and Ag, and succeeded by Sb, Te, and I: lead comes in series 11; it is preceded by Tl, Hg, and Au, and succeeded by Bi and two unknown elements. The highest salt-forming oxides of the elements preceding and succeeding the tin elements in the series are: series 5, CuO, ZnO, Ga₂O₃; Geo.; As.O., SeO., —; series 5, Cuo, 2no, Ga.2., Geo.; As.O., SeO., —; series 5, Ag.O., GdO., In.O.; Sno.; Sb.O., TeO., ?I.O., — series 11, Au.O., HgO, Tl.O.; PbO.; Bi.O., —, —. The preceding oxides GeO. are, on the whole, more basic than the oxides preceding SnO. or PbO.; and the oxides succeeding GeO, are on the whole more acidic than the oxides succeeding SnO2 or PbO. Hence we should expect the three oxides, GeO. SnO. and PbO. to be about equally acidic and equally basic; SnO. perhaps, being the most acidic of the three. Looking at the position of the tin elements in the general periodic system, we may express the relationship of these Vot. IV.

odd-series elements to the first member of the odd series of their group (Si) by the following statements: (1) Pb:Si = Tl:Al = Hg:Mg = Au:Na = Bi:P. — (2) Sn:Si = In:Al = Cd:Mg = Ag:Na = Sb:P. — (3) Ge:Si = Ga:Al = Zn:Mg = Cu:Na = As:P. On the whole, Bi is more unlike P than Tl is unlike Mg, or Au is a public Na: She is more unlike P than In is unlike Na; Sb is more unlike P than In is unlike Al, Cd is unlike Mg, or Ag is unlike Na; and As is more unlike P than Ga is unlike Al, Zn is unlike Mg, or Cu is unlike Na. Hence the differences between Pb and Si, tin and Si, and Ge and Si, respectively, will be very decided. If the relations expressed in the preceding statements are worked out in detail, they show that the differences between Pb and Si will be more marked than those between tin and Si, and still more marked than those between Ge and Si; but at the same time there will not be any very great differences between these three pairs of differences. Lead, then, will on the whole be the most metallic, and Ge the least metallic, of the tin elements. But the three elements will resemble each other closely in so far as their general positive or negative character is concerned. Then, considering the resemblances and differences between consecutive pairs of elements in (1) series 3 and 5 (i.e. Si and Ge series), and (2) series 5 and 7 (i.e. Ge and tin series), and (3) series 7 and 11 (i.e. tin and lead series), it is evident that Ge and Si will be more alike than Ge and tin, and that Ge and tin will more closely resemble one another than tin and lead. Hence, on the whole, we may expect that Ge will form several compounds wherein the element acts distinctly as a non-metal; it is possible that GeH, may be isolated (GeHCl_s exists). Ge will form volatilisable organic compounds; GeO₂ will probably be found to be di- or trimorphous.

In connection with this article v. Carron group of elements, vol. i. p. 682; and Titanium group of elements, this vol. p. 749.

M. M. P. M.

TITANATES, and derivatives of, v. p. 747.

TITANIC ACIDS v. p. 746.

TITANIFLUORHYDRIC ACID H₂TiF₆ v
TITANIFLUORIDES, Combinations, No. 1, p. 742.

TITANIFLUORIDES M₂TiF₆ v. TITANIFLUORIDES, Combinations, No. 2, p. 742.

TITANIUM. At. w. 47.9. Mol. w. not known. Has not been fused. S.G. 5.3 3.5888 (K. Hofmann, B. 26, 1025); some older determinations gave S.G. c. 5.3 (v. Fritz, M. 1892. 772; Wilson, Pr. 32, 457 [1881]). S.H. 0° to 100° 1135, 0° to 211° 1288, 0° to 301° 1485, 0° to 440° 162 (Nilson a. Pettersson, Z. P. C. 1, 84); Fritz (M. 1892. 772) gives 13. S.V.S. 13.4.

Historical.—In 1791 Gregor found indications of a new element in a Cornish ironsand called menaccanits, from the village near which it was found (Crell's Ann. 1791 [1] 40, 103). The element was called menachin by Kirwan. In 1795 Klaproth found a compound of a new element in a mineral known as 'rother Schoerl su Boinik in Hungarm,' and called the element titanium (derived from Titan). In 1797 K. recognised that the chief constituent of the Hungarian mineral was the same as the characteristic constituent of menaccanits examined by

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Gregor (K., Beiträge sur Kentniss der Mineral-körper). The element was isolated for the first

time by Berzelius in 1824 (P. 4, 8).

Occurrence.-Ti never occurs native. The chief ore of Ti is more or less pure TiO2, which occurs in different crystalline forms as rutile and anatase (tetragonal) and brookite (rhombic). Various minerals contain TiO2, combined with SiO₂ and alkaline earths, e.g. titanite or sphene and ferrotitanite; compounds of titanates and niobates are found in some rare minerals, e.g. polycrase, euxenite, and pyrochlore. Varying quantities of TiO₂ occur in many titaniferous iron ores. The ores of Ti are found in small quantities. Some of them, especially the titaniferous iron ores, are widely distributed (for analyses of Ti ores v. H. Rose, P. 3, 163; 14, 501; 15, 276; Mosander, P. 19, 211; Hermann, J.
25, 368; Berzelius, J. pr. 43, 50; Rammelsberg,
P. 1858, 507; Scheerer, P. 64, 489; Groth, Tabellarische Uebersicht der Mineralien [2nd ed.] 85; O. v. d. Pfordten, B. 22, 1485). Mazade (C. R. 34, 952) found traces of Ti compounds in the mineral spring of Negrac in France. For analyses and descriptions of anatase, brookite, and rutile v. Klein (J. M. 1875. 337); G. vom Rath (ibid. 1876. 64); Schrauf (ibid. 1877. 403); Koch (ibid. 1878. 652). Cornu (C. R. 86, 101, 983) obtained spectroscopic indications of the occurrence of Ti in the sun's atmosphere. The bright-reddish crystals that are often found in blast furnaces wherein titaniferous iron ore has been smelted were supposed for many years to be Ti; but Wöhler (A. 73, 34) showed that they contain C and N besides Ti (v. TITANIUM CARBONI-TRIDE, p. 739).

Formation.—1. By heating K₂TiF₆ with K or Na, out of contact with air (v. Preparation) .-2. By heating TiCl, with Na in a closed iron crucible heated in a wind furnace (Nilson a. Pettersson, W. 4, 554; cf. Kern, C. N. 33, 57). According to Robinson a. Hutchings (Am. 6, 74), Ti is formed by heating TiCl, with Na to 130° in a sealed tube; but O. v. d. Pfordten (A. 237, 201) says that very little, if any, Ti is obtained, and that the main product is TiCl_x.—3. By fusing K_zTiF_e with an equal weight of iron filings, and dissolving away the iron by HClAq (Wehrlin a. Giraud, C. R. 85, 288).—4. By heating TiCl, to 180° with Ag obtained by reducing a salt of Ag; Ti₂Cl₈ is also formed (Friedel a. Guérin, A. Ch. [5] 7, 24).

Preparation.—Finely-powdered rutile mixed with double its weight of K2CO, and the mixture is fused in a Pt orucible placed inside an earthenware crucible. The fused mass is powdered and dissolved in the necessary quantity of HFAq in a Pt dish; water is added to dissolve K.TiF. which begins to separate, the liquid is boiled and filtered while boiling; the K2TiFe, which separates from the filtrate in lustrous crystalline crusts, is pressed between filter-paper, washed repeatedly with cold water, again pressed, and recrystallised several times from boiling water (Wöhler, A. 74, 212). The dry K₂TiF, is placed in a porcelain boat in a porcelain (or hard glass) tube connected with a supply of pure dry H; another boat, containing Na, is also placed in the tube; H is passed through the apparatus, and the Na is gradually vaporised and driven over the K_TiF, which is heated.

When the action is completed and the tube has cooled (the H stream being maintained), the solid is treated with water, which dissolves out KF, NaF, and Na (W. a. Deville, A. 103, 280). The Ti is finally washed with ether and dried over H₂SO₄ (v. Kern, C. N. 33, 57). Merz (P. 73, 48) prepared Ti by heating a mixture of six pts. K₂TiF_s and three pts. NaCl with two pts. Na in a glass bulb placed in a sand-bath, while a stream of H was passed through the bulb, then adding zinc-powder and fusing, and dissolving

out the zinc by cold dilute HClAq.

Properties.—A dark-grey or black lustrous powder; has not been obtained crystallised. Ti resembles iron reduced from Fe₂O₂ by H at a

low temperature.

According to Junot (J. 1853. 336), Ti was obtained by him as a silver-white lustrous deposit by electrolysing a solution prepared by dissolving titanate of K in boiling H₂SO,Aq, evaporating to a syrup, and extracting with Na₂SO₄Aq. J.'s experiments do not seem to have been repeated. Lévy (C. R. 110, 1368) obtained hard, steel-white, cubicle crystals, which he took to be nearly pure Ti, by passing TiCl, vapour over Si, B, or several metals heated to bright redness. L. found only 80 p.c. Ti in the crystals. Ti burns brilliantly in a flame. When heated to redness in O, it is oxidised with production of an intensely brilliant white light. also burns in Cl. It decomposes water at 100°. Warm HClAq dissolves Ti easily; it is also at once dissolved by HFAq. Heated by electric sparks, Ti gives an emission-spectrum very rich in lines (v. Thalen, A. Ch. [4] 18, 289; Troost a. Hautefeuille, C. R. 73, 620; Cornu, C. R. 86, 101, 983; Liveing a. Dewar, Pr. 32, 402).

The atomic weight of Ti has been determined (1) by analyses of TiCl, (H. Rose, P. 15, 145 [1829]; Pierre, A. Ch. [3] 20, 257 [1847]; Thorpe, C. J. 47, 108 [1885]); (2) by determining the weight of TiO₂ got from a determinate weight of TiCl. (H. Rose, l.c.; Thorpe, l.c.); (3) by analyses of TiBr, (Thorpe, Lc.); (4) by determining the weight of TiO, got from a determinate weight of TiBr, (Thorpe, Lc.); (5) by measuring S H of Ti TiBr₄ (Thorpe, l.c.); (5) by measuring S.H. of Ti (Nilson a. Pettersson, Z. P. C. 1, 34); (6) by determining V.D. of TiCl₄ and TiI₄ (v. these compounds). Rose's determinations gave values for the at. w. of Ti varying from 47.92 to 48.32; Pierre's values varied from 50.2 to 50.29. The very accurate series of determinations made by Thorpe has settled the at. w. to be almost exactly 48 (48.01 if O=16; 47.9 if 0 = 15.96).

The atom of Ti is tetravalent in the gaseous molecules TiCl, and TiI,. Ti is metallic in many of its chemical properties; it decomposes water, giving off H; the oxides Ti₂O₂ and TiO₂ form corresponding salts; the chloride and iodide (and ? also bromide) can be gasified without decomposition; the sulphide TiS, does not show any acidic characters so far as it has been examined. The chlorides give pps. of corresponding hydrated oxides by reacting with alkalis, &c. On the other hand, Ti exhibits distinctly non-metallic or chlorous properties; hydrates of TiO, act as feeble acids, producing salts wherein Ti forms part of the negative radicle. The acids H.TiF, and H.TiO.F, are known, and also salts derived therefrom. TiOl, is very easily decomposed by

water, giving HClAq and TiO, xH2O, &c. As might be expected from the two-sided character of the reactions of Ti, most of the salts obtained by replacing the H of oxyacids by Ti are basic salts. The oxide TiO, has been isolated; it is probably a superoxide. Ti is closely related to Zr, Ce, and Th; these elements, with Ti, form the even-series family of Group IV. (v. TITANIUM GROUP OF ELEMENTS, p. 749). Ti is also closely related to Ge, Sn, and Pb, which form the oddseries elements of Group IV. (v. Tin GROUP OF ELEMENTS, p. 785). Ti also shows distinct relationships to C and Si, which are the first and second members of Group IV. (v. CARBON GROUP OF ELEMENTS, vol. i. p. 682). The similarity between Ti and Si is shown very distinctly in the titanic and silicic acids, the relations between which may be expressed by such a general equation as $nMH_4O_4 - mH_2O = M_nH_{4n-2m}O_{4n-m}$ (M = Ti or Si). Ti is the only element of Group IV. which is known to form compounds of the

type M₂TiO₂F₄ = Ti(OM)₂F₄ = TiX₄.

Reactions and Combinations.—1. Heated in air or in oxygen, burns brilliantly to TiO2-2. Burns when heated in chlorine, forming Ti₄Cl.—3. TiI₄ is formed by passing vapour of iodine over hot Ti.-4. Combines with nitrogen when heated therewith (v. Nitrides, p. 743).-5. Ti is violently oxidised to TiO, by heating with cupric oxide or red lead .- 6. Interacts with water at 100°, forming TiO2 and H.-7. Dissolves in warm hydrochloric acid, forming Ti₂Cl₆ (Glatzel, B. 9, 1831; Rammelsberg, B. B. 1874. 490).—8. Ti dissolves easily in hydrofluoric acid (? forming Ti.F.); also in warm dilute nitric or sulphuric acid, according to Glatzel (l.c.), the solution in H2SO4Aq deposits crystals of Ti₂(SO₄), on cooling. Conc. hot H₂SO₄ reacts with Ti, giving off SO₂ (? forming Ti(SO₄)₂); conc. hot HNO₄ oxidises Ti to metatitanic acid (q. v., p. 747) according to Weber (P. 120, 287).

Detection and Estimation .- Compounds of Ti are not reduced to metal by heating on char-coal in the blowpipe flame. Ti compounds, if not containing elements which give a colour to microcosmic salt, form a colourless glass when melted in the microcosmic salt bead in the outer blowpipe flame; in the inner flame they give a glass which is colourless when hot, but becomes violet on cooling; the delicacy of the reaction is much increased by adding a little zinc to the bead (Riley, C. J. 12, 13). Solutions of TiO₂.xH₂O in HClAq or H₂SO₄Aq give white pps. of TiO₂.xH₂O on boiling; addition of a piece of tin to a solution of TiO₂.xH₂O in HClAq causes evolution of H and produces a violet-blue solution which becomes rose-coloured on dilution with water (von Kobell, P. 62, 599). A solution of TiO2xH2O in HClAq gives a violet-red liquid (Ti₂Cl₂ solution) by digestion for some time with Cu (Fuchs, A. 56, 819). Ti is estimated as TiO2, ppd. from solutions by a slight excess of NH.Aq. Insoluble compounds of Ti are generally fused with KHSO₄, the fused mass is dissolved in water, and TiO₂.xH₂O is ppd. by long continued boiling.

Titanium, alloys of. The alloys of Ti have not been much examined. Wöhler a. Michel (A. 113, 248; 115, 102) obtained an alloy with aluminium approaching in composition to TiAl by heating a mixture of TiO, KCl, NaCl, and cryolite, and treating with HClAq. By heating a mixture of Ti and Al with KCl or NaCl in an earthenware crucible in a stream of H, Lévy (C. R. 106, 66) obtained brittle steel-grey crystals which he took to be an isomorphous mixture of TiAl, and SiAl. Ti alloys with iron; it is often present in pig-iron, but rarely in bar-iron or steel (v. Riley, C. J. 16, 387).

Titanium, bromides of. Only one bromide

has been isolated, TiBr.
TITANIC BROMIDE TiBr. (Titanium tetrabromide.) Mol. w. has not been determined, but the formula is almost certainly molecular from

analogy of TiCl, and TiL.

Prepared by passing Br vapour over a mixture of TiO2 and charcoal heated to full redness, collecting the liquid that distils over and solidifies in the receiver, and redistilling it from Hg (to remove free Br) (Duppa, Pr. 8, 42). Also prepared by passing HBr gas over TiCl, kept a few degrees below its boiling-point in a distilling flask connected with an upright condenser; the whole of the Cl of the TiCl, is expelled (Thorpe, C. J. 47, 126). Thorpe used 350 g. TiCl, and continued the passage of HBr for three days. The HBr is conveniently obtained by the interaction of Br and amorphous P in a little water; it is freed from Br by passing through a U tube containing broken glass and amorphous P. and is dried by passing through another U tube containing CaCl.

TiBr, forms dark-yellow or orange crystals, S.G. 2.6, melting at c. 40°, and boiling at c. 229° (D., l.c.; T., l.c.). It is very hygroscopic; easily decomposed by water, forming a perfectly clear solution containing HBr and TiO, xH₂O.

Titanium, bromochlorides of. By adding Br

to TiCl₂ (q. v., p. 741) Friedel a. Guérin (A. Ch. [5] 8, 24) obtained a colourless liquid, fuming in the air, boiling at 176°, which they said was probably TiBr₂Cl₂; and by the reaction of Br with Ti₂Cl₂ they obtained a liquid boiling at 154°, probably TiBrCl₂.

Titanium, carbonitride of, Ti, CN. (Titanium nitrocyanide, or cyanonitride Ti, (CN), N. This compound is often found in blast furnaces wherein titaniferous ironstone has been smelted. It was supposed by Wollaston to be Ti (G. A. 75, 220), and was for many years generally mistaken for the metal. In 1850 Wöhler (A. 73, 84) proved it to contain C and N besides Ti.

The compound is found in blast furnaces as lustrous, opaque, copper-coloured, prismatic crystals. By treating these crystals with conc. HClAq, then with conc. H2SO4, washing, treating with HFAq, and again washing, the compound is freed from admixed Fe, &c.; small leaflets of graphite generally remain mixed with the car-bonitride. Ti₆CN₄ can be prepared by heating a mixture of dry K₄FeCy₅ and TiO₅ in a closed crucible for an hour or two at c. the melting-point of Ni (1400° to 1450°), dissolving out Fe, &c., by conc. HClAq, washing, and drying. By examining the brown powder thus obtained with a magnifying power of 300 diameters coppercoloured crystals of Ti,CN, can be detected Wöhler, A. 73, 84). It is also obtained by heating a mixture 1 part C and 5 parts TiO, in a closed carbon crucible for some hours at 1400° to 1450°; also by passing N over a mixture of C and TiO, in the ratio 1:52 heated in a carbon boat placed in a carbon tube to the melting-point of Pt (c. 1800°) (W. a. Deville, A. 103, 231).

of Pt (o. 1800°) (W. a. Deville, A. 103, 281).

When Ti₂CN₄ is fused with KOH it gives off NH₂; heated to redness in a stream of water vapour TiO₂ is formed, and H, NH₃, and HCN are given off (W., l.c.); heated in Cl it gives TiCl₄ and TiCl₄.2HCN (W., l.c.; also p. 219); the compound is burnt by heating with PbO, CuO, or HgO, giving off CO₂ and leaving TiO₂ and Pb, Cu, or Hg (much heat is produced in the reaction) (W., l.c.).

The simplest formula that expresses the composition of Ti carbonitride is Ti₂CN₄ (W., l.c.; cf. Reinhardt, Zeit. für anorgan. Chemie, 1, 124). This formula is generally doubled and written Ti(CN)₂STi₂N₃, and the substance is regarded as a compound of cyanide and nitride of Ti; Joly (C. R. 82, 1195) regards it as a compound of carbide and nitride of Ti, and writes the formula

TiC.4TiN (or 2TiC.8TiN).

Titanium, chlorides of. Ti and Cl combine to form TiCl₄; by the regulated action of H on TiCl₅ one-fourth of the Cl can be removed and TiCl₅ produced; and by the further action of H on TiCl₅ one-third of the Cl can be removed and TiCl₅ produced. The V.D. of the tetrachloride has been determined, and the formula TiCl₄ is molecular; from the analogy of C₂Cl₆ and Si₂Cl₆ it is probable that the molecular formula of the trichloride is Ti₂Cl₆; from the analogy of PbCl₂ and SnCl₂ the formula TiCl₂ may be supposed to be molecular, but if the analogy of C₂Cl₄ is to be followed, the molecular formula must be Ti₂Cl₄.

TITANIC CHLORIDE TiCl₄. (Titanium tetrachloride.) Mol. w. 189·38. Melts at -25° (Haase, B. 26, 1052). Boils at $136\cdot41^{\circ}$ at 760 mm. (Thorpe, C. J. 37, 329). S.G. $\frac{66}{45}$ 1·76041; S.G. at b.p. 1·52223 (Thorpe, l.c.). V.D. 98·8 at 143° (Dumas, A. Ch. [2] 83, 388). S.V. 124·47 (Thorpe,

Lc.

Preparation.—1. TiO₂ is intimately mixed with c. half its weight of lampblack previously heated to redness in a stream of Cl, and the mixture is heated to full redness in a tube of hard glass connected with a receiver, while a stream of dry Cl is passed over it. The TiCl₄ which collects in the receiver is freed from Cl by shaking with Cu, Hg, or Na amalgam (Merz, Bl. [2] 7, 401), or by boiling for some time (Thorpe, C. J. 47, 119), and is then distilled. Thorpe (l.c.) conducted the final distillation in dry N.—2. Vapour of CCl₄, or a mixture of equal volumes of Cl and CO, is passed over TiO₂ in small lumps, heated to bright redness in a tube of hard glass connected with a receiver. The TiCl₄ is purified as described above (Watts a. Bell, C. J. 33, 443).

Properties.—A colourless liquid, solidifying at -25° (Haase, B. 26, 1052). Does not conduct electricity (O. v. d. Pfordten, A. 237, 201). Fumes in the air, rapidly absorbing moisture and solidifying to the hydrate TiCl₄.5H₂O (Demoly, A. 72, 218). (For an apparatus for collecting TiCl₄ for analysis, v. Thorpe, l.c.) TiCl₄ is violently decomposed by water, forming HClAq, and TiO₂.xH₂O some of which ppts. (v. Reactions, No. 8).

Reactions.—1. Hydrogen reduces TiOl, at a high temperature, to Ti₂Ol, and then to TiOl₂ (v. trichloride and dichloride).—2. Mixed with to form yellow crystals of TiOl₂ CNOl; sublimable crystal and passed through a red-hot porcelain at 100° (W., A. 73, 219).—6. With phosphorus

tube, Ti₄O₂Cl₆ is formed (v. Oxychlorides, p. 748). 3. Heated to 130° in a sealed tube with sodium, the chief product is TiCl, according to O. v. d. Pfordten (A. 237, 201); Robinson a. Hutchings (Am. 6, 74) say that Ti is produced in this reaction. Nilson a. Pettersson (W. 4, 554) obtained Ti by reducing TiCl, by Na at a very high temperature.—4. Sodium-amalgam produces Ti₂Cl₈ and TiCl₂ (O. v. d. P., l.c.).—5. Reduced silver gives Ti₂Cl₈ and Ti when heated with TiCl₄ to 180° (Friedel a. Guérin, A. Ch. [5] 7, 24).—6. By passing vapour of TiCl, over silicon, boron, or several metals, heated to bright redness, Lévy (C. R. 110, 1368) obtained hard, steel-white, cubical crystals which he supposed to be nearly pure Ti.-7. Pure, dry, hydrogen sulphide reacts at the ordinary temperature, forming Ti2Ols; at a higher temperature a pp. is produced which is probably a thiochloride of Ti (O. v. d. P., A. 234, 257).—8. Cold water reacts violently with TiCl, with production of much heat and formation of a turbid liquid containing HCl and holding in suspension 2TiO2.H2O (v. Metatitanic acid under TITANIUM OXYACIDS, AND SALTS AND DERIVATIVES THEREOF, p. 747); after 30 or 40 hours the liquid becomes clear, but ppn. occurs again on heating to c. 85° when about 7 pts. water are present to 1 pt. TiCl, used (v. Thorpe, C. J. 47, 120; and cf. TITANIUM OXYACIDS, p. 746). According to Merz (Bl. [2] 7, 401), an oxychloride, approximately TiCl, 3TiO, 16H, O, is obtained by evaporating TiCl, Aq, or TiCl, that has deliquesced in air, over H₂SO₄ and CaO (v. Oxychlorides, p. 748). TiO₂, in the form of brookite, is obtained by passing vapour of TiCl, and water, mixed with CO2, through a red-hot porcelain tube (Daubrée, C. R. 29, 227; 39, 153.—9. With hydrochloric acid the compounds TiCl₂.OH and TiCl₂(OH)₂ are formed (v. TITANIUM HYDROXYL CHLORIDES, p. 743).—10. Chlorosulphonic acid (SO₂.OH.Cl) reacts with TiCl, at the ordinary temperature to form a yellow, amorphous, deliquescent powder, having the composition TiCl.SO, = TiCl₂.OSO₂.Cl) (Clausnizer, B. 11, 2011).—11. By passing vapour of TiCl4, mixed with H, through a white-hot tube containing titanium dioxide the oxychloride Ti2O2Cl2 is obtained (v. Oxychlorides, p. 748).

Combinations.—1. With water to form a pentahydrate and a dihydrate. Demoly (A. 72, 213) obtained TiCl₄.5H₂O, as deliquescent crystals, by allowing TiCl, to absorb moisture from the air, then adding a little more water, and evaporating under reduced pressure. By drying the pentahydrate over H₂SO₄ in vacuo, D. obtained TiCl₄·2H₂O.—2. With ammonia, to form TiCl.4NH, according to H. Rose (P. 16, 57), TiCl. 6NH, according to Persoz (A. Ch. [2] 46, 315); easily decomposed in air; heated in a tube gives off NH₂, then NH₂ and HCl, and a yellowish sublimate of TiCl₂.8NH₄Cl, and leaves Ti; heated in NH₃ gives Ti₂N₄ (v. Titanium nitrides, p. 743). 3. With hydrogen phosphide, to form a brown powder, which when heated gives a sublimate of BTiCl, 2PH, Cl according to H. Rose (P. 42, 527). 4. With hydrogen cyanide to form TiCl. 2HCN (Wöhler, A. 73, 226); by vaporising this compound through a red-hot tube Ti, N, and C are pentachloride to form TiCl₄.PCl₅; obtained by heating TiCl₄ and PCl₅ to 150° in a sealed tube (Bertrand, Bl. [2] 33, 565); also by heating TiO. and PCl, in the ratio TiO,:3PCl, or by saturating a mixture of TiCl, and PCl, with Cl (Tüttscheff, A. 141, 111), and removing excess of TiCl, by heating in a stream of Cl (Weber, P. 132, 452). A citron-yellow, loose, semi-crystalline solid; sublimes without melting; deliquescent in air .--7. With phosphorus trichloride to form yellow crystals of TiCl. PCl., melting at 85.5° (Bertrand, l.c.).—8. With sulphur tetrachloride, to form 2TiCl. SCl.; obtained by saturating a mixture of TiCl, and S2Cl2 with Cl, and warming in a stream of Cl (Weber, P. 132, 454).—9. By mixing TiCl, and sulphurous chloride H. Rose obtained large yellow crystals, probably a compound of the two chlorides (P. 42, 527).—10. With ammonium chloride, to form TiCl. 3NH Cl (v. supra, Combinations, No. 2).—11. With phosphoryl chloride to form TiCl, POCl, obtained by gradually adding POCl, to TiCl, and gently warming (Weber, P. 132, 453); colourless, deliquescent crystals, melting at 110°, and boiling at 140° (Wehrlin a. Giraud, C. R. 85, 288).—12. With selenion oxychloride to form a yellow powder, decomposed by heat, having the composition TiCl₄.2SeOCl₂; obtained by adding SeOCl₂ to TiCl₄, and drying the solid that separates on cooling on a tile over H₂SO₄ (Weber, B. B. 1865. 154).—13. With nitrosyl chloride; obtained by saturating TiCl₄ with NO₂, and subliming (Hampe, A. 126, 47), also by passing the vapours from sets. 126, 170, 170 (Weber, B. 118, 476). from aqua regia into TiCl, (Weber, P. 118, 476). Hampe (l.c.) gives the composition 3TiCl, 4NOCl, Weber (l.c.) the composition TiCl, 2NOCl. A deep citron-yellow, crystalline mass; decomposes in air; sublimable out of air; with water gives off NO.—14. With sulphur trioxide to form TiCl₄.SO₂ (or TiCl₂.OSO₂.Cl), formed by the interaction of TiCl₄ and SO₂.OH.Cl at the ordinary temperature; a yellow, amorphous, deliquescent powder (Clausnizer, B. 11, 2011).-15. Demarçay (Bl. [2] 20, 127) obtained a great many compounds of TiCl, with esters, mercaptans,

and ethyl sulphide (cf. Bedson, C. J. 29, 309).

TITANIUM TRICHLORDE Ti.Cl. (Titanium sesquichloride. Titanous chloride.) Mol. w. not determined; but from analogy of C.Cl. and Si.Cl. the formula Ti.Cl. is probably molecular.

Formation.—1. By reducing TiCl, by H (v. Preparation).—2. By reducing TiCl, by sodiumamalgam (O. v. d. Pfordten, A. 237, 201); or by pure, dry H.S (O. v. d. P., A. 234, 257).—3. By heating TiCl, with reduced Ag to 180°; Ti is also formed, and probably TiCl, also (Friedel a. Guérin, A. Ch. [5] 8, 24).—4. A solution of Ti.Cl. is obtained by digesting a solution of Ti.C. Ti.Cl. is obtained by digesting a solution of Ti.C. XH_O in HClAq with Cu at 40°-50° (Fuchs, A. 56, 319); or with Ag ppd. by zinc from AgCl (von Kobell, P. 62, 599). Ti.Cl. is also probably present in the greenish liquid obtained by adding Na amalgam to K_TiF, in HClAq (O. v. d. P., A. 237, 201).

Preparation.—TiCl, is placed in a retort connected with a tube of porcelain or hard glass (if a glass tube is used it should be wrapped in copper-foil) which is placed horizontally in a furnace; the other end of the tube projects considerably out of the furnace and is connected with a receiver to collect undecomposed TiCl.

The delivery tube from an apparatus evolving pure and dry H passes into the retort beneath the surface of the TiCl. H is passed through the apparatus until it is entirely filled with the gas; the tube is then heated to redness, and the retort is gently warmed. Ti₂Cl₃ is formed and condenses as dark-violet scales on the part of the tube that projects out of the furnace. When all the TiCl₄ has been driven out of the retort, the Ti₂Cl₅ in the cooler part of the tube is gently warmed, while the passage of H is continued, to remove any adhering TiCl₄; the Ti₂Cl₅ is allowed to cool in the stream of H (Ebelmen, A. Ch. [3] 20, 385).

Properties.—A dark-violet, very lustrous solid, deliquescing in air, and dissolving in water to form a violet liquid, which gradually decolourises with ppn. of TiO₂xH₂O. Ti₂Cl₆ is easily soluble in alcohol, forming a violet or green liquid; it is insoluble in ether, and sparingly soluble in HClAg (O. v. d. P., A. 237, 201).

soluble in HClAq (O.v. d. P., A. 237, 201).

Reactions.—1. Heated in hydrogen to c. 440°,
TiCl, distils over and TiCl₂ remains (Friedel a.
Guérin, A. Ch. [5] 8, 24).—2. Heated in air,
TiCl₄ vaporises and TiO₂ remains.—3. Dissolves
in water, forming a violet, or green-violet, liquid
which slowly becomes colourless with ppn. of
TiO₂xH₂O.—4. Alkalis, or alkali carbonates, in
solution ppt. Ti₂O₃xH₂O from Ti₂Cl₄Aq; ammonium sulphide forms the same pp.; hydrogen,
sulphide gives no pp. (Ebelmen, l.c.).—5. Ti₂Cl₄Aq
reduces solutions of gold, silver, or platinum
salts with ppn. of the metals; ferric and cupric
salts are reduced to ferrous and cuprous salts;
sulphurous acid solution is reduced by heating
with Ti₂Cl₄Aq with ppn. of S (E., l.c.).—6. Ti₂Cl₄
reacts with bromine to form a liquid boiling at
154°, probably TiBrCl₄ (F. s. G., l.c.).

154°, probably TiBrCl₂ (F. a. G., *i.c.*).

Combinations. — With water to form Ti₂Cl₆:8H₂O; obtained by Glatzel (B. 9, 1829) by evaporating a solution of Ti in HClAq to the crystallising point, filtering off ppd. TiO₂:xH₂O from time to time. The hydrate is a green solid; it dissolves in water, forming an opalescent violet solution containing a little suspended TiO₂:xH₂O; the solution becomes colourless on standing, and then gives the reactions of titanic

acid.

TITANIUM DICHLORIDE TiCl₂. (Sometimes called titanous chloride.) Mol. w. is not known. From the analogy of SnCl₂ and PbCl₂, the formula TiCl₂ would be molecular; but from the analogy of C₂Cl₄ the molecular formula would be Ti₂Cl₄.

Formation.—1. By reducing Ti₂Cl₆ by H (v. Preparation).—2. TiCl₂ is probably formed, along with Ti₂Cl₆, by heating TiCl₄ with reduced Ag (Friedel a. Guérin, A. Ch. [5] 8, 24).—3. Along with Ti₂Cl₆, by the action of Na amalgam on TiCl₄; also by heating TiCl₆ with Na to 180° in a sealed tube (O. v. d. Pfordten, A. 237, 201).

Preparation.—Ti,Cl_e reacts with H at a red heat to give TiCl₂ and TiCl₄. Friedel a. Guérin (l.c.) proceed as follows. The tube containing Ti₂Cl₆, prepared from TiCl₄ (v. supra), while still filled with H, is connected with an apparatus evolving pure, dry CO₂, and the whole of the H in the apparatus is driven out by CO₂. The Ti₂Cl₄ is then shaken into a small tubulated retort, placed on a sand-tray, and previously filled with CO₂; the CO₂ is displaced by pure, dry H; the retort is then heated to dull red-

ness, while a stream of H is passed through it, until TiCl, ceases to distil off; the retort is allowed to cool in the stream of H, and CO2 is then passed in until all the H is removed (if this is omitted the TiCl, takes fire in the air), and the black TiCl, in the retort is quickly shaken into small dry tubes filled with CO2, which are at once sealed off.

Properties.—A black, light powder; very rapidly absorbs water, forming a kind of mud; insoluble in CS2, CHCl3, or Et2O. Volatilised without melting by heating to full redness in H. Shaken with Et₂O and KCNSAq, the Et₂O becomes dark brown; this is a delicate test for TiOl, (O. v. d. Pfordten, A. 237, 201).

Reactions.-1. Takes fire when touched with a drop of water (? with formation of Ti2O2Cl2; v. F. a. G., l.c.); thrown into water, it hisses like red-hot iron, decomposing the water with evolution of H and formation of a yellowish liquid.—2. Decomposes absolute alcohol, giving off H and forming a yellowish liquid.—8. Heated in air, TiCl₂ burns like tinder, giving off TiCl₄ and leaving TiO₂—4. With bromine it forms a liquid boiling at 176°, probably TiCl₂Br₂ (F. a.

Titanium, chlorobromides of, v. TITANIUM

BROMOCHLORIDES, p. 739.

Titanium, cyanonitride of, v. TITANIUM

CARBONITRIDE, p. 739.

Titanium, ferrocyanides of, v. vol. ii. p. 337. Titanium, fluorides of. The only compound that has been isolated with certainty is TiF. There are indications of the existence of Ti2H and compounds of this fluoride with alkali fluorides are known.

Tranio fluoride Tif. (Titanium tetra-fluoride.) Formula probably molecular, from analogy of TiCl, and Til. By heating a mixture of TiO₂ and CaF₂ with H₂SO₃, Unverdorten (P. 7, 320) obtained a colourless distillate, which he supposed to contain a compound of Ti and F. Berzelius (P. 4, 1) obtained crystals (? TiF₄) by dissolving TiO₂ xH₂O in HFAq. The crystals were decomposed by water, giving an acid solution (? H.TiF.Aq), and an insoluble, 'so to say basic,' compound. TiO2 is obtained by passing vapour of TiF4, mixed with H2O, through a tube heated to c. 800° or 1000° (Hautefeuille, A. Ch. [4] 4, 181; cf. TITANIUM DIOXIDE, p. 741). Heated to redness in H, perhaps gives Ti₂F₆ (H.,

Combinations.—1. TiF₄ probably combines with hydrogen fluorids to form H₂TiF₄. By dissolving TiO₂.xH₂O in HFAq, evaporating, adding water to the crystals (? TiF₄) thus formed, and filtering from an insoluble solid (? oxyfluoride), Berzelius (P. 4, 1) obtained an acid solution from which, by neutralisation with KOHAq and evaporation, he obtained the salt K, TiF. acid solution probably contained H, TiF. compound (if it exists) is best called titanifluorhydric acid. It is analogous to H_*SiF_* , H_*SnF_* , and other acids of the type H_*MX_* , where M= an element of Group IV. (except C), and X= a halogen.—2. With various metallic fluorides to form TiF.2MF = M.TiF. salts were obtained from the K salt, which was got by neutralising the acid solution prepared as described under 1. Salts of NH4, Ca, Cu, Fe, Pb, Mg, and Ni are described (B., Lc.; Weber,

P. 120, 291). These salts are best called titanifluorides; they are similar to stanni-fluorides.—3. TiF₄ also combines with am-monium fluoride to form TiF₄.3NH₄F (besides

TiF₄.2NH₄F) (Baker, C. J. 35, 768).

By adding NH, Aq, drop by drop, to warm (NH,),TiF,Aq, Piccini (G. 17, 479) obtained a compound of the form TiO₂F₂xNH₂F; from this he prepared TiO₂F₂BaF₂, and by carefully decomposing this salt by H₂SO₂Aq he obtained a solution of the acid TiO2F2.2HF. P. regards this acid as a F derivative of the hypothetical pertitanic acid H₂TiO₄ (the soid of TiO₃); he calls the soid fluoroxypertitanic acid, and the salts fluoroxypertitanates. Following the plan adopted in this Dictionary, the soid will be called titanoxyfluorhydric acid H.TiO.F. and the salts, M'2TiO2F4, will be called titan-oxyfuorides. H.TiO2F4Aq is a yellow liquid, not ppd. by Ba salts, giving a pp. of TiO, with alkali carbonates.

TITANIUM TRIFLUORIDE Ti2F6. This compound is said to be formed by strongly heating K2TiF4 in a stream of H with a little HCl, according to Hautefeuille (C. R. 59, 189; cf. 57, 451). According to H., the compound is a purple-red solid, soluble in water. By treating Ti₂Cl₆Aq with KHF₂ or NH₄F, Piccini (G. 16, 104) obtained violet pps. of Ti₂F₆.2MF, where M=NH₄ or K; he also obtained Ti₂F₆.3NH₄F. The K salt is slightly soluble in water; soluble in dilute acids, forming green liquids, from which alkalis throw down azure-blue pps. that gradually oxidise to TiO₂xH₂O on exposure to air. The salt TiF₄:8NH₄F, when exposed to air, gradually oxidises to TiO₄F₂:3NH₄F.

Titanium, haloid compounds of. Ti combines with the halogens directly to form compounds TiX4; these tetrahalides are generally prepared by passing the halogen as gas over a red-hot mixture of TiO₂ and C. TiI₄ is best prepared by heating TiCl₄ in HI gas, and TiBr₄ is very conveniently prepared by heating TiOl, in HBr gas; TiF, is prepared by dissolving TiO₂xH₂O in HFAq and evaporating. The two tetrahalides TiCl, and TiI, have been gasified. These formulæ are molecular, hence the formulæ TiF₄ and TiBr₄ probably also represent the compositions of the gaseous molecules of these compounds. By the action of H, or certain other reducing agents, on TiCl4, two chlorides are obtained, $Ti_x Cl_{xx}$ and $Ti_x Cl_{xx}$. The V.D. of neither has been determined, but from the analogy of C_2Cl_a and Si_2Cl_a , the value of x in $Ti_x Cl_x$ is probably 2; from the analogy of C_2Cl_a the molecular formula of the lowest chloride is probable. of SnCl₂ and PbCl₂ the formula Tick analogy of SnCl₂ and PbCl₂ the formula TiCl₂ would be given to it. Ti₂F₆ or TiF₅ probably exists. No bromide or iodide of either type Ti₂X₆ or Ti₂X₄ has been isolated. Two bromochlorides of the types TiX, and Ti₂X, probably exist. TiCl, and TiBr, dissolve in cold water; the solutions contain HXAq and TiO2xH2O; dilution, and more quickly heating, ppts. insoluble TiO_{*}xH_{*}O from the solution of TiOl_{*}. TiF_{*} is decomposed by water, giving H_{*}TiF_{*}Aq (and ? an oxyfluoride). TiI_{*} is also decomposed by water; on heating insoluble TiO, xH₂O separates. TiOl₄ combines with many non-metallic chlorides and oxychlorides. TiF₄ combines with metallic fluorides,

forming salts M.TiF, and also some more complex salts xTiF.yMF. A few oxyhaloid compounds have been isolated, but a great number can doubtless be formed. The acid H2TiO2F4 and salts of this acid have been isolated. Ti₂Cl_s dissolves in water, and after a time TiO₂xH₂O separates; the solution is an energetic reducing agent; alkalis ppt. Ti,O,xH,O. TiCl, reacts violently with water, giving off H, and probably

forming an oxychloride.

Titanium, hydroxyl chlorides of. By the reaction of the proper quantity of 36 p.c. HOlAq with TiCl₄, the compound Ti(OH)Cl₃ was obtained by Koenig a.O. v. d. Pfordten (B. 21, 1708); using the calculated quantity of 36 p.c. HClAq, the compound Ti(OH)2Cl2 was produced; and the prolonged action of ordinary air on either of the preceding compounds resulted in the formation of Ti(OH), Cl. These compounds are regarded by K. a. O. v. d. P. as derived from Ti(OH), by replacement of OH by Cl; the compounds may be called orthotitanic chlorhydrins. The compounds are yellow or white deliquescent solids; they dissolve in water; Ti(OH)Cl, very rapidly with a hissing noise, Ti(OH)Cl, less violently, and Ti(OH), Cl with some difficulty. The aqueous solutions give pps. of TiO2.xH2O on boiling. When the compounds are heated they give (1) TiCl, and TiO, xH,O, (2) TiO, xH₂O and HCl, and (3) TiO₂,xH₂O, HCl, and H₂O. By reducing Ti(OH)Cl, by Na amalgam, in H, Ti₂O, is produced (K. a. O. v. d. P., B. 22, 2070; cf. B. 22, 1485).

Titanium, hydroxyl fluoride of. This name may be given to the compound Ti(OH)2F4; it is described as titanoxyfluorhydric acid,

under TITANIC FLUORIDE (p. 742).

Titanium, iodides of. Only one compound

has been isolated, Til.

TITANIO IODIDE TII. (Titanium tetraiodide.) Mol. w. 554.02. V.D. at 440° = 261.2
(Hautefeuille, Bl. [2] 7, 201).
Formation.—1. Vapour of I is passed over

Ti heated to redness (Weber, P. 120, 287).—2. A mixture of TiCl, vapour, I vapour, and H is passed through a tube heated to dull redness; TiI, mixed with free I, condenses on the colder part of the tube (Hautefeuille, Bl. [2] 7, 201) .-3. By decomposing TiCl, by HI.

Preparation. - Dry HI gas is passed into TiCl, which is gradually heated to its b.p. and kept at that temperature until the reaction is completed; the TiI, is freed from traces of I, which give it a violet tinge, by a few distillations

in H (H., l.c.).

Properties and Reactions .- A reddish-brown, lustrous, brittle solid; when melted and cooled, crystallises in octahedra, changing after some days to tufts of silky, prismatic crystals. Melts at 150°, and boils a little above 360°; can be distilled without decomposition; melted Til, remains liquid till cooled below 100°. Fumes strongly in the air. Dissolves easily in water; solution turns brown on exposure to air and ppts. TiO, xH_O. When vapour of TiI, is heated in air it burns to TiO, and IO, (H., l.c.).

Titanium, nitrides of. Four compounds of

Ti and N have been described; but according to

later experiments only two seem to exist.

TRITITANIUM TETRANITRIDE Ti₂N₄. Mol. w. un-known. This compound is obtained by heating

TiCl. 4NH, (v. TITANIC CHLORIDE, Combinations. No. 2, p. 740), loosely packed in a tube of hard glass, in a stream of dry NH, until the glass begins to melt, and allowing to cool in NH. (Wöhler, A. 73, 43; Friedel a. Guérin, A. Ch. [5] 8, 24). Indigo-blue powder, or golden-yellow (? rhombohedral) crystals; strongly heated in H gives Ti₂N₂ (F. a. G., l.c.).

DITITANIUM DINITRIDE Ti.N. Mol. w. un-known. Prepared by placing two porcelain boats, one containing Na and the other K, TiF, in a hard-glass tube previously filled with N, passing a stream of dry N through the tube, heating the K.TiF, to full redness while the Na is volatilised over it, and allowing to cool in N after all the Na has been volatilised; boiling with HClAq, washing, and drying (Wöhler a. Deville, A. 102, 234). Also obtained by heating K₂TiF₆ with K, KCl, NaCl, and Al in N; also by passing TiCl, vapour and H over heated Al, in N; also by passing vapour of TiCl, over heated NH, Cl (W. a. D., l.c.). F. a. G. (l.c.) obtained the same compound by strongly heating TiO, in a stream of dry NH₂ for some hours; W. (A. 73, 43) said that TiN₂ was formed by this reaction. F. a. G. also obtained Ti₂N₂ by strongly heating Ti₂N₄ in H; W. said that Ti₂N₆ was formed by this reaction. An amorphous, brown-yellow, very hard powder; S.G. 5.28 at 18° (F. a.

G., l.c.).
The nitrides of Ti are not decomposed by heating, out of air, to c. 950°-1,000°; fused with KOH, or strongly heated in steam (F. a. G., l.c.), they give off NHs; heated in Cl, TiCl, is produced (F. a. G., l.c.); heated with CuO, PbO, or HgO, the nitrides burn, and Cu, Pb, or Hg is

produced (W., A. 123, 84).

TITANIUM DINITIBLE TiN₂. This compound is formed, according to Wöhler (A. 73, 43), by heating TiO₂ in NH, for some time. Friedel s. Guerin (A. Ch. [5] 8. 24) say that the substance obtained by W. contained Ti₂O₁, and that if the heating is continued until a definite nitride is formed the compound so produced is Ti, N,

PENTATITANIUM HEXANITRIDE Ti₂N₆. Formed, according to W. (*l.c.*), by heating Ti₂N₄ to full redness in H; or, mixed with C, by heating TiO₂ in CN or HCN gas. F. a. G. (*l.c.*) say that the product of either reaction is Ti.N., in the latter reaction mixed with C.

Titanium, nitrocyanide of, v. TITANIUM

CARBONITRIDE (p. 739).

Titanium, oxides and hydrated oxides of (cf. TITANIUM OXYACIDS AND BALTS AND DERIVATIVES THEREOF, p. 746). Ti burns when heated in O, forming TiO₂; by the reaction of reducing agents on TiCl, Aq, and addition of alkalis, Ti₂O₂ is obtained; when the reduction is carried further it is probable that TiO.xH2O is formed. By the reaction of H₂O₂Aq in presence of alkali on salts of TiO₂, a pp. of TiO₂ is obtained. Oxides intermediate between TiO₂ and Ti₂O₂ probably exist. TiO₂ is a basic oxide, forming salts TiX₄, where X = NO₂, ½SO₄, &c.; many of the salts are basic. A few salts corresponding with Ti₂O₄ have been isolated; no salt corresponding with TiO and derived from an oxyacid has yet been obtained. Some of the hydrates of TiO, react as feeble acids.

TITANIUM DIOXIDE TiO. (Titanic oxide. Titanic anhydride.) Mol. w. unknown.

Occurrence.—As rutile, anatase, and brookite; rutile occurs in dimetric prisms, isomorphous with tinstone; anatase in dimetric octahedra; and brookits in trimetric octahedra.

Formation.—1. By burning Ti in air or O.-2. By ppg. TiCl, Aq by alkali, and washing, drying, and strongly heating the ppd. TiO₂xH₂O.— 8. By ppg. K₂TiF₄Aq by NH₄Aq, and strongly heating the pp. after washing and drying.—
4. Vapour of TiCl, or TiF, mixed with steam is passed through a red-hot tube. - 5. TiS2 is heated in dry CO_2 ($\tilde{T}iS_2 + 2CO_2 = TiO_2 + 2CO + 2S$; O. v.

d. Pfordten, B. 22, 2070).

Preparation.—A. Amorphous titanic oxide.—1. Pure TiCl, is added, little by little, to water; sufficient NH, Aq to neutralise the HCl produced is added, the solution is evaporated to dryness, and the residue is heated to somewhat above 700° until it ceases to lose weight. TiCl, may be obtained from rutile by mixing the finely-powdered mineral with c. half its weight of dry charcoal-powder, making into little pellets with starch-paste, drying the pellets, heating them to full redness in a covered crucible, then heating in a tube of hard glass (to get quite dry), and then passing dry Cl over the pellets while they are heated to bright redness, and a dry receiver is connected with the end of the hard glass tube. The TiCl, may be purified (from SiCl, FeCl, &c.) by repeatedly distilling, best in a stream of N (v. Merz, J. pr. 99, 161).— 2. K2TiFe is dissolved in water, NH2Aq is added, the pp. of TiO2.xH2O is thoroughly washed, dried, and heated to somewhat above 700°. (For a method of getting K₂TiF₆ from rutile v. Preparation of titanium, p. 738.) For other methods of preparing TiO, from Ti ores v. Lévy (A. Ch. [6] 25, 511), Austen a. Wilber (Am. 4, 211), Hempel (Zeit. f. anorg. Chemie, 3, 193), Jones (Fr. 9, 41, 380). B. Crystalline titanic oxide.-3. By long-continued heating to whiteness TiO₂ is said to become crystalline (v. Rammelsberg, B. 5, 1006).—4. Amorphous TiO₂. α H₂O is heated to incipient redness in HCl gas, at a pressure of three atmospheres; the crystals have the form of anatase (Hautefeuille a. Perrey, C. R. 110, 1038).-5. Crystalline TiO2 is also formed, according to Hautefeuille (A. Ch. [4] 4, 127), by passing the mixed vapours of TiF, HF, and HCl—obtained by decomposing molten K₂TiF₆ by HCl gas—through a Pt tube into which a stream of moist H is also passed; at c.860° (boiling-point of Cd) crystals of anatase are produced, and at temperatures between 860° and 1,000° brookite is formed.—6. Heating amorphous TiO, in an atmosphere of HF causes crystallisation; at an incipient red heat anatase forms, at a higher temperature brookite, and at very high temperature rutile (H., l.c.).-7. Crystals of rutile are formed by heating amorphous TiO, with microcosmic salt (Ebelmen, A. Ch. [3] 38, 34), or with borax (G. Roce, B. B. 1867. 129; Knop, A. 157, 365), in a porcelain oven. For other methods of producing crystals of TiO₂ v. Deville (C. R. 53, 161, 163), Senarmont (A. Ch. [3] 30, 129), Michel (C. R. 115, 1020), Wöhler (A. 78, 35), Daubrée (C. R. 29, 227; 39, 153), H. Bose (A. 56, 127; 68, 163).

Properties. - TiO, obtained by strongly heating TiO, xH2O forms reddish-brown lumps, more nearly resembling rutile in colour and lustre the

higher the temperature to which it is heated. The dehydration by heat of TiO, xHO is accompanied by the appearance of shades of colour from white, though grey and greenish, to black, according to Wagner (B. 21, 960); these colours are similar to those shown by anatase, brookite, and rutile. Nilson a. Pettersson (Z. P. C. 1, 38 note) prepared TiO₂ as a white powder with faintest yellow tinge, by decomposing TiCl₄ by NH, Aq, evaporating, and heating to whiteness. When melted by the O-H flame the colour is blue to black. S.G. TiO₂ increases as the oxide is heated; heated to c. 700° S.G. is c. 8.95 (Karsten, S. 65, 894), after being very strongly heated S.G. rises to 4.25 (Ebelmen, J. 4, 15; 12, 14). S.G. of anatase is 8.75 to 3.9 (Breithaupt, J. 2, 730; Damour, J. 10, 666); S.G. of brookite is 4 to 4.2 (Rammelsberg, J. 2, 730; Beck, J. 3, 704); S.G. of rutile is c. 4.3 (Scheerer, P. 65, 296; Müller, J. 5, 847). After fusion in the O-H flame and cooling S.G. is 4.1, according to Hautefeuille (A. Ch. [4] 40, 140). S.H. 0° to 100° = 1785, 0°-211° = 1791, 0°-301° = 1843, 0°-440° = 1919° (Nilson a. Pettersson, Z. P. C. 1, 27; TiO, prepared by decomposing TiCl, by NH,Aq, evaporating to dryness, and heating to white heat). TiO2 crystallises in dimetric prisms (rutile), dimetric octahedra (anatase), and trimetric octahedra (brookite); it is isotrimorphous with SnO₂ (Wunder, J. pr. [2] 2, 206). TiO₂ is somewhat hygroscopic, even after prolonged and intense heating (Thorpe, C. J. 47, 125). It is insoluble in water, and in all acids except conc. H₂SO₄; when powdered TiO₂ is heated with conc. H₂SO₄ until the excess of acid is removed, the solid thus produced dissolves in water. TiO₂ melts in the O-H flame. According to Moissan (C. R. 115, 1034), when heated in an electric furnace-to c. 2500° TiO, forms black crystals of TiO.

Reactions.—1. According to Ebelmen (A. Ch. [3] 20, 394), TiO₂ is reduced to Ti₂O₃ by heating to redness in hydrogen; but O. v. d. Pfordten (A. 237, 201) says the product has the composition Ti,O12.—2. A mixture of TiO2 and magnesium powder heated to redness in H gives MgTiO₂, and a brown powder which is probably TiO (Winkler, B. 23, 2657).—3. A mixture of TiO₂ and carbon strongly heated in chlorine gives TiCl.-4. By long-continued heating in dry ammonia Ti₂N₂ is formed (Friedel a. Guérin, A. Ch. [5] 8, 24).—5. The compound TiCl₄.PCl₅ is obtained by heating an intimate mixture of TiO₂ and phosphorus pentachloride, in the ratio TiO₂:3PCl, in a retort until the POCl, formed is driven off (Weber, P. 132, 452).-6. Heated to redness in carbon tetrachloride vapour, TiCl, is formed (Watts a. Bell, C. J. 33, 443). Demarcay (C. R. 104, 111) says that the first product is TiOCl, —7. TiCl, is formed by heating TiO₂ to redness in a mixture of equal volumes of chlorine and carbon monoxide (W. a. B., Lc.).— 8. Ti₂N₂ mixed with C is formed by heating TiO₂ in cyanogen or in hydrogen cyanide (Friedel a. Guérin, A. Ch. [5] 8, 24).—9. Heating in a mixture of hydrogen sulphide and carbon disulphide forms Ti₂S₂ or TiS₂, according to the temperature (Thorpe, C. J. 47, 491).—10. When TiO, is fused with sodium or potassium carbonate the weight of CO₂ expelled corresponds with the formation of M₂TiO₂; on treating the fused mass

with water MOHAq isformed, and an acid Na or K titanate which is insoluble in water (v. Hermann, J. pr. 38, 92).—11. Fusion with potassium hydrogen sulphate forms a substance soluble in water; by evaporating the fused mass with conc. H₂SO₄, and then treating with water, the double salt Ti(SO₄)₂.K₂SO₄.3H₂O is obtained (Warren, P. 102, 449; Glatzel, B. 9, 1833; Hermann, l.c.).—12. Heating powdered TiO₂ with conc. sulphuric acid produces a substance that dissolves wholly in water (?forming Ti(SO₄)₂.xH₂O or TiO.SO₄; v. Merz, J. pr. 99, 157). For reactions of TiO₂.xH₂O v. infra, hydrated titanic oxide; and v. also titanic acid under Titanium oxyacides and salts and derivatives thereof, p. 746).

HYDRATED TITANIC OXIDE. Hydrates of TiO, are obtained by gradually adding TiCl, or TiBr, to cold NH,Aq; by fusing TiO₂ with KHSO₄, dissolving in water, diluting, and boiling for some time; by adding water to TiCl₄ and boiling; by fusing TiO₂ with K₂CO₂, washing the fused mass with small quantities of water till all KOH is removed, adding a little conc. HClAq, filtering cold, and allowing the liquid to stand, when it gradually deposits gelatinous $TiO_2xH_2O_6$ (O.v. d. Pfordten, B. 17, 727); by decomposing $Ti(OEt)_4$ by H_2O (Demarçay, G. B. 80, 51); and by other reactions. The isolation of a great many definite hydrates of TiO₂ has been announced from time to time. By decomposing TiOl, POl, (v. TITANIC CHLORIDE, Combinations, No. 6, p. 740) by the action of moist air, and drying over H2SO4, Tüttscheff (A. 141, 111) said that the dihydrate TiO_{2.2}H₂O was formed, and by drying this at 110°-120° he said that the *monohydrate* TiO_{2.}H₂O was obtained. According to Carnelley a. Walker (C. J. 53, 66, 81), dehydration of TiO₂.xH₂O (obtained by adding TiCl, to cold NH, Aq), air-dried for 14 days, proceeds continuously from 15° to 710° whereat the whole of the water is removed. A very large number of hydrates probably exists, and these pass one into the other as temperature rises without any of them remaining unchanged through more than a few degrees.

Two classes of hydrates of TiO₂ exist; one obtained by ppg. TiCl₂ by NH₂Aq, or warming TiO₂ with conc. H₂SO₄, dissolving in water, diluting, and ppg. by alkali; the other obtained by dissolving hydrates of the former class in dilute acid, and boiling. Hydrates of the former class dissolve easily in dilute acids, while those of the latter class are insoluble in dilute acids (for more details v. TITANIUM OXXGIDS, p. 746).

Hydrated TiO₂, obtained by ppn. by alkali in the cold, dissolves in dilute acids, forming salts TiX₄, where X = NO₃, ½SO₄, &c.; not many salts of this form have been isolated, and most of those that are known are basic salts. TiO₂xH₂O dissolves in molten H₂PO₄; on cooling crystals of TiO₂·P₂O₄(-TiP₂O₅) are obtained (Hautefeuille a. Margottet, C. R. 102, 1017). By heatfiold, and the solution after a few days, and evaporating it in vacuo, Koenig a. O. v. d. Pfordten obtained crystals approximating to Ti(OH)₂Cl₂, but differing from Ti(OH)₂Cl₂ prepared by the interaction of TiCl₄ and HClAq (B. 22, 1485; cf. Titanium hydroxyle chlorides, p. 743).

TITANIUM SESQUINORIDE Ti.O. (Titanous

oxide.) Mol. w. unknown. By digesting a solution of TiO, xH2O in HClAq with Cu at 40° to 50°. Fuchs (A. 56, 319) obtained a violet solution which gave a brownish black pp. with NH,Aq; the pp. quickly reacted with water, giving off H and forming TiO2xH2O. Von Kobell (P. 62, 599) obtained similar reactions by using reduced Ag in place of Cu. Ebelmen (J. pr. 42, 73) obtained a nearly black pp. by adding alkali to Ti₂Cl₈Aq. According to Ebelmen (J. pr. 42, 76) Ti2O, can be prepared by heating TiO, to redness in perfectly dry H. Ti₂O₃ thus prepared is described as a black solid; unacted on by HNO₃ or HClAq; oxidised to TiO2 only by heating to a very high temperature; soluble in H.SO.Aq, forming a violet solution. According to O. v. d. Pfordten (A. 237, 201), the product of reducing TiO₂ in H is not Ti₂O₃ but has the composition Ti,O12. Friedel a. Guérin (A. Ch. [5] 8, 38) obtained Ti2O2, but not free from other compounds, by passing TiCl, vapour mixed with H over TiO, at a red heat. They describe Ti,O, as a copper-red, metal-like, microscopically crystalline powder; not acted on by boiling HNO, Aq, but oxidised to TiO, xH,O by boiling H,SO, Aq; soluble in HFAq or warm aqua regia; giving off NH, and H with hot alkali solutions; oxidised to TiO₂ by heating to redness in air. Koenig a. O. v. d. Pfordten (B. 22, 2070) failed to obtain Ti_2O_3 by the method of F. a. G.

When Ti is dissolved in hot HClAq, in an atmosphere of H, the solution contains Ti₂Cl₄ (Rammelsberg, J. pr. 99, 176); a solution of TiO₂:xH₂O in HClAq reduced by zinc also contains Ti₂Cl₄ (K. a. O. v. d. P., l.c.); in these solutions alkalis give black pps., probably Ti₂O₂:xH₂O; the pps. soon change in contact with H₂O to TiO₂:xH₂O, H being given off. According to O. v. d. P. (A. 237, 201), a solution of K₂TiF₄ in a little HClAq gives a pp. of Ti₂O₂:xH₂O on

treatment with Na-amalgam.

Sabatier a. Senderens (C. R. 114, 1429; 115, 236) say that Ti₂O₃ is oxidised to TiO₂ by heating

to c. 500° in NO, or to c. 300° in NO2.

TITANIUM MONOXIDE TiO. This oxide has not been isolated with certainty. According to Moissan (C. R. 115, 1034) black prisms, which he took to be TiO, are formed by heating TiO, to c. 2500° in an electric furnace, and at a higher temperature the (?) TiO melts and then volatilises. By heating a mixture of TiO, and Mg powder to redness, in H, Winkler (B. 23, 2657) obtained a brown powder which probably contained some TiO, mixed with MgTiO,. cording to Berthier (A. Ch. [2] 54, 874) TiO, lost 6 p.c. O when heated strongly in a carbon crucible, 13 to 16 p.c. O when heated with 12 p.c. C, and 20 p.c. O when heated with 24 p.c. C (TiO₂ loses 20 p.c. O in becoming TiO); the product may have contained TiO, or it may have been a mixture of TiO, TiO, and Ti.

Hydrafed thanium monoxide TiO.H.O = TiO.H.c. o.v.d. Pfordten (A. 237, 201) obtained a black pp., said by him to be TiO.H.g. by the continued action of Na-amalgam on a solution of K.TiF. in a fair amount of HOlAq (if there is little HOlAq, Ti.O., is ppd.); the solution became green, then colourless, and then the black pp. was formed. The black pp. formed by NH.Aq, alkali carbonates, or (NH.).SAq, in solutions of TiOl. (q. v., p. 741) is TiO.H.g. according

to O. v. d. P. (l.c.). With Na, HPO, Aq, and NaO, H,O, Aq, TiOl, is said to give bluish-black and greenish-black pps., which may be salts of

TiO (O. v. d. P., l.c.).

TITANIUM PEROXIDE. TiO, (Titanium trioxide or superoxide.) In 1882 (Ätti dei Lincei, 1882. 1) Piccini dropped H₂O₂Aq into solution of TiO₂.xH₂O; from the quantity of H₂O₂ that reacted he concluded that an oxide TiO, was produced. In the same year (B. 15, 2599) Weller obtained a yellow pp., which reacted with HClAq, giving off Cl, by the interaction of H₂O₂Aq and freshly ppd. TiO₂xH₂O or a solution of TiO₂xH₂O. In 1888, Classen (B. 21, 870) added TiCl, drop by drop, to dilute alcohol, then added to this solution a very large excess of H2O,Aq, and then KOHAq, NH,Aq, or (NH₄)₂CO₂Aq; in each case he obtained a yellow liquid from which a yellow pp. separated after some time; after syphoning off the liquid, washing the pp. by decantation, and drying it on a tile, he obtained a yellow solid approximating to the composition TiO_{8.3}H₂O. In 1889, Levy (C. R. 108, 294) approximately determined the composition of the pp. obtained by adding H,O,Aq to TiO, xH2O in H2SO,Aq, by dropping in H.O.Aq of known concentration, and determining the quantity of H₂O₂ used in the reaction; L. concluded that the results could be accounted for by supposing the yellow pp. to be TiO_s, TiO_s·H₂O₂, or Ti₂O_s·H₂O₂, but that probably the pp. was TiO_s. In 1893 Bailey a. Dawson (Studies from the Phys. and Chem. Laboratories of the Owens College, vol. i. p. 216) obtained yellow to orange solids, agreeing fairly in composition with the formula TiO₂. B. a. D. added (1) H₂O₂Aq, (2) freshly ppd. BaO₂, to TiO₂xH₂O in dilute H₂SO₄Aq, diluted the deep-red liquids thus obtained with twice their volumes of alcohol, added conc. alcoholic solution of KOH, washed the pps. with alcohol (to remove H2O and H.O.), and then with ether, and, in some cases, dried in the air. According to B. a. D., TiO. dissolves in water, forming a deep-red liquid; this solution decolourises KMnO, Aq; TiO, dissolves in HClAq, giving off Cl. By allowing ppd. TiO, to stand for some time, B. a. D. say that a modification is formed insoluble in water.

Oxides of titanium intermediate between

TiO, AND Ti₂O₃.

I. By heating TiO₂ with HOl gas in a reducing atmosphere, Deville (C. R. 53, 168) obtained a blue, crystalline solid, to which he gave the formula Ti₂O₃. A similar solid (? same composition) was obtained by Friedel a. Guérin (A. Ch. [5] 8, 44) by passing H and HCl over TiO, strongly heated in a porcelain tube. This blue solid is said to decompose water, in presence of strong

bases, giving off H and forming TiO,.

II. By very strongly heating TiO, in H,
O. v. d. Pfordten (A. 237, 228) obtained a dark indigo-blue solid, to which he gave the formula Ti,O₁₂; according to Ebelmen (J. pr. 42, 76), Ti2O, is formed by this reaction. Heated in air

the compound burns to TiO2.

III. By adding alcoholic solution of KOH to a solution of TiO.xH.O in dilute H.SO.Aq, after treating the solution with BaO, Piccini (Atti dei Lincei, 1882. 1) obtained yellow pps. with compositions varying from Ti₂O₃₁ to Ti₂O₃.

Titanium oxyacids and salts and derivatives thereof. The hydrates of TiO, react as feeble acids, besides reacting with the stronger acids as feeble bases. The salts wherein Ti forms part of the acidic radicles are generally obtained by fusing TiO2 with salts of the metals which are to be converted into titanates. Only a very few titanates are soluble in water; several dissolve in HClAq, but on diluting and boiling most, if

not all, of the TiO_{2.x}H₂O is ppd.
Titanic acids. Hydrates of TiO₂ are obtained by various reactions (v. Hydrated titanic oxide, p. 745). By drying under different conditions solids are obtained approximately corresponding with the formula $x \text{TiO}_2 y \text{H}_2 \text{O}$, where x = 1, 2, or 3, and y = 1, 2, 3, 4, or 5. (For a list of the hydrates obtained by different experimenters v. Tüttscheff, A. 141, 111.) There seems to be a series of hydrates of TiO₂, all of which may be classed together under the name titanic acids, none of them being stable through more than a few degrees of temperature (v. Carnelley a. Walker, C. J. 58, 66, 81); the relations of composition of these acids may be expressed by such a general equation as $n \text{TiH}_1 O_4 - m H_2 O = \text{Ti}_n H_{4n-2m} O_{4n-m}$.

The titanic acids belong to two classes: those which are formed by decomposing TiCl, or TiBr, by dilute cold alkali solutions, or by warming TiO₂ with conc. H₂SO₄, dissolving the product in water, and ppg. by dilute cold alkali solutions; and those which are formed by decomposing TiCl, or TiBr, by water and boiling, or by dissolving titanic acids of the former class in dilute soid and boiling, or by fusing TiO, with KHSO4, dissolving in much water and boiling. The acids of the former class are generally called titanic acids or ortho-titanic acids, and those of the latter class meta-titanic acids. Orthotitanic acids dissolve easily in dilute acids; metatitanic acids are insol. dilute acids; by prolonged heating with conc. H2SO4 compounds are produced which dissolve on adding water. Metaacids are also formed by heating ortho-acids to c. 100°. Experiments made by Merz (J. pr. 99, 166) show that the meta-acids lose water, on heating, more easily than the ortho- acids.

ORTHOTITANIC ACIDS. According to Wagner (B. 21, 960), a clear solution of an orthotitanic acid is obtained by adding water, a drop at a time, to TiCl, with constant shaking; HCl and TiCl, are given off (by the heat of the reaction), a solid is produced, then a greenish-yellow liquid, and finally a clear solution. By adding 1 pt. TiCl, to c. 6 to 7 pts. water, Thorpe (C. J. 47, 120) obtained an opalescent liquid which became clear after standing for 40 hours, and then contained orthotitanic acid; when this solution was heated ppn. of metatitanic acids began at c. 87°, and at c. 90° most of the Ti was ppd. A solution of TiBr, in water forms a perfectly clear liquid containing orthoacid (T., L.c., p. 126). The white flocculent solid obtained by adding TiCl, to dilute HClAq, ppg. by NH, Aq when cold, washing with cold water, and drying in the air is TiO₂.2H₂O (-TiO₄H₄), according to Merz (J. pr. 99, 166); this composition is also assigned to the solid formed by keeping TiCl, PCl, under a bell-jar, with water and CaO, for some days, and then drying over H.SO, (Tüttscheff, A. 141, 111). When TiO, H. is dried in vacuo (Tüttscheff, Lc.), or over H.SO.

(Mers, l.c.), the product is said to be TiO₂·H₂O (- H₂TiO₂ = TiO.O₂H₂). The composition 2TiO₂·H₂O (- H₂Ti₂O₃ - Ti₂O₄·O₂·H₂) is given to the product dried at 100° (Merz, l.c.; Demoly, C. R. 1849. 325), dried at 140° (Tüttscheff, l.c.), or dried in vacuo over H₂SO₄ (H. Rose, A. 52, 268); after drying at 100° or upwards, meta-acids are probably produced.

Orthotitanic acids are insoluble in water or alcohol; but dissolve easily in dilute acids; said also to dissolve in alkali carbonate solutions. The solutions in acids probably contain salts of the type TiX_4 , X = Cl, NO_3 , $\frac{1}{2}SO_4$, &c.; but compounds of the form $Ti(OH)_xX_y$, where X is a monovalent acidic radicle, and x is not greater than 3, may be formed, similar to the compounds Ti(OH), Cl. Ti(OH)2Cl2, and Ti(OH)Cl3, obtained by O. v. d. Pfordten by the interaction of HClAq and TiCl₄ (B. 21, 1708; v. TITANIUM HYDROXYL CHLORIDES, p. 753). By adding a little conc. cold HClAq to orthotitanic acid, allowing to stand for some days, pouring off the clear liquid from undissolved acid, and evaporating in vacuo, Koenig a. O. v. d. P. (B. 22, 1485) obtained a white solid containing Ti and Cl in the ratio 1:1.47; they regarded this as consisting chiefly either of TiCl2(OH)2-but different in some properties from TiCl₂(OH)₂ obtained from TiCl,—or chiefly of TiCl(OH). HCl. White pps. are obtained by adding H.PO.Aq, H.AsO.Aq, or H.C.O.Aq to solutions of ortho-titanic acids in dilute HClAq, HNO.Aq, or H₂SO₄Aq, after making nearly neutral by NH₃Aq. When acid solutions of orthotitanic acids are diluted considerably and boiled for some time the whole of the Ti is ppd. as meta-acids. Ortho- acids are also partly changed to metaacids by keeping under water for a long time (Wagner, B. 21, 960), or by washing with hot water. When an orthotitanic acid is heated strongly TiO2 is produced, with vivid incandescence; the change from a meta- acid to TiO2 is not accompanied by incandescence. According to Wagner (l.c.), the dehydration of moist ortho-acids by gently warming is accompanied by changes of colour from white to grey, green, and black.

METATITANIC ACIDS. Formed, as white powders, by dissolving ortho-acids in dilute acid and boiling for some time; also by fusing TiO, with KHSO, and boiling with much water; also by adding water to TiCl, or TiBr, and boiling; also by oxidising Ti by HNO, S.G. 1.25 (Weber, P. 120, 287). According to Weber (l.c.), freshly prepared dilute TiCl, Aq is not rendered turbid by HClAq, HNO, Aq, or H₂SO, Aq, but these acids at once ppt. metatitanic acids when added to dilute TiCl, Aq that has been boiled even for a few seconds. The pp. obtained by boiling a diluted solution of an orthotitanic acid in dilute H₂SO, is said to have the composition TiO₂2H₂O (=TiO₂H₁), when dried over H₂SO, the composition STiO₂.4H₂O (=Ti₂O₁B₂=Ti₂O₂O₂H₃) when dried at 120°, and the composition STiO₂.2H₂O (=Ti₂O₂H₄=Ti₃O₂O₄H₃) when dried at 140° (Tüttscheff, A. 141, 111); Merz (J. pr. 99, 166) gives the composition TiO₂H₄O (=TiO₂H₂=TiOO₂H₂) to the air-dried pp., and the composition 2TiO₂H₂O (=Ti₂O₃H₂ = TiO₂O₄H₂) to the pp. dried over H₂SO₄, or at 40°.

Metatitanic acids are white powders, insoluble

in water or dilute acids; after heating with conc. H₂SO₄ the products dissolve in water; when strongly heated TiO₂ is formed without incandescence.

COLLOIDAL TITANIC ACIDS. Graham (T. 1861. 213) obtained a gelatinous, probably orthotitanic, acid, insoluble in water, by dialysing a solution of $\text{TiO}_{x}x\text{H}_{z}\text{O}$ in HClAq; by dissolving this gelatinous acid in such a quantity of cold dilute HClAq that not more than 1 p.c. titanic acid was present in the solution, and dialysing for several days, G. (C. J. 17, 325) obtained a dilute aqueous solution of colloidal titanic acid. Colloidal, insoluble, titanic acids have also been prepared by Knop (A. 123, 351), Rose (G. A. 73, 76 [1823]), and O. v. d. Pfordten (B. 17, 727).

TITANATES. These salts have not been thoroughly investigated. Those which have been best examined are either derivatives of H₂TiO₂ or H₃TiO₄, or are basic salts of the type xMO.TiO₂, where x>1. Some acid salts MO.xTiO₂, where x>1, are also known. The older investigations were made chiefly by H. Rose (P. 61, 507) and by Hautefeuille (A. Ch. [2] 4, 129).

Barium titanates. An acid salt 2BaO.8TiO₂ (=2BaTiO₂.TiO₂ or Ba₂TiO₂.2TiO₂) was obtained by Bourgeois (C. R. 103, 141) in lustrous microscopic crystals, by heating to full redness a mixture of equivalent parts of TiO₂ and BaCO₂ with excess of BaCl₂, and washing with very dilute HClAq.

Calcium titanates. The normal salt CaTiO_s occurs native as perowskite. The same salt was formed by heating to bright redness a mixture of equivalent parts of TiO₂ and CaCO₂ with excess of CaCl₂, and washing with very dilute HClAq (Bourgeois, *l.c.*).

Iron titanates. The mineral titaniferous iron or ilmenite is more or less pure $xFe_2O_2.yTiO_2$. By fusing a mixture of 2 pts. TiO₂ and 5 pts. FeF₃ with a large excess of NaCl, washing with water, and then with very dilute acid, Hautefeuille (C. R. 59, 733) obtained dark purple-violet, lustrous crystals of ferrous titanate Fe₂TiO₄ (=2FeO.TiO₂); but according to Koenig a. O. v. d. Pfordten (B. 22, 1485) the salt produced is ferric titanate Fe₄(TiO₄)₈ (=2Fe₂O₈.8TiO₂). K. a. O. v. d. P. say that the salt dissolves in water after warming with conc. H₂SO₄, forming a green solution containing Fe₂(SO₄)₈ and titanic acid.

Magnesium titanates. The normal salt MgTiO₁(= MgO.TiO₂) was obtained (by Hautefuille, l.c.) by heating to whiteness, for a short time in a closed crucible, a mixture of 1 pt. TiO₂ and 10 pts. MgCl₂ with a little NH₄Cl, washing with extremely dilute acetic acid, and then with water; lustrous, six-sided (probably trimetricy crystals, S.G. 3-91. The same salt is formed, according to Winkler (B. 23, 2657), by heating a mixture of TiO₂ and Mg powder. By heating a mixture of 2 pts. TiO₂, 1 pt. MgO, and 40 pts. MgCl₂, and washing with very dilute acetic acid, the salt Mg₂TiO₄ (= 2MgO.TiO₂) is said to be formed in hard, lustrous, regular octahedra, S.G. 3-42 (H.).

Potassium titanates. When TiO, is fused with excess of K₂CO, the quantity of CO, given off corresponds with the formation of the

normal salt K.TiO.; after fusion two layers are obtained, the lower of which is said to consist of K₂TiO₃, while the upper contains the undecomposed K₂CO₃. K₂TiO₃ is described as a yellowish, fibrous, easily fused solid (H. Rose, P. 61, 507). Water resolves it (according to Bose) into an insoluble acid salt, and a soluble basic salt; but according to Hermann (J. pr. 38, 92) no trace of titanic acid goes into solution in water, but all remains in the insoluble acid salt.

Sodium titanates. The normal salt Na₂TiO₂ was obtained by H. Rose (l.c.) similarly to the K salt; water resolves it into an insoluble acid salt and NaOHAq. By heating Na₂WO₄, and in some cases also WO₂, with mixtures of TiO₂ and Na. CO, previously fused, Cormimbouf (C. R. 115, 823) obtained three acid salts:

(1) 2Na₂O.8TiO₂ (= 2Na₂TiO₂.TiO₂, or Na₄TiO₂.2TiO₂), (2) Na₂O.2TiO₂(Na₂TiO₂.TiO₂, or Na₂Ti₂O₃), (3) Na₂O.8TiO₂ (= Na₂TiO₃.2TiO₂, or Na₂Ti₂O₃).

Strontium titanates. An acid salt

2SrO.3TiO₂ (= 2SrTiO₃.TiO₂ or Sr₂TiO₄.2TiO₂) was obtained, in pale-greenish yellow cubes, S.G. 5·1, by Bourgeois (C. R. 103, 141), by heating equivalent parts of TiO₂ and SrCO₃ with excess of SrCl₂, and washing with very dilute HClAq.

Zinc titanates. The normal salt ZnTiO, was obtained by Lévy (O. R. 107, 421), by heating to redness a mixture of 2 pts. TiO, 8 pts. ZnSO, and 3 pts. K₂SO₄, and washing with dilute HClAq; pale-violet, silky needles, S.G. 3·17, scarcely acted on by boiling conc. acids or conc. KOHAq. By varying the proportion of TiO2, ZnSO,, and K,SO, used, L. (i.c.) obtained the normal salt Zn,TiO,, which may also be regarded as a basic salt ZnTiOs.ZnO; the basic salt $8ZnO.2TiO_2$ (= $2ZnTiO_3$, ZnO); and the acid salt $4ZnO.5TiO_2$ (= $4ZnTiO_3$, TiO_3).

FLUOTITANATES; Titanifluorides, v. under TITANIC FLUORIDE, Combinations, No. 2

(p. 742)

FLÚOROXYTITANATES; v. Titanoxyfluorides, under Titanic fluoride (p. 742).

MOLYBDOTITANATES. A few compounds of the form TiO₂.12MoO₂.2M₂O. xaq, where M=NH₄ and K, are described by Péchard (C. R. 117, 788). By shaking with ether a solution of the NH, salt, acidified by HClAq, and allowing the ethereal liquid to evaporate, P. (l.c.) obtained molybdotitanic acid TiO. 12MoO. 22aq, in golden yellow octahedra, melting at c. 60°, very soluble in water.

OXALOTITANATES. By dissolving

TiO2.xH2O in hot KHC2O4Aq and cooling, Péchard (C. R. 116, 1513) obtained triclinic crystals of 2KHC₂O₄.TiO₂.H₂O. By treating a solution of this salt with BaCl₂Aq, crystals of Ba(HC₂O₄)₂TiO₂H₂O were obtained; and by decomposing this by an equivalent quantity of dilute H₂SO₄Aq, filtering, and evaporating in vacuo, long needles of oxalotitanic acid 2H.C.O., TiO., 2H.O were obtained SILICOTITANATES; v. this heading,

p. 464.

Titanium, oxychlorides of. Several oxychlorides are known. (1) TiOCl. According to Demarcay (C. R. 104, 111), this is the first product of heating TiO, and CCl, in a sealed tube; it is described as a yellow crystalline solid. Heated with CCl, it gives TiOl, and COCl, (2) TiOCl. Obtained by passing a mixture of TiCl, vapour and dry H over TiO, in a tube heated white hot; Ti,O, and Ti,Ol, are also produced in the reaction (Friedel a. Guérin, Bl. [2] 22, 481). Brown, orthorhombic leaflets; heated in air burns to TiO, and TiOl,. (3) Ti₄O₂Ol₄. Formed by passing TiCl₄ vapour through a redhot tube containing fragments of porcelain (Troost a. Hautefeuille, C. R. 73, 563). (4) By allowing TiCl, to deliquesce in air, and then evaporating over H₂SO₄ and CaO, Merz (Bl. [2] 7, 401) obtained a solid approximating in composition to Ti₄O₆Cl₄.16H₂O. In connection with oxychlorides cf. TITANIUM HYDROXYL CHLOR-IDES (p. 743).

Titanium, oxyfluoride of. By decomposing by water the crystals formed by dissolving TiO_xxH_2O in HFAq and evaporating. Berzelius (P.4,1) obtained a white solid, of which he said it was an insoluble, 'so to say, basic,' com-

pound.

The titanoxyfluorides TiO,F2.2MF may be regarded as compounds of the oxyfluoride TiO.F.

(v. TITANIC FLUORIDE, p. 742).

Titanium, oxythiochloride of. The compound TiCl, OSO, Cl formed by the reaction of TiCl, with SO2.OH.Cl may be regarded as TiO2SCl4 (v. TITANIC CHLORIDE, Reactions, No. 10, p. 740).

Titanium, salts of. Not many salts have been prepared by replacing H of oxyacids by Ti; most of those that have been isolated are basic salts derived from the oxide TiO2. One salt, Ti₂(SO₄)_s, corresponding with the oxide Ti₂O_s has been isolated. For the individual salts v. Nitrates, vol. iii. p. 517; Phosphates, this vol. p. 112, TiP₂O₇, prepared by dissolving TiO₂xH₂O in molten H₂PO₄ (Hautefeuille a. Margottet, C. R. 102, 1017) should be added; and Sulphates, this vol. p. 580.

Titanium, sulphides of. Three compounds of Ti and S have been isolated: TiS, Tis, and TiS, corresponding with the three oxides TiO,

Ti₂O₃, and TiO.

TITANIUM DISULPHIDE TiS₂. (Titanic sulphide.) Mol. w. unknown. H. Rose (P. 8, 177) said that this compound was obtained by passing vapour of CS₂ over very strongly heated TiO₂; but O. v. d. Pfordten (B. 17, 727) and Thorpe (C. J. 47, 491) have found that it is not possible by this method to obtain TiS, free from Ti oxides. TiS, is prepared by passing perfectly dry H2S into TiCl, kept somewhat below its b.p., and sending the mixed vapours through a glass tube heated to incipient redness; TiS₂ deposits in the tube, and HCl is given off (Ebelmen, A. Ch. [8] 20, 285; confirmed by O. v. d. Pfordten, B. 17, 727). According to O. v.d. P. (A. 234, 257), the H₂S should be passed through CrCl₂Aq, to remove traces of O, and dried by means of P₂O₄; and the tube should be filled with H₂S before the mixed vapours are passed into it.

TiS, forms large, brass-yellow, lustrous,

metal-like scales.

TiS, is decomposed to Ti,S, and S by heating in hydrogen or nitrogen; it is not changed by heating in hydrogen in presence of excess of hydrogen sulphide; heated to redness in carbon dioxide it is completely oxidised to TiO₂ (O. v. d. P., A. 234, 257). Ebelmen (l.c.) gives the following reactions: with dry chlorine gives TiOl₄ and S₂Ol₂; heated in air, TiS₂ is burnt to

TiO, and SO,; exposure to moist air decomposes TiS2 gradually, with evolution of H2S; heated to redness with steam gives TiO2.xH2O, H2S, and H; nitric acid produces TiO2.xH2O, S, and NO; aqua regia oxidises it to TiO, xH,O and H₂SO₄Aq; not acted on by hydrochloric acid (H. Rose said that TiS₂ dissolved slowly in HClAq giving off H₂S); insoluble in potassium sulphide solution; digestion with potash solution produces K titanate and KHSAq (Rose).

TITANIUM SESQUISULPHIDE Ti2S. Mol. w. unknown. Thorpe (C. J. 47, 491) obtained this compound by passing the vapours of moist H2S and CS2 over powdered TiO2 heated to very bright redness in a porcelain tube; dry H₂S and CS₂ had no action on TiO₂. T. describes Ti₂S₃ as a greenish-black powder; he did not obtain it quite free from TiS2, as by heating at a lower temperature than full redness in H2S and CS2 it is slowly changed to TiS₂. O. v. d. Pfordten (A. 234, 257) obtained Ti₂S₂ by heating TiS₂ to full redness in dry H or N (S being set free); he describes it as a grey, metal-like solid; insoluble in NaOHAq; soluble in HNO, Aq or conc. H2SO. forming green solutions.

TITANIUM MONOSULPHIDE TiS. Mol. w. unknown. Formed by very strongly heating TiS₂ in perfectly dry H quite free from O (T., *l.c.*; O. v. d. P., *l.c.*). A black powder (T., *l.c.*); forms dark-red crystals (O. v. d. P., I.c.). Slowly acted on by HNO, Aq or aqua regia; insoluble in NaOHAq.

Titanium, sulphochloride of, v. TITANIUM THIOCHLORIDE, infra.

Titanium, thiochloride of. The pp. formed by passing pure dry H2S into hot TiCl, is probably a thiochloride, according to O. v. d. Pfordten (A. 234, 257). M. M. P. M.

TITANIUM GROUP OF ELEMENTS. four elements, titanium, sirconium, cerium, and thorium form, with carbon, the even-series family of Group IV. in the periodic classification of the elements. There is yet an element to be discovered between Ce and Th; this unknown element will come in series 10 of Group IV. The titanium family of elements is closely allied to the tin family-Ge, Sn, Pb-which, with Si, form the odd-series members of Group IV. Carbon is the first even-series member, and silicon the first odd-series member, of the group; these two elements are more like one another than they are like the rest of the group.

For the properties and relations of C and Si v. Carbon group of elements, vol. i. p. 682; and for the tin family v. TIN GROUP OF ELEMENTS,

this vol. p. 735.

The presence of an element, before unknown, in a Cornish mineral was recognised by Gregor in 1791, and by Klaproth a few years later; the element—called titanium by Klaproth—was isolated by Berzelius in 1824. Klaproth, in 1789, recognised the presence of a new earth in zirconite from Ceylon; the metal zirconium was isolated by Berzelius in 1824. In 1803 the presence of a new earth in a Swedish mineral was announced by Klaproth, who gave the name of cerium to the metal of the earth he had discovered; Mosander isolated the element in 1826. Berzelius discovered a new earth, in a Norwegian mineral, in 1828, and isolated the metal of the earth, which he called thorium, in the same

None of the four elements Ti, Zr, Ce, or Th occurs native; the compounds of these elements are comparatively rare, especially those of Ce and Th.

The most frequently found compounds are the oxides MO_n generally in combination with SiO_2 and with alkaline earths. The metals are obtained by reducing their double fluorides by Al, K, or Na; also in some cases by reducing the chlorides by K or Na; Ce has also been isolated by electrolysing molten CeCl. The table on the following page presents some of the physical and chemical properties of the elements.

General formulæ and characters of com-Ceneral jornales and characters of compounds.—Oxides.—MO₃, probably for all (? Th₂O₃); MO₂, for all; M₂O₃, when M=Ti or Ce; perhaps TiO. Hydrates of MO₂ exist.

Sulphides.—MS₂, when M=Ti or Th, ? Zr; M₂S₃, when M=Ti or Ce; TiS.

Haloid compounds.-MX, when X=Br, Cl, F or I (only CeF, prepared); MX, when M = Ti, and X = Cl, also when M = Ce and X = Br, Cl, F or I; TiCl, Salts.—MR, R = NO, $\frac{1}{2}SO_4$, $\frac{1}{3}PO_4$, &c.; most are basic when M = Ti or Zr, but when M = Ce many are normal, and when M=Th most are normal; several salts Ce_2R_6 are known, and a few Ti_2R_6 ; the Ce_2R_6 salts are more stable than the CeR_4 salts. Salts containing M in the acidic radicle .- M_2TiO_2 and others of general form $xMO.yTiO_2$; M₂ZrO₃, M₂ZrO₄, and others of general form $xMO.yZrO_2$. No cerates or thorates isolated. Salts X₂MF₈ known when M = Ti, Zr, or Th; 3CeF.3KF known. Acids.—H₂TiO₂, H₂TiO₃, H₄TiO₄, H₂TiF₈; various hydrates $xZrO.yH_2O$ act as feeble acids; H,ThO, and H,Th,O, are perhaps feeble acids, but it is said that ThO, does not decompose alkali carbonates when fused therewith.

The oxides MO₂ are basic, and also feebly acidic when M is Ti or Zr; they interact with some acids to form corresponding salts, and, when M = Ti or Zr, with alkalis to form salts wherein M forms part of the acidic radicle; no salts of oxyacids with Ce or Th in the negative radicles have yet been isolated. Several hydrates of MO₂ exist, they easily pass one into the other with small changes of temperature; some of these hydrates react as feeble acids towards strong bases, forming salts of the types X12MO, $X_{14}MO_{4}$, and, generally, $xX_{12}O.yMO_{2}$; these salts are not referable to such definite forms as the stannates or silicates are, but they resemble these classes of salts fairly closely.

The salts which are derived from the oxides MO2, by the interactions of these oxides with acids, are of the form MX_4 , where $X = NO_4$, $\frac{1}{3}SO_4$, $\frac{1}{3}PO_4$, &c.; when M = Ti most of the salts that are known are basic; when M = Zr, many are basic, but a fair number of normal salts is also known; when M = Ce or Th, most of the salts that have been isolated are normal. Little is known of the oxides MO_1 ; as they are obtained by adding H_2O_2Aq and an alkali to solutions of salts MX4, they are probably super-oxides; the formula Th₂O, that is assigned to the oxide of Th formed in this way cannot be accepted as final. The oxides Ti,O, and Ce,O, are basic; the most definite and stable salts of

_	TITANIUM.	Zirconium.	ORRIUM.	THORIUM.				
Atomic weights.	48	90	139-9	282				
	Molecular weights unknown. Mol. weights of one or more compounds of each element, except Ce, have been determined by Avogadro's law; spec. heats determined directly.							
Melting-points.			c. 800°	Has not been fused				
Spec. gravities (approx.)	8.6	4.2	6.7	11.0				
Spec. heats. Atomic weight.	·13	-066	·0448	•0276				
Spec. gravities (approx.)	13.4	21.4	21·1	21.1				
Physical pro- perties.	Obtained only as a powder; black, lus- trous, apparently amorphous. The powder has not been fused.	A black amorphous powder; resembling lampblack, but much heavier. Also obtained as a very lustrous, hard, brittle solid, resembling Sb. Said to melt above m.p. of Si.	Steel-grey, very lustrous metal; malleable, and ductile. Melts much more easily than any other member of the family.	Dark-grey, lustrons powder; also obtained in micros ropic octahedral crystals. Has not been fused.				
Occurrence and Preparation.	The four metals occur chiefly as silicates, Ti and Zr in moderate quantiti and the others only in a few rare minerals. The metals are prepared reducing the chlorides, or alkali salts of the form K ₂ (or Na ₂)MF ₂ , by heating with K or Na; Ce also prepared by electrolysing molten CeCl.							
Chemical properties.			Burns when heated in air, or O, to CeO ₂ Decomposes cold water, giving off H. Combines directly with Cl, Br, I vapour, S vapour, and P vapour. Dissolves easily in HClAq, HNO ₂ Aq, H ₂ SO ₄ Aq, or HFAq. Forms two classes of salts, Ce ₂ X ₈ and CeX ₄ , X=NO ₉ , \$SO ₄ , are more stable than CeX ₄ , the latter are easily reduced to Ce ₂ X ₈ . No cerates, where Ce forms part of negative radicle, have been isolated. Salts of form M ₂ CeF ₈ not isolated. CeO ₂ , probably a superoxide, seems to exist.					

Ce are derived from this oxide. Several of the haloid compounds MX, have been gasified without decomposition, viz. TiCl4, TiI4, ZrCl4, and ThCl. The haloid compounds MX, form many double compounds with other halides; some of these are certainly best regarded as salts of hypothetical acids containing M and X in their acidic radicles; the salts of this class R.MF. are characteristic of the elements of Group IV.; all the elements of the group, except C and Ce, give these salts R_2MF_6 , Ce is said to form $K_3Ce_2F_{11}(=2CeF_4.3KF)$, but further examination will most probably lead to the isolation of salts R.CeF. Ti and Ce form chlorides MX, or more probably M₂X₆ from the analogy of C₂Cl₆ and Si₂Cl₆; Ce also forms Ce₂Br₆, Ce₂F₆, and Ce₂I₆, indeed the only haloid compound of Ce belonging to the form MX, that has yet been isolated is CeF4. The sulphides have not been thoroughly examined; Ti is the only member of Group IV. which is known to form three sulphides corresponding with the oxides, TiS, Ti₂S₂, and TiS₂; one sulphide of Zr is known, probably ZrS₂; ThS₂ is the only sulphide of Th that has been isolated; and the only known sulphide of Ce is Ca,S3. So far as investigation has gone, the sulphides of Ti, Zr, Ce, and Th are basic.

A comparison of the titanium family with the tin family, which comprises the odd-series members of Group IV. (C and Si being omitted), shows that Ti, Zr, Ce, and Th are more metallic, on the whole, than Ge, Sn, and Pb. Th is certainly the most markedly positive element of Group IV., and Ti is at least not more negative than Ge; Zr is more metallic than Ge, and Ce more metallic than tin. The following formulæ show that, so far as composition goes, there is about an equal similarity between C and Si and the Ti elements, as between C and Si and the

tin elements:

there is between the two families of Group VI. Considering that the titanium and the tin elements belong to even and odd series, respectively, of Group IV., which group comes midway in the general periodic scheme of classification, we should expect the titanium (even-series) family to be rather more metallic than the tin (odd-series) family, but at the same time we should expect these two families to be very like one another; this is exactly what a study of the two families shows to be the case.

In connection with this article v. TIN GROUP OF ELEMENTS, this vol. p. 735, and CARBON GROUP OF ELEMENTS, vol. i. p. 682).

M. M. P. M. TITANOFLUORHYDRIC ACID H.TiF. v. Titanifluorhydric acid, under TITANIC FLUORIDE, Combinations, No. 1, p. 742.

TITANOFLUORIDES M2TiF6 v. Titanifluorides, under Titanic fluoride, Combinations, No. 2, p. 742.

TITANOXYFLUORHYDRIC ACID H2TiO2F4 v. TITANIC FLUORIDE, p. 742.

TITANOXYFLUORIDES M2TiO2F4 v. TITANIC FLUORIDE, p. 742.

TOLALLYL SULPHIDE C28H20S2. [174°]. 350°-360° i.V.). A product of distillation of benzyl sulphide (Märcker, A. 136, 75; Forst, A. 178, 370; Baumann a. Klett, B. 24, 8318). Plates (from ether).

TOLANE v. DI-PHENYL-ACETYLENE.

Tolane dibromide v. DI-BROMO-PHENYL-ETHYL-

Tolane chlorides v. Chloro-Phenyl-Ethane and Chloro-PHENYL-ETHYLENE.

TOLENE v. TOLU BALSAM.

v-TOLENYL-AMIDINE C,H10N, C_sH₄Me.C(NH)(NH₂). [102°]. Formed by the action of alcoholic NH₃ on the hydrochloride of

Oa	rides		Sulphides	3	(Thlorides	, &c.
CO ? C ₂ O, ? SiO	CO ₃ SiO ₂	CS ?SiS	? C ₂ S ₃	CS ₂ SiS ₂	?SiCl ₂	C ₂ Cl, Si ₂ Cl,	CCl. SiCl.
? TiO Ti ₂ O ₃ Ce ₂ O ₃	TiO ₂ TiO ₃ ZrO ₂ ZrO ₃ CeO ₂ CeO ₃ ThO ₂ ? ThO ₃	TiS	Ti ₂ S ₃ Ce ₂ S ₃	TiS ₂ ZrS ₂ (?)	TiCl ₂	Ti ₂ Cl _e Ce ₂ Cl _e	TiCl, ZrCl, CeF, ThCl,
GeO SnO ? Sn ₂ O ₃ PbO Pb ₂ O ₃	GeO ₂ SnO ₂ PbO ₂	GeS SnS PbS	? ? Sn ₂ S,	GeS ₂ SnS ₂ ?PbS ₂	?GeCl ₂ SnCl ₂ PbCl ₂		GeCl ₄ SnCl ₄ PbCl ₄ (?)

A comparison of the odd-series families of | p-tolenyl-imido-ethyl-ether (Glock, B. 21, 2653). the different groups with the even-series families (omitting series 2 and 3) shows that, speaking broadly, the members of the even-series families are more metallic in their chemical properties than the members of the odd-series families, and that the general difference between even and odd families becomes more marked in the higher than in the lower groups (v. Table in vol. iii. p. 811). For instance, the even family of Group VI.—Cr, Mo, W, and U—is decidedly more metallic than the odd family of the same group-S, Se, and Te; but although the even family of Group II.—Ca, Sr and Ba—is more metallic than the odd family of the same group -Mg, Zn, Od and Hg-there is not nearly so great a difference between these two families as

Pearly plates, sol. alcohol and ether.

Reactions.—1. CO.Cl, in toluene and NaOH give (C.H., C(NH).NH).CO and C.H., C.N.C(C.H.) N [over 800°] (Pinner, B.

25, 1425).—2. Oxalacetic ether and NaOH form C,H,.C(NH).NH.CO.CH,.CO.CO,Et [190°] and $C_7H_7.0 \stackrel{\text{N.C(GO,H)}}{\sim} OH [252^\circ] \text{ (Pinner, } B. 25,$ 1422).

Salts.—B'HCl saq. [218°]. — B'₂H₄PtCl₂. [225°].—B'HNO₂. [133°]. Needles, v. sol. water (Kirschnick, A. 265, 167).—B'HNO₂ 2aq. [95°].—B'.H.SO. 2aq. [240°], v. sol. water. o-TOLÉNYL-AMIDOXIM C₈H₁₀N₂O

C,H,C(NOH).NH, [149.5°]. Formed by heat-

ing o-toluic nitrile with hydroxylamine hydroshloride, alcohol, and Na₂CO₃ (Schubart, B. 22, 2438). Needles (from hot water), v. sol. alcohol and ether.

Ethyl ether C,H,C(NOEt).NH, [140°].

Prisms, v. sol. alcohol.

Bensoyl derivative C,H,.C(NOBz).NH2. [145°]. Needles, sol. alcohol. Conc. H₂SO₄ forms $C_1H_1.0 < NO > OPh [80°]$

o-Toluyl derivative C,H,.C(NO.CO.C,H,).NH₂. [118°]. Converted at 180° into C,H, $C \leqslant \frac{N.0}{N} \geqslant C.C,H$, [59°] (Stieglitz, B. 22, 3156). Sodium diazobenzene sul-C,H,O(NH2).O,H, phonate forms [110°]

p-Tolenyl-amidoxim C_eH₁₀N₂O i.e. C.H. Me.C(NOH).NH2. Toluamidoxim. Formed by combination of toluic nitrile with hydroxylamine (Schubart, B. 19, 1487). White plates, sol. alcohol, ether and hot water. Yields NaA', a hygroscopic mass, and HA'HCl [187°],

crystallising in prisms.

Reactions.-1. Benzene sulphonic chloride forms $C,H,C \leqslant_{N}^{N,O} \geqslant SO.C_{\bullet}H_{s}$ [89°] (Pinner, B. 24, 4173).-2. Aceto-acetic ether gives rise to $C,H_1,C \leqslant_N^{N,O} > C.CH_2Ac$ [97°] (Schubart, B. 22, 2438).—8. Acetic aldehyde forms the compound $C_7H_7.C < NO > CHMe [127.5°].-4.$ Hot HOAc forms C,H,.C NO. NO. C.C,H, [135°].-5. COCl. produces C,H,.C NO [220°].-6. ClCO₂Et yields C,H,.C(NH2):NO.CO2Et [180°].-7. Phenyl C,H,.C(NOH).NH.CO.NHPh cyanate gives C,H,.C(NOH).NH.CO.NHPh [155°].—8. Phenyl thio-carbimide reacts forming C,H,.C(NOH).NH.CS.NHPh.—9. Potassium cyanate acting on the hydrocloride gives rise to C,H,.C(NOH).NH.CO.NH, [170°]. - 10. Ac,O and NaOAc acting on the hydrochloride form NH(C(NH).C,H,), [153°] (Glock, B. 21, 2657).— 11. Succinic anhydride forms the compound $C_rH_r.O < \stackrel{N.O}{\sim} C.CH_r.CH_r.CO_2H$ [188·5°].—12.

Boiling Ac_2O forms $C_7H_7.0 < N.O > CMe$ [80°]. 18. CS₂ and alcohol form, on long boiling, C,H,C(NSH)NH.CS.SHH,N.C(NSH).O,H, (Crayen, B. 24, 390).—14. CS₂ and alcoholic

potash yield C,H,CONNSCS [165°].

Methyl ether C.H., C(NOMe).NH₂. [85°]. Ethyl ether EtA'. [64°]. Needles. Converted by HBr and NaNO, into C.H., CBr: NOEt, en oil decomposed at 155° (Calulation C.C.). an oil, decomposed at 155° (Schubart, B. 22, 2434), while hydrogen chloride and NaNO, form C,H,.CCl:NOEt (200°).

Bensoyl derivative C,H, J(NOBz).NH2. [178°]. Converted by heat into

 $C_{r}H_{r}.C \leqslant_{N}^{N.O} \geqslant CPh.$

References .- NITRO- and OXX-TOLUAMIDOXIM. p-ŤOLENYL-IMIDO-ETHYL ETHER

C,H, C(NH).OEt. The hydrochloride B'HCl [161°] formed by the action of dry HCl on p-toluic nitrile dissolved in other-alcohol forms prisms, yielding B'₂H,PtCl₆ 2aq, converted at 200° into p-toluic amide, by NH₂ into p-tolenylamidine, by aniline into di-phenyl-tolenyl-amidine [168°], and by Ac₂O into the compound C_cH₄Me.C(NH).OAc [147°] (Glock, B. 21, 2650). The free base is liquid and changes on keeping into a polymeride [260°].

TOLENYL-PHENYLENE-DIAMINE

 $C_0H_0 < NH > C.C_7H_7$. [268°]. Formed by reduction of tolyl-o-nitro-aniline and also from ptoluyl chloride and o-phenylene-diamine in benzene (Hübner a. Hanemann, A. 210, 328). Prisms, sl. sol. water, sol. alcohol.

TOLENYL-PHTHALAMIDONE C18H16N2O i.e.

tolenyl-benzenyl-amidine o-carboxylic acid (Bistrzycki, B. 25, 1984).

TOLENYL-XYLYLENE-DIAMINE C15H16N i.e. [1:4] $C_6H_4Me < N > C_6H_2Me_2$ 6:1:8 [217°]. Formed by reducing the p-toluyl derivative of nitro-xylidine (Brückner, A. 205, 125; Hübner, A. 210, 333). Long crystals (from dilute alcohol).—B'HCl.—B'HNO₂.—B'₂H₂SO₄.

TOLIDINE v. DI-AMIDO-DITOLYL.

TOLIL v. DI-TOLYL-DIRETONE.
TOLIL-BENZOIN v. BENZOÏN, Reaction 9.

TOLINDOLE v. METHYL-INDOLE. TOLISATIN v. Methyl-ISATIN.

TOLU-ACET-ALDEHYDINE C₁₁H₁₄N₂ i.e. $C_1H_4(N:CHMe)_2$ or $C_1H_4 < N_{Et} > CMe$ (?) [91° uncor.]. Formed together with a small quantity of ethenyl-tolylene-diamine, by mixing tolylene o-diamine (1 mol.) with aldehyde (2 mols.)

in cold HOAc (Hinsberg, B. 20, 1588).

TOLU-AMIDOXIM v. TOLENYL-AMIDOXIM. Obtained from incisions TOLU BALSAM. in the stem of Myroxylum toluiferum growing in Columbia, South America. It contains a terpene, tolene C₁₀H₁₆ (154°-170°), benzoic and cinnamic acids, benzyl cinnamate and two resins: C16H18O4 [60°] v. sol. alcohol and ether and $C_{18}H_{29}O_5$ [above 100°], insol. alcohol and ether (Deville, A. 44, 804; Scharling, A. 97, 71; E. Kopp, A. 64, 372; Busse, B. 9, 830; Baillon, Ph. [8] 4, 385).

TOLUBENZYLAMINE v. METHYL-BENZYL-AMINE.

TOLUENE C,H, i.s. C,H,CH, Methylbensene. Retinaphtha. Phenyl-methane. Mol. V.D. 3.20 (calc. 3.29). (109°) (R. Schiff, w. 92. A. 220, 91); (111°) (Wilbrand a. Beilstein,
 A. 128, 259; Tollens a. Fittig, A. 131, 304).
 S.G. 20 8656 (Brühl); 24 8566 (Gladstone,
 C. J. 59, 290); 13 8708 (S.). C.E. (13°-109°) 001242. S.V. 118 (S.). μ_p = 1 4893 (G.). $\mu_{\rho} = 1.507$ (B.). $R_{\infty} = 50.06$. H.F.p. -8520. H.F.v. -5260 (Thomsen, Th.). H.C. 983,762 (Stohmann, J. pr. [2] 35, 41). Critical tempera-(Stohmann, J. pr. [2] 35, 41). Critical tempera-ture 321° (Pawlewski, B. 16, 2634). Occurs in cold tar (Mansfield). Obtained by dry distillation from balsam of tolu (Deville, A. Ch. [8] 3, 168; Muspratta. Hofmann, A.54, 9), from dragon's blood (Glénard a. Boudault, C. R. 19, 505), from the resin of Pinus maritima (Pelletier a. Walter,

A. Ch. [2] 67, 278), and from wood (Völckel, A. | tavson, B. 11, 2151). Oil.—(C,H_e)₂AlBr₂. S.G. 2 86, 334).

Formation.—1. By distilling vulpic acid with KOH (Möller a. Strecker, A. 113, 69).—2. By the action of sodium on a mixture of bromobenzene and MeI (Fittig a. Tollens, A. 131, 303).—3. By passing petroleum vapour over red-hot charcoal (Letny, B. 11, 1210).—4. By heating petroleum with AlCl, in presence of PbO and air (Friedel a. Crafts, Z. 1878, 1166).— 5. By passing MeCl or CH2Cl2 into benzene containing AlCl, (Friedel a. Crafts, A. Ch. [6] 1, 460; 11, 264).-6. Together with CH, by heating benzene with MeI and I (Rayman a. Preis, A. 223, 317).—7. By distilling cresol with P₂S₈ (Geuther, A. 221, 58).—8. By distilling toluic acid with baryta (Noad, A. 63, 305).

Properties.—Oil, smelling like benzene.

Reactions.—1. When passed through a redhot tube it yields benzene, styrene, naphthalene, diphenyl, anthracene, phenanthrene, and other products (Ferko, B. 20, 661; cf. Berthelot, Bl. 7, 218; Graebe, B. 7, 48).—2. Distillation over red-hot PbO gives benzene, di-phenyl-ethylene, diphenyl, phenanthrene, and anthracene (Lorenz, B. 7, 1097; Vincent, C. R. 100, 908).—3. By electrolysis of toluene to which alcohol and H2SO4 have been added benzoic aldehyde and phenose are formed (Rénard, C. R. 92, 965).-4.—When electric sparks are passed through toluene, acetylene and hydrogen are given off (Destrem, Bl. [2] 42, 267).—5. AlCl, at 200° in sealed tubes forms benzene, and m- and p-xylene (Friedel a. Crafts, C. R. 100, 692; Anschütz, A. 235, 178).-6. Oxygen passed through toluene containing AlCl, forms cresol (Friedel a. Crafts, C. R. 86, 884).—7. A mixture of toluene and ethylene passed through a red-hot tube yields benzene, styrene, and anthracene (Ferko).— 8. CH₂Cl₂ and AlCl₃ yield di-tolyl-methane (280°-290°), di-methyl-anthracene [232°], and m-, and p-xylenes (Friedel a. Crafts, Bl. [2] 41, 322).-9. Bromine acts upon toluene in the dark as readily as in diffused daylight, with production of o- and p-bromo-toluene. The addition of iodine greatly hastens the reaction, but the same products are formed. In direct sunlight the substitution takes place entirely in the side-chain, with production of benzyl bromide. But if iodine (even 2 p.c.) is present the effect of the sunshine is entirely counteracted, and the substitution takes place wholly in the nucleus (Schramm, B. 18, 606; cf. Zakrzewski, M. 8, 304).—10. Chlorine, in presence of I, acting even on boiling toluene, forms chloro-toluene and benzyl chloride (Beilstein a. Geitner, A. 231, 170; cf. Aronheim, B. 8, 1401).—11. CrO_2Cl_2 added slowly to a solution of toluene in CS_2 ppts.C,H,2CrO,Cl,,which slowly absorbs moisture, being converted into benzoic aldehyde (Etard, A. Ch. [5] 22, 223).—12. H₂SO₄ yields the o- and p-surphonic acids, converted by potash-fusion into o- and p-cresol (Wurtz, A. Ch. [4] 25, 108).—13. Bz₂O₂ forms C₁₄H₁₂ (c. 260°) isomeric with the di-phenyl-ethylenes, and yielding benzoic acid on oxidation (Lippmann, M. 7, 521).-14. HNO, forms o- and p-nitro-toluene. aldehyde and conc. H₂SO₄ form ditolylethane and C₂₈H₂₈ (350°-360°) (O. Fischer, B. 7, 1194).

Compounds (C,H,),AlCl, S.G. 21.08 (Gus-

Vor. IV.

Dihydride C_7H_{10} . (105°-108°). Formed by heating toluene with PH₄I at 350° (Baeyer, A.

155, 271; Z. [2] 4, 445).

Tetrahydride C, H₁₂. (104°). S.G. ¹³ ·797.

Occurs in the product of distillation of colophony (Rénard, A. Ch. [6] 1, 231). Oil, sol. alcohol and ether. Rapidly absorbs oxygen. In contact with water it forms crystalline C, H16O2. H₂SO₄ forms two polymerides (230°-235°), one only being oxidisable by air.

Hexahydride C,H_{II}. (97°). S.G. § ·7741. C.E. (0°-20°) ·00113. S.V. 142 (Lossen, A. 225, 109). H.C. 1,092,800 (Louguinine, C. R. 93, 275). Formed by heating toluene with conc. HIAq at 280° (Wreden, A. 187, 161; cf. Berthelot, Bl. [2] 7, 124; 26, 146). Occurs in oil of resin. Completely oxidised by a hot mixture of

HNO₃ and H₂SO₄.

References. — TRI-AMIDO-, BROMO-, BROMO10DO-, BROMO-10DO-NITRO-, BROMO-NITRO-, CHLORO-, CHLORO-IODO-, CHLORO-NITRO-, IODO-, IODO-NITRO-, NITROSO-, NITRO-, and OXY- TOLUENE.

TOLUENE ARSONIC ACID v. ARSENIC.

TOLUENE-AZIMIDO-TOLUENE v. AZIMIDO-

COMPOUNDS. TOLUENE-AZO- compounds v. Azo- com-POUNDS and DISAZO- COMPOUNDS.

TOLUENE-AZOXY- compounds v. Azoxy-COMPOUNDS.

TOLUENE CARBOXYLIC ACID v. Toluic

 $C_0H_8O_4$ Toluene dicarboxylic acid C₆H₅Me(CO₂H)₂[4:2:1]. Methyl-phthalic acid. Mol. w. 180. [152°]. Formed from (a)-amido-ptoluic nitrile by Sandmeyer's reaction, the resulting nitrile being saponified (Niementowski, M. 12, 623). V. e. sol. water and alcohol. With o-amido-phenol it forms C₅H₃Me(CO)₂N.C₅H₄OH [205°], whence boiling aqueous sodium carbonate gives rise to C₆H₃Me(CO₂H).CO.NH.C₆H₄.OH [ॅ200°].

 $Amide\ C_6H_8Me(CONH_2)_2$. [188°]. Formed

from the imide and NH,Aq.

m the imide and NH3A4.

Anhydride. [92°]. Needles.

Imide C₄H₃Me.C₂O₂NH. [196°]. Got by fusing the anhydride with urea.

Nitrile C₆H₃Me(CN)₂. [120°] (N.); [117°] (Glock, B. 21, 2663). Needles. Converted by alcohol and HCl into C,H,Cy.C(NH,Cl).OEt [199°].

Toluene dicarboxylic acid

 $\rm C_6H_3Me(CO_2H)_2$ [4:3:1]. (β)-Xylidic acid. Methylisophthalic acid. [320°–330°]. Formed by oxidation of m-xylene carboxylic acid (Jacobsen, B. 14, 2112) and of C_sH₃Me(CH₂·OH)₂ (Hjelt a. Gadd, B. 19, 868). Formed also by fusing potassium toluene m-disulphonate or the salt $C_eH_sMe(SO_2NH_2).CO_2K$ with sodium formate (Hakansson, B. 5, 1088; Remsen a. Iles, Am. 1, 119). The same acid (?) is got by oxidation of C_cH_sMe_cCO₂H_c[1:4:2] by HNO₂ (S.G. 1·12) at 150° (Claus a. Wollner, B. 18, 1858). Small crystals, sl. sol. hot water. May be sublimed. -Ag A": crystals, v. sol. hot water.

Toluene dicarboxylic acid

C_cH₂Me(CO₂H)₂ [2:4:1]. Methyl-terephthalic
acid. (a)-Xylidic acid. [c. 282°]. Formed by boiling pseudocumene with dilute HNO, (Fittig a. Laubinger, A. 151, 276). Got also by fusing C₆H₅Me(SO₂NH₂)CO₂K with sodium formate (Remsen a. Îles, Am. 1,114). Needles (by sublimation). Sl. sol. Aq. Oxidised by KMnO, to trimellitic and isophthalic acids (Krinos, B. 10, 1494).— *ZnA". S. 36 at 0°; .735 at 100°, .5 at 130° (Jacobsen, B. 10, 859).

Toluene dicarboxylic acid CoH, Me(CO, H)2. [310°-315°]. Formed by fusing potassium toluene (γ) -disulphonate with sodium formate (Senhofer, A. 164, 134). Minute needles, v. sl. sol. cold water, v. sol. alcohol and ether. May be sublimed.—BaA" 2aq.—Ag₂A": amorphous

pp.
Toluene ω o-dicarboxylic acid

CO₂H.CH₂.C₆H₄.CO₂H. [2:1]. Homophthalic acid v. o-Carboxy-phenyl-acetic acid.

Nitrile v. o-Cyano-Benzyl Cyanide.

Toluene & m-dicarboxylic acid

CO2H.CH2.C6H4.CO2H[3:1]. Formed by oxidation of m-di-ethyl-benzene (Allen a. Underwood, Bl. [2] 40, 100). Needles (from alcohol). Sublimes without melting at 200°-210°.—Ag₂A".

Toluene ω p-dicarboxylic acid CO₂H.CH₂.C₄H₂.CO₂H. [285°-288°]. S. 1 at 50°. S. (alcohol) 14·3 at 30°. Formed by the action of dilute (25 p.c.) HCl on the amide (Mellinghoff, B. 22, 3215). V. sl. sol. ether and benzene.— The same (?) acid Ag₂A": crystalline pp. formed by oxidation of C.H.PrPr [1:4]

BaA" $1\frac{1}{2}$ aq (Paterno a. Spica, B. 10, 1746). A mide CO(NH₂).CH₂.C₄H₄.CO.NH₂. [235°]. Formed from CH₂Cy.C₆H₄.Cy and H₂SO₄. Small

hard nodules.

AmicacidCO(NH₂).CH₂.C₆H₄.CO₂H. [261°]. Formed from CH₂Cy.C₆Ĥ₄.CO₂H and H₂SO₄. Sl. sol. ether, hot water and alcohol.— AgA': crystalline.

Iso-amic acid CO2H.CH2.C6H4.CO.NH2. [229°]. Formed from p-cyano-phenyl-acetic acid. Crystalline, v. sol. alcohol and hot water.

-AgA'. Crystalline.

Nitrile CH₂Cy.C₆H₄.Cy. p-Cyano-benzyl cyanide. [100°]. (above 360°). Formed from CH2Cl.CgH4Cy and KCy (Mellinghoff, B. 22, 3209). Needles, v. sl. sol. hot water, sol. alcohol and ether.

Semi-nitrile CH2Cy.C6H4.CO2H. [201°]. Formed from w-chloro-p-toluic acid by treatment with KOHAq and alcoholic KCy (Mellinghoff, B. 22, 3213). Crystalline, v. sol. alcohol

and ether.—AgA'.

Semi-nitrile CO₂H.CH₂.C₆H₄.Cy. [152°]. Formed from the nitrile and HCl at 105°. Prisms, v. sol. alcohol.—AgA'. Crystalline.

CH2Cy.C6H4.CO.NH2 Nitrile-amide [182°]. Formed from w-chloro-p-toluic amide and KCy. Plates.

Amide - nitrile CO(NH₂).CH₂.C₆H₄.Cy. [196.5°]. Formed from the nitrile and HCl at 70°. Crystalline, v. sol. alcohol.

Toluene tri-carboxylic acid

C,H,(CO,H)2.CH2.CO2H. Formed in small quantity by oxidation of s-tri-ethyl-benzene (Friedel a. Balsohn, Bl. [2] 34, 635). Needles. Sublimes before melting.—Ag₄A". Tables (from hot

TÓLUENE O-PHOSPHINIC ACID

CeH4Me.P(OH)2. Obtained by decomposing its chloride with water (Michaelis a. Paneck, A. 212,223). Oil. Monobasic acid.—CaA'₂ aq. Chloride C₂H₄Me.PCl₂. (244°). Formed

by the action of PCl, on mercury ditolyl [107°] and by warming toluene with AlCl, and PCl. Liquid.

Toluene p-phosphinic acid C₆H₄Me.P(OH)₂. [105°]. Formed in like manner. Plates, v. si. sol. water. Oxidised by HNO, to the phosphonic acid. Decomposed on heating into tolylphosphine and toluene phosphonic acid.—KA'.
—NH,A'.—BaA'2 aq.—PbA'2.—CuA'2 4aq.

Ether C.H.Me.P(OEt)2. [280°] Formed from the chloride and dry NaOEt.

Chloride $C_6H_4Me.PCl_2$. [25°]. (245°). Formed, together with the o-isomeride, from toluene, PCl_s, and AlCl_s (Michaelis a. Paneck, A. 212, 203). Formed also from Hg(C₇H₇)₂ [235°] and PCl. Needles, v. sol. ether and benzene.

References .- NITRO- and OXY-TOLCENE PHOS-PHINIC ACID.

TOLUENE PHOSPHINIC ANHYDRIDE v. Phosphino-toluene.

TOLUENE o-PHOSPHONIC ACID

 $C_6H_4Me.PO(OH)_2$. [141°]. Formed by the action of water on its chloride (Michaelis a. Paneck, A. 212, 231). Crystals, v. sol. water.—Ag₂A".

Chloride C,H,.PCl,. This is formed from C,H,.PCl, and chlorine. Yellow solid.

Toluene p-phosphonic acid $C_8H_4Me.PO(OH)_2$. [189°]. Formed from C,H,PCl, and cold water. Woolly needles. Oxidised by alkaline KMnO. to CO₂H.C₆H₄.PO(OH)₂. Bromine forms bromotoluene. — KH3A"2: néedles. — BaH2A"2 — AgHA". -Ag₂A".

Chloride C,H,.PCl₄. [42°]. Formed from C,H,PCl₂ and chlorine. Yellow mass, converted by dry SO₂ into liquid C,H,.POCl₂ (285°).

Toluene ω -phosphonic acid

C_eH_s.CH₂.PO(OH)₂. [166°]. Formed, together with (C_rH_r)₂PO(OH) [191°], from benzoic aldehyde and PH₄I (Litthauer, B. 22, 2145). Stellate groups of prisms (from HOAc).

Reference.—OXY-TOLUENE PHOSPHONIC ACID. TOLUENE ω-SELINIC ACID C,H,SeO, i.e. C₆H₅.CH₂.SeO.OH. [85°]. Formed by oxidising benzyl diselenide with HNO₃ (Jackson, A. 179, 8). Needles, sl. sol. cold water, v. sol. alcohol, nearly insol. ether. -AgA': slender crystals (from hot water).

TOLUENE SULPHAMINE v. AMIDO-TOLUENE SULPHINIC ACID.

TOLUENE o-SULPHINIC ACID C,H,SO, i.e. C.H.Me.SO.OH. [80°]. Formed by boiling C.H., N.2H. SO. C.H., with baryta-water (Limpricht, B. 20, 1241). Long needles, v. sol. ether.-BaA', 8aq. Nodules, v. sol. water.

Toluene p-sulphinic acid C.H.Me.SO.H. [85°]. Formed in like manner (Limpricht), and also from toluene p-sulphonic chloride and sodium-amalgam, zinc-dust, or Na₂SO₃ (Otto, A. 142, 92; 145, 19; B. 9, 1586; Blomstrand, B. 3, 965). Plates, v. sol. ether. Furning HNO, forms crystals [190°]. The Na salt heated with CH₃.CHCl₂ and alcohol at 150° gives a small quantity of CH₃.CHCl.SO₂.C,H₇ [48°] (Otto, J. pr. [2] 40, 519). The Na salt with ClCO₂Et gives the ether (Otto, B. 26, 308).—BaA'₂. plates.—CaA', 4aq.—ZnA', 2aq.—AgA'.

Methyl ether. Formed from the Na salt

and ClCO, Et in MeOH. Oil (Otto, J. pr. [2] 47, 166).

Ethyl ether EtA'. Oil. Oxidised by KMnO, in HOAc to toluene p-sulphonic ether

(Otto a. Rossing, B. 19, 1226).

Toluene exo-sulphinic acid CeH3.CH2.SO2H. Prepared by reducing C_sH_s.CH₂.SO₂Cl with sodium-amalgam (Otto a. Lüders, B. 13, 1288). Very unstable.—NaA': leaflets, sol. hot alcohol.

Reference.—Amido-toluene sulphinic acid. TOLUENE o-SULPHONIC ACID C, H, SO, i.e.

C_eH₄Me.SO₃H [1:2]. Formation.—1. In small quantity together with the p-isomeride, by dissolving toluene in hot fuming H₂SO₄ (Engelhardt a. Latschinoff, Z. [2] 5, 617; Anna Wolkoff, Z. [2] 6, 321).-2. Together with the m- and p- isomerides by the action of Cl.SO..OH on toluene (Claesson a. Wallin, B. 12, 1848; Noyes, Am. 8, 176).— 3. From o-toluidine by diazotisation followed by treatment with H₂SO₃ (Müller a. Wiesinger, B. 12, 1348).—4. By reducing p-bromo-toluene sulphonic acid with sodium-amalgam (Terry, A. 169, 27).—5. From (4,1,2)- nitro-toluene sulphonic acid by elimination of NO₂ (Jenssen, A.

Preparation.—By adding toluene (60 g.) to ClSO₃H (180 g.) in the cold, and pouring the

product into water (Noyes).

Properties .- Thin leaflets (containing 2aq), v.

sol. water. Deliquescent.

Salts (C. a. W.; Hubner a. Post, A. 169, 1).

Amide C,H,.SO,NH₂. [154°]. S. ·1 at 9°; S. (alcohol) 3·5 at 5°. Octahedra, sl. sol. water. Yields a benzoyl derivative [112°], which forms $C_7H_7.SO_2.NKBz$ $1\frac{1}{2}aq$ and $(C_7H_7.SO_2.NBz)_2Ba$ aq. (Wolkoff, Z. [2] 6, 57). Oxidised by KMnO, in acid solution to o-sulpho-benzoic acid and in neutral solution to C₆H₄ $< \frac{SO_2}{CO} > NH$ (Fahlberg

a. List, B. 21, 242)

Anilide C,H,SO2NHPh. [136°]

Toluide C,H,.SO,NHC,H,. [134°] Toluene m-sulphonic acid C,H4. C₆H₄Me.SO₃H [1:3]. Formation.—1. By the action of H₂SO, on m-diazo-toluene salts (Müller a. Wiesinger, B. 12, 1349).-2. A product of the action of ClSO₃H on toluene (Claesson a. Wallin, B. 12, 1848).—3. From o-bromo-toluene m-sulphonic acid and sodium-amalgam (Müller, A. 169, 47). 4. By the diazo-reaction from p-toluidine sulphonic acid and from o-toluidine sulphonic acid (Pechmann, A. 178, 202; Pagel, A. 176, 297; Nevile a. Winther, C. J. 37, 628; Klason, B. 19, 2887).

Properties .- Thin deliquescent scales (con-

taining aq), v. sol. water.

Salts.-KA'aq.-NaA'aq.-NH,A': scales, v. sol. water.—CaA', 2aq: needles, v. sol. water.— BaA', 2aq.—BaA', aq. S. 22.75 at 12° (C. a. W.); 15.5 at 15° (Vallin, B. 19, 2952).—PbA', aq.— PbA'₂2aq.—PbA'₂3aq.—MgA'₂8aq.—ZnA'₂6aq. —ZnA'₂7aq.—CdA'₂5aq.—CdA'₂6aq: crystals.

MnA', 7aq.—CuA', 4aq.—AgA': v. sol. water. Chloride C,H, SO, Cl. Oil. Amide C,H, SO, NH, [108°]. S. 4 at [108°]. S. 4 at 9° Amide C,H, SO₂NH₂. [108°]. S. 4 at 9° (C. a. W.); 265 at 14° (V.). S. (alcohol) 18 at

5° (C. a. W.); 17.4 at 14° (V.). Monoclinic tables and octahedra (from alcohol). Melts at 91° according to Noyes. Oxidised by alkaline K₃FeCy₆ to CO₂H.C₆H₄.SO₂NH₂ (Noyes a. Walker, Am. 8, 187).

Anilide C,H,.SO,NHPh. [72°]. m-Toluide C,H,.SO,NHC,H,. [103°].

Toluene p-sulphonic acid C₆H₄Me.SO₈H [1:4.] The chief product of sulphonation of boiling toluene (Deville, A. Ch. [3] 3, 172; Fittig a. Tollens, A. 131, 310; Märcker, A. 136, 85; Engelhardt a. Latschinoff, Z. [2] 5, 617; Jaworsky, Z. 1865, 221; Otto a. Grüber, A. 142, 92; 145, 10; Chrustshoff, B. 7, 1167; Fahlberg B. 12, 1048).—2. From toluene and CISO H (Claesson a. Wallin, B. 12, 1848).—3. By the action of H2SO, on p-diazo-toluene salts (M. a. W.).

Properties .- Thick leaflets or flat prisms (containing aq). Deliquescent. Melts at 92° (Norton a. Otten, Am. 10, 140). Fusion with sodium formate yields p-toluic acid (Remsen, B. 8, 1412). Potash-fusion gives p-cresol. Hydrolysis by steam begins at 150° (Armstrong a.

Miller, C. J. 45, 148).

Salts.—KA' aq. Trimetric crystals; a:b:c = 864:1:3:237.—NaA' 3aq.—NaA' 2aq.—NH₄A'.
—CaA'₂4aq.—BaA'₂.—BaA'₂aq. S. 21 at 12°.—BaA'₂3aq (Kelbe, B. 16, 621).—PbA'₂.—MgA'₂6aq.—ZnA'₂6aq.—CdA'₂6aq.—MhA'₂6aq.—MhA'₂6aq.—MhA'₂6aq.—SaA': long plates, v. sol. water.

Methylamine salt [125°].-NMe,H,A'. [78°].—NMe,HA'. [92°]. Aniline salt [223°].—o-Toluidine salt [180°]

(N. a. O.).

Chloride $C_7H_7.SO_2Cl.$ [69°]. Triclinic a:b:c=.768:1:1.114; $\alpha=97^{\circ}$ 24'; crystals; $\beta = 117^{\circ}$ 6'; $\gamma = 84^{\circ}$ 28'. Acetoxim and NaOH form C,H, SO, O.N:CMe₂ [89°] (Wege, B. 24, 3538). In ethercal solution it is reduced by sodium-amalgam to toluene sulphinic acid and the compound C₉H₁₀SO₂ [76°]. In presence of isoamyl ether $(C_5H_{11})_2^2O$ the compound $C_{12}H_{16}S_2O_4$

[36°] is formed (Otto, A. 143, 216).

Bromide C.H., SO.Br. [96°].

Iodide C.H., SO.I. [85°]. Formed by adding an alcoholic solution of I to aqueous C,H,SO,Na (Otto a. Tröger, B. 24, 479). Yellow crystalline powder, v. sol. ether, turning brown in air. Heated with water at 100° it forms C,H,S,O,C,H, Reduced silver acts in like manner. ZnEt₂ forms zinc toluene sulphinate.

Methyl ether MeA'. [c. 80°]. Ethyl ether EtA'. [32°]. S.G. 32 1.174. Thick prisms (Kraft a. Roos, B. 25, 2259).

Phenyl ether C,H, SO.OPh. [95°]. Formed by the action of NaOPh on the chloride dissolved in benzene (Otto, B. 19, 1832). Trimetric crystals; a:b:c=289:1:476. Not affected by alcoholic NH₂₁ even at 200°.

A mide C,H,SO₂NH₂. [186°]. S. ·2 at 9°. S. (alcohol) 7·4 at 5°. Leaflets. Yields C,H,SO₂NHFaq. BzCl forms C,H,SO₂NHBz [147°-150°], which gives KA', CaA', aq, BaA', AgA', and AgA'NH, and is converted by PCl, into C,H,SO₂N:CCl.C₆H₅ [100°], whence ammonium carbonate solution forms C₁,H₁,N₂SO₂ [114°] (Wolkoff, B. 5, 140). Succinyl chloride forms crystalline C,H,SO₂NC,H₂O₂, converted by NH2Aq into C,H,SO2.N2H2(C,H4O2) [180°] and

(C,H,SO₂),N,H₄(C,H,O₂), which crystallises from alcohol (Wolkoff, Z. 1870, 580).

Methylamide C,H,SO2.NHMe. [75°]. Rectangular plates, sl. sol. hot water. Yields

C,H,,SO,,NBZMe [58°].

Ethylamide C,H,,SO,NHEt. [58°].

Anilide C,H,,SO,NHPh. [103°]. BzOl forms C,H,,SO,2NBZPh [149° cor.] (Remsen a. Palmer, Am. 8, 242).

Methylanilide C,H,SO,NMePh [95°].

Monoclinic crystals (Otto, J. pr. [2] 47, 371).

Ethyl-anilide [88°]. Prisms, insol. water.
p-Toluide C.H., SO₂NHC,H., [118°].

Toluene exc-sulphonic acid C₆H₅, CH₂, SO₃H. Formed by boiling benzyl chloride with aqueous K₂SO₂ (Böhler, A. 154, 50; 221, 215; Mohr, A. 221, 216; Otto a. Lüders, B. 13, 1286). Formed also by oxidising benzyl disulphide with HNO₃ (Barbaglia, B. 5, 270, 688), and also, together with acetic acid, by heating benzyl methyl ketone with H₂SO₄ (Krekeler, B. 19, 2625). Very hygroscopic crystals. The K salt heated with KCy yields C.H..CH2.CN.

Salts.—NH,A'.—KA'aq. Trimetric prisms. -BaA', 2aq. Plates, sl. sol. water.—CaA', 2aq.-

PbA'(OH): crystals.—PbA'₂.—AgA': crystalline. Chloride C₆H₈.CH₂.SO₂Cl. [93°]. Decomposed by heat into SO₂ and benzyl chloride. A m i d e C₆H₅.CH₂.SO₂NH₂. [102°] (O. a. L.) [105°] (Pechmann, B. 6, 534). Needles, sol. water

and alcohol.

Toluene (a)-disulphonic acid CoH3Me(SOBH) [1:2:4]. Formed by sulphonation of toluene and of toluene o- or p-sulphonic acid (Hakansson, B. 1084; Senhofer, A. 164, 129; Gnehm a.
 Forrer, B. 10, 542; Claesson a. Berg, B. 13, 1170; Klason, B. 20, 354). Formed also by heating toluene p-sulphonic chloride with H2SO4 (Fahlberg, B. 12, 1052; Am. 1, 170; 2, 182). Thick liquid.—(NH₄)₂A"aq.—K₂A"aq.—BaA"aq. S. 75 at 17°. Insol. alcohol.—Ag₂A" 2aq.

 $C\ h\ l\ o\ r\ i\ d\ e\ C_7H_6(SO_2Cl)_2.$ [52°]. $A\ m\ i\ d\ e\ C_7H_6(SO_2NH_2)_2.$ [186°].

Toluene (β)-disulphonic acid C_eH_sMe(SO₃H)₂ [1:2:5]. Formed by heating toluene σ -sulphonic acid with fuming H_2SO_4 at 160° (H.), or toluene m-sulphonic acid with H_2SO_4 at 180° (Klason, B. 19, 2889; 20, 352).—K₂A" aq.—BaA" aq. S. 3.9 at 15° (K.).

Chloride $C_eH_sMe(SO_2Cl)_2$. [96°]. Amide $C_eH_sMe(SO_2NH_2)_2$. [224°].

A mide C₆H₃Me(SO₂NH₂)₂. [224°]. Toluene disulphonic acid C₆H₃Me(SO₂H)₂ [1:2:3]. Formed by the action of sodium amalgam on p-iodo-toluene disulphonic acid (Limpricht a. E. Richter, B. 18, 2179; A. 230, 326), and by heating toluene m-sulphonic acid with fuming H₂SO₄ at 180° (Klason). Slender needles, v. sol. water and alcohol.—K₂A" aq. needles, v. sol. water and acceptable of the control of the contro

A mide C.H.Me(SO2NH2)2. [214°].
Toluene o-disulphonic acid C.H.Me(SO4H)2. [1:2:6]. Formed by reduction of p-bromo-toluene disulphonic acid with sodium-amalgam (Kornatzki, A. 221, 199).--K2A".--BaA" 4aq. sol. water.

Chloride [86.5°]. Prisms (from ligroin). Amide [above 260°]. Needles, v. sol. Aq. Toluene disulphonic acid C.H.Me(SO,H), [1:3:4]. Formed from p-toluidine sulphonic acid by conversion into C.H.Me(SH).SO.H and oxidation of the product (Klason, B. 20, 356).—

K₂A" aq.—BaA" 2aq. S 15. Prisms.

Chloride. [111°]. V. sol. CHCl₂.

A mide. [235°-239°]. V. sol. water.

Toluene-s-disulphonic acid

C_oH_oMe(SO_oH)₂ [1:3:5]. Formed from o-iodo-(or bromo-) toluene disulphonic acid or o-diazotoluene disulphonic acid by long boiling with conc. HI (Limpricht a. Hasse, B. 18, 2177; A.

230, 295).—K_A'' 2½aq.—(NH₄)₂A''.

Chloride [132°]. Long prisms.

Amide [over 240°]. Small plates.

Toluene trisulphonic acid C₆H₂Me(SO₂H)₃. Formed by heating sodium toluene (a)-disulphonate with ClSO₃H at 240° (Claësson, B. 14, 307). Slender needles (containing 6aq), v. sol. water. Salts.—K₃A''' 3½aq.—Pb₃A'''₂ 8aq. water. Salts.—K₃A''' 3\frac{1}{2}aq.—Pb₃A
Ba₃A'''₂ 14aq. Crystals, v. sol. water.
Chloride C₆H_Me(SO_Cl)₃. [155]

[153°].

Amide C₆H₂Me(SO₂NH₂)₈. [above 300°]. Minute crystals, nearly insol. water.

References.—Amido-, Bromo-, Bromo-amido-, BROMO-NITRO-, CHLORO-, IODO-, IODO-AMIDO-, NITRO-AMIDO-, NITRO- and OXY- TOLUENE SUL-PHONIC ACID.

TOLUENE p-THIOSULPHONIC ACID

C.H.Me.SO2.SH. Formed by heating a solution of a salt of toluene p-sulphinic acid with sulphur (Otto, B. 15, 129; 20, 2087) or by adding a mixture of Na₂S and C,H,SO₂Na to an alcoholic solution of iodine (Otto a. Troger, B. 24, 1132). A solution of Na salt gives a white pp. of cuprous salt on adding CuSO₄.

Reactions.--1. Iddine added to an alcoholic solution of the K salt forms the three compounds (C,H,SO₂)₂S [134°], (C,H,SO₂)₂S₂ [109°], and (C,H,SO₂)₂S₃ [182°]. The compound (C,H,SO₂)₂S is also formed by the action of I on a mixture of C,H,SO,SNa and C,H,SO,Na, and crystallises from benzene in monoclinic forms a:b:c=2.829:1:3.221; $\beta=60^{\circ}7'$. The compound (C,H,SO2)2S2 is split up by boiling HOAc into (C,H,SO₂)₂S and (C,H,SO₂)₂S₃ (Otto a. Tröger, B. 24, 1126).—2. Cl.CO₂Et acting on the Na salt forms the ethyl and tolyl ethers and the compound (C₇H₇SO₂)₂S₈ [182°] (Otto a. Rossing, B. 24, 1148).

Salts.-NaA' 2aq. Trimetric KA' 2aq.—AgA' aq: small tables, sl. sol. water.

Ethyl ether C,H,SO2.SEt. Tolyl ethyl disulphoxide. Got from NaA' and EtI. Oil.

Ethylene ether C₂H₄A'₂. [77°]. Needles. Decomposed by zinc-dust into the zinc salts of toluene sulphinic acid and ethylene mercaptan

(Otto a. Heydecke, B. 25, 1478). Tolyl ether C,H,,SO₂,SC,H,. Tolyl disulphoxide. [78°]. Formed by oxidation of p-tolyl mercaptan (Märcker, A. 136, 83). Formed also from toluene p-sulphonic acid and water at 100° (Otto a. Tröger, B. 24, 480). Monoclinic prisms (from alcohol). Decomposed on saponification by alkalis into toluene p-sulphonic acid and di-tolyl disulphide [41°] (Otto a. Rössing, B. 19, 1240). Reduced by zinc and dilute H.SO, to tolyl mercaptan. Bromine forms $(C_{14}H_{14}S_2O_2)_2Br_2$ (Otto a. Grüber, A. 149, 105). Alcoholic H2S forms, on warming, toluene p-sulphinic acid, di-tolyl disulphide, and tolyl tetrasulphide.

Reference.—AMIDO-TOLUENE THIOSULPHONIC

TOLUENYL- v. Tolenyl-.

o-TOLUIC ACID CaH, Me. COaH [1:2]. Mol. w. 136. [104°]. H.C.v. 928,800. H.C.p. 929,400. H.F. 98,600 (Stohmann, J. pr. [2] 40, 133).

Formation.-1. By oxidation of o-xylene with dilute HNO₃ (Bieber a. Fittig, Z. [2] 6, 496; A. 156, 242).—2. From toluene o sulphonic acid via C,H,.CN (Ramsay a. Fittig, Z. [2] 7, 584; A. 168, 246).—3. By the action of sodiumamalgam on a mixture of ClCO2Et and o-iodotoluene and saponification of the resulting C,H,.CO,Et (Kekulé, B. 7, 1007).-4. By boiling phthalide (3 mols.) with HIAq (127°) and yellow P (2 at.), the yield being 97 p.c. (Hessert, B. 11, 238; Racine, A. 239, 72).—5. From o-toluidine vid the nitrile (Cahn, A. 240, 280).

Properties .-- Plates (containing 2aq), volatile with steam, sl. sol. cold water. Oxidised by alkaline KMnO, to phthalic acid (Weith, B. 7, 100° forms Bromine-vapour above phthalide; in the cold Br forms bromo-toluic

acid $C_6H_3MeBr.CO_2H$ [167°].

plates.—NaA' 2aq. [228°].
plates.—CaA'₂ 2aq.—BaA'₂ 2aq.
Ethyl ether Eth'. (220°).
Anido-ethyl ether
C,H,CO.O.CH.CH Efflorescent

C,H,.CO.O.CH2.CH2.NH2. Formed from o-tolyloxazoline and HClAq. Oil. Yields B'HBr [156°], B'C₆H₃N₅O₇ [188°].

Amido-propyl ether C,H,.CO.O.CHMe.CH2NH2. Formed by evaporating tolyl-methyl-oxazoline with HClAq. Yields B'HBr [140°], B'C₆H₃N₃O₇ [192°], and a platinochloride [214°].

Benzyl ether C,H,A'. (315°). S.G. 17 1.12.

Oil (Hodgkinson, C. J. Proc. No. 103, p. 167). Chloride C.H.OCl. (211° at 733 mm.). Anhydride (C.H.,CO). [c. 38°]. (above 325°). Formed from the Na salt (6 mols.) and POCis (1 mol.). Insol. water, sol. ether and benzene.

Amide C,H,.CONH₂. [140°] (Hutchinson, B. 24, 174; C. J. 57, 957; cf. Weith, B. 6, 420). On reduction in acid solution it yields o-tolyl alcohol and a small quantity of the dihydride C_7H_9 .CO.NH₂ [156°], which on boiling with water yields an acid [68°]. The dihydride is also formed by reducing the amide in alkaline solution.

Anilide C,H,CO.NHPh. [125°]. Formed from anti-phenyl-o-tolyl ketoxim and PCl, followed by water (Smith, B. 24, 4047).

Xylide C,H,.CO.NHC,H,Me₂ [1:2:4]. [165°]. Formed by heating o-tolyl xylyl ketone with hydroxylamine at 120° (Smith).

Bromo-ethylamide C,H,.CO.NH.CH,.CH,Br. [71°] (Salomon, B. 26, 1322).

Chloro-ethylamide C,H,.CO.NH.CH,CH,Cl. [73°]. White needles.

Bromo-propyl-amide[86°]. C,H,.CO.NH.CH,.CHBr.CH,. Needles (from benzene-ligroïn) (Salomon, B. 26, 1323).

Chloro-propyl-amide C,H,.CO.NH.CH,.CHMeCl. [84°]. Formed by evaporating o-tolyl-methyl-oxazoline with excess of HClAq. Needles (from ligroin).

o-Nitro-benzyl-amide C₇H,.CO.NH.CH₂.C₆H₄.NO₂. [135°]. Needles (from alcohol). May be reduced to the o-amido-

benzyl-amide [116°], which yields B'HCl [214°]

(Wolff, B. 25, 3034).

Imide (C,H,CO),NH. [148°].

Nitrile C,H,CN. (204° cor.). V.D. 4·03 (obs.). H.F. -34,800 (Berthelot a. Petit, A. Ch. [6] 17, 123). Formed by heating KCy with potassium toluene o-sulphonate or with tri-otolyl phosphate (Heim, B. 16, 1776), by heating o-tolyl thiocarbimide with finely-divided copper (Weith, B. 6, 419), by boiling o-tolyl carbamine (Weith, B. 7, 722), by heating o-iodo-toluene with AgCy at 850° (Merz a. Weith, B. 10, 751), by boiling the formyl derivative of o-toluidine with zinc-dust (the yield being 18 p.c.) (Gasiorowski a. Merz, B. 17, 74; 18, 1004), and by Sandmeyer's reaction from o-toluidine (Cahn, B. 19, 756). Liquid. H₂SO, at 60° to 70° forms C,H,C(NH).NH.CO.C,H, [103°], which is converted, by warming its acid solution, into the imide (C,H,CO)2NH (Krafft a. Kerstens, B. 25, 455). Conc. HIAq forms C₇H₇.CI₂.NH₂ [98°] (Biltz, B. 25, 2540).

Hexahydride v. METHYL-HEXAMETHYLENE

CARBOXYLIC ACID.

m-Toluic acid C₆H₄Me.CO₂H [1:3]. [110°] (Jacobsen, B. 14, 2347; Bornemann, B. 20, 1382). (263°). H.C.v. 928,500. H.C.p. 929,100. H.F. 98,900 (Stohmann, J. pr. [2] 40, 184). S. 09 at 15°; 1.66 at 100°.

Formation.—1. By oxidising bromo-xylene and reducing the resulting bromo-toluic acid with sodium-amalgam (Ahrens, Z. [2] 5, 106; Richter, B. 5, 424; Böttinger a. Ramsay, A. 168, 255).—2. By distilling uvitic acid with lime (Fittig, B. 5, 268).—3. By oxidation of m-xylene with dilute HNO₃ (Tavildaroff, B. 4, 410; Brückner, B. 9, 406; Reuter, B. 17, 2028).— 4. By saponifying the nitrile.

Preparation.—1. By boiling ω -chloro-xylene with HNO₃ (1 vol. of S.G. 1·4) and water (8 vols.) (Senff, A. 220, 247).—2. By sulphonating crude xylene, then preparing C₆H₃Me₂,SO₂NH₂, oxidising by KMnO₄, heating the resulting C₆H₄Me(SO₂NH₂).CO₂H with HCl at 230°, and distilling with steam (Jacobsen, B. 14, 2347).

Properties. — Long needles, volatile with steam, v. sol. alcohol and ether. Oxidised by chromic acid mixture to isophthalic acid.

Salts.—CaA'23aq. S. 317 at 15°; 8.2 at 100°.—BaA'22aq: trimetric plates or tables.— AgA'.

Ethyl ether EtA'. (226°). Chloride (218° at 724 mm.) (Ador a. Rilliet, B. 12, 2300).

Nitrile C,H,.CN. (209°). Formed by heating m-tolyl thiocarbimide with Cu (Weith a. Landolt, B. 8, 720), and in other ways (Buchka

a. Schachtebeck, B. 22, 841).
p-Toluic acid C₆H₄Me.CO₂H [1:4]. [179°].
(275° cor.) (Fischli, B. 12, 615). H.C.v. 926,800. H.C.p. 927,400. H.F. 100,600 (Stohmann, J. pr.

[2] 40, 184).

Formation.—1. By the action of dilute NHO₃ on cymene (Noad, P. M. [8] 32, 19; A. 63, 289) on p-xylene (Beilstein, A. 187, 302; Brückner, A. 205, 113), and on terpenes (Hirzel, Z. 1866, 205).—2. By saponification of the nitrile.—3. By heating p-bromo-toluene with ClCO. Et and sodium amalgam and saponifying the product (Wurtz, C. R. 68, 1298).—4. From p-bromo-toluene, CO₂, and sodium (Kekulé, A. 187, 184).-5. By the action of POCl₂ (40 g.) on a mixture of toluene (40 g.), HOAc (80 g.) and ZnCl₂ (80 g.) at 110° (Frey, J. pr. [2] 43, 116).— 6. By reducing bromo-p-toluic acid with sodiumamalgam (Remsen a. Morse, Am. 1, 138) .-7. From the chloride which is formed by the action of COCl2 on toluene in presence of AlCl2 (Ador a. Crafts, B. 10, 2176).—8. From the amide which is formed by the action of AlCl, on a mixture of toluene, CS2, and NH2.CO.Cl (Gattermann a. Schmidt, B. 20, 859).—9. By fusing C.H., CO.C.H., CO.2H with potash (Friedel a. Crafts, Bl. [2] 35, 508), and also by passing gaseous cyanic acid and HCl into toluene at 100° containing AlCl. (Gattermann a. Rossolymo, B. 23,

Properties.—Needles, v. sol. hot water, v. e. sol. alcohol and ether. Volatile with steam. Oxidised by CrO, or KMnO, to terephthalic acid.

Yields a di-nitro-derivative [158°].

Salts.—KA'.—BaA'22aq: needles (Buchka a. Irish, B. 20, 1764).—CaA', 3aq.—MgA', 3aq. CuA'2. -AgA'.

Methyl ether MeA'. [32°]. (217°). Ethyl ether EtA'. (228°). Oil. A mido-ethyl ether C,H,CO.O.C,H,NH2.

Yields B'HBr [167°] and B'C, H₃N₃O, [180°].

p-Amido-propyl ether C,H,CO.O.C,H,NH2. The The hydrobromide B'HBr is formed by evaporating a solution of the bromo-propyl-amide. It yields B'C, H, N, O, [186°] and B',H,PtCl,

Phenyl ether PhA'. [72°]. Chloride (225° at 720 mm.).

Amide C,H,CO.NH₂ [159°]. Formed by heating the NH₄ salt in sealed tubes at 230° (Hallemann, R. T. C. 6, 79).

Methylamide C,H,.CO.NHMe. [143°]. Formed by the action of NHMe.COCl on toluene in presence of AlCl_s (Gattermann a. Schmidt, B. 20, 120; A. 244, 51). Tables. Ethylamide C,H,.CO.NHEt. [96°].

Bromo-ethyl-amide

C,H,.CO.NH.CH,CH,Br. [129°] (Salomon, B. 26, 1325). Formed from p-toluic chloride and bromo-ethylamine. Plates. Converted by boiling water into the amido-ethyl ether.

Chloro-ethyl-amide C,H,.CO.NH.CH,.CH,Cl. [122°]. Needles (from

ligroïn).

Bromo-propyl-amide C,H,CO.NH.CH,CHBr.CH₃. Melts at 74° becomes solid, and melts again at 158° (B. 26, 1326)

Chloro-propyl-amide C,H,.CO.NH.CH,.CHMeCl. [78°].

Anilide C,H,.CO.NHPh. [145°]. Formed from toluene, phenyl cyanate, and AlCl3 (Leuckart, J. pr. [2] 41, 306). o - Nitro - anilide.

[110°]. Yellow

prisms.

p - Toluide C,H,.CO.NHC,H,. [160°]. Formed by the action of sodium amalgam and HOAc at 60° on an alcoholic solution of di-ptolyl ketoxim (Goldschmidt a. Stöcker, B. 23, 2747; 24, 2799). Needles.

Nitro-toluideC,H,CO.NHC,H,Me(NO2)

[4:1:8]. [166°].

Diphenylamide C,H,.CO.NPh. [155°]. Prisms (from alcohol) (Lellmann a. Bonhöffer, B. 20, 2118).

Xylide C,H,.CO.NHC,H3Me2. [139°]. Yields C_7H_7 .ČO.NH. $\dot{C}_8\dot{H}_2Me_2(NO_2)$ [187°] Phenyl-methyl-amide C,H,.CO.NPhMe.

[70°] (Lellmann a. Benz, B. 24, 2114).

o-Nitro-benzyl-amide

 C_7H_7 .CO.NH.CH₂. C_6H_4 .NO₂. [142°] (Wolff, B. 25, 3036).

Nitrile C.H.Me.CN. [28·5°]. (218°). Formed by distilling potassium toluene p-sulphonate with KCy (Merz, Z. [2] 4, 33), by heating C,H,.N.CS with Cu (Weith, B. 6, 421), by heating tri-p-tolyl phosphate with KCy (Heim, B. 16, 1775), and by passing C.H., NH.CHO over heated zinc-dust (G. a. M.). Obtained also from p-toluidine by Sandmeyer's reaction (Glock, B. 21, 2650). Sodium added to its boiling alcoholic solution forms a small quantity of w-amido-p-xylene, the greater part being saponified (Bamxylene, the greater part being bary H₂SO₄ gives berger a. Lodter, B. 20, 1710). H₂SO₄ gives CNH\NH.CO.C.H. [145°]. Easily converted into the imide (C,H,.CO),NH [155°] (Krafft a. Karstens, B. 25, 454). Conc. HIAq forms C₇H₇.CI₂.NH₂ [115°-120°], decomposed by fusion (Biltz, B. 25, 2539).

(252°). Formed sodium-amalgam (Einhorn a. Willstälter, B. 26, 2009). Needles, melting below 0°. Yields CaA'₂4aq, CuA'₂2aq, MeA'. (210°-220°), and an amide [158°]. Boiling NaOHAq converts the acid into a solid isomeride [47°] (254°-260°) which yields CaA'24aq and an amide [135°]. Hexahydride (Serebojakoff, B. 25, 3355).

References .- AMIDO-, BROMO-, BROMO-NITRO-, CHLORO-, NITRO-, NITRO-AMIDO-, OXY-, OXY-AMIDO-, and SULPHO- TOLUIC ACID.

a-Toluic acid is PHENYL-ACETIC ACID.

o-TOLUIC ALDEHYDE C₆H, Mc.CHO[1:2]. o-Toluylic aldehyde. (200°). Prepared by boiling ω-chloro-o-xylene with lead nitrate solution (Lauth a. Grimaux, Bl. [2] 7, 233; Rayman, Bl. [2] 27, 498). Prepared also from o-xylene by the successive action of CrO₂Cl₂ and water (Bornemann, B. 17, 1467), and by oxidising $C_6H_4Me.CH_2OH$ with chromic acid mixture (Kröber, B. 23, 1029). Oil, smelling like bitter almonds. Yields a crystalline compound with NaHSO₃. Reduced by sodium-amalgam to o-tolyl-carbinol.

Oxim C₆H₄Me.CH:NOH. [49°]. Is an antioxim.White crystals (from ether), v. sol. Yields a crystalline hydrochloride. alcohol. Ac₂O yields an acetyl derivative [56°]. AcCl acts in like manner. The acetyl derivative is decomposed by alkalis with regeneration of the oxim, no nitrile being formed (Dollfus, B. 25, 1921). The oxim is converted in ethereal solution by PCl_s into the nitrile (204°).

m-Toluic aldehyde C₆H₄Me.CHO[1:3]. (199°). S.G. ² 1.037. Formed by boiling ω-chloro-mxylene or ω-bromo-m-xylene with lead nitrate solution (Gundelach, Bl. [2] 26, 44; Müller, B. 20, 1213), and also by the action of water on C,H,CrO,Cl, obtained from m-xylene (Etard, B. 14, 848; Bornemann, B. 17, 1464). Oil, smelling like almonds. Combines with NaHSO.

Aniline forms C,H,CH:NPh. (314°).

Phenyl-hydraside C,H,Me.CH:N,HPh.
[91°]. (B.); [88°] (Rudolph, A. 248, 100).

Prisms (from ligroïn), v. sol. ether.

p-Toluic aldehyde C₆H₄Me.CHO[1:4]. (205°). Formed by distilling calcium p-toluate with calcium formate (Cannizzaro, A. 124, 254). Formed also by the action of water on the product of combination of p-xylene with CrO₂Cl₂ (Bornemann). Oil, with peppery smell. Oxidised by air to toluic acid. Alcoholic potash forms potassium toluate and tolyl-carbinol.

Reference.—Nitro- and Oxy- toluic alde-

TOLUIC CHLORIDE v. Chloride of Toluic

TOLUIDES. Compounds derived from toluidine by displacement of H in NH₂ by an acid

o-TOLUIDINE C₆H₄Me.NH₂ [1:2]. Mol. w. 107. (198°). S.G. ²⁰ ·9986 (Brühl, A. 200,189). H.C.p. 964,700. H.C.v. 963,750. H.F. 3,800 (Petit, C. R. 107, 266). μ_B 1·5895. R_∞ 57·56. S.V. 126·6. Formed from o-cresol, ZnBr₂, NH₃, and NH₄Cl at 335° (Merz a. Müller, B. 20, 547). Prepared by reduction of o-nitro-toluene. May be separated from p-toluidine by means of the acidoxalates (Bindschedler, B. 6, 448; Ihle, J. pr. [2] 14, 449; Miniati, Booth a. Cohen, S. C. I. 6, 418), by freezing (Rosenstiehl, Bl. [2] 17, 7), by means of the nitrates (Schad, B. 6, 1861), hydrochlorides, sulphates, or phosphates (Lewy, Fr. 23, 269; B. 19, 1717, 2728; cf. Wülfing, B. 19, 2132). o-Toluidine can be prepared from C₆MeHBr(NH₂)H, C₆MeHBrIRH, followed by C₆Me(NO₂)BrIBrH, and C₆Me(NH₂)BrIBrH. It can also be got from C₆Me(NH₂)BrIBrH. Via C₆MeIBrIBrH, followed by C₆MeIBrIBrH, followed by C₆MeIBrIBrH, via C₆MeIBrIBrH, 101, 21:2] is identical with C₆H₄Me(NH₅) [1:2] is identical with C₆H₆Me(NH₆), [1:6] (Wroblewsky, A. 192, 213).

C_eH₄Me(NH₂) [1:6] (Wroblewsky, A. 192, 213). Properties.—Oil. A solution in H₂SO₄H₂O is coloured orange by HNO, and blue by CrO, the solution in the latter case becoming reddishviolet on dilution. Forms a crystalline compound with liquid CO2 at 8° (Bitte, C. R. 105, 614). With PbO, and aqueous acetic acid gives a green colour, while PbO₂ and alcoholic acetic acid gives a reddish-violet colour (Lauth, C. R. 111, 975). Unlike p-toluidine, it forms a crystalline compound with alloxan (Pellizzari, C. C. 1887, 1288, 1396). If an ethereal solution of o-toluidine is shaken with very dilute bleachingpowder solution the aqueous layer becomes yellow or brown, and the ethereal layer, if decanted and shaken with dilute H2SO4, colours this reddish-violet. A solution of p-tolylene diamine hydrochloride mixed with a little otoluidine gives a green colour on addition of

FeCl₃ or K₂Cr₂O₇ (Nietzki, B. 10, 1157).

Reactions.—1. POCl₃ forms PO(NHC₂H₁)₃
[225°] (Michaelis a. Rudert, B. 26, 565), v. sol.
water, whence Br forms PO(NHC₄H₃BrMe)₃
[253°]. Another product of the action of POCl₂
on o-toluidine is POCl(NHC₄H₄Me)₂, whence water
forms HO.PO(NHC₂H₂)₂ [95°]. PSCl₂ forms
PS(NHC₂H₃)₃ [135°].—2. Sicl₄ reacts forming
SiCl₄(NHC₂H₃)₂, a white granular powder (Harden,
C. J. 51, 40).—3. On heating with malic acid it
yields the mono-toluide [178°], di-toluide [181°],
and a tolyl-imide [116°] of malic acid (Bischoff,
B. 23, 2043).—4. Chloro-citryl chloride forms
C₂H₂N₃ (CO.CH₂CO.NHC₂H₄) [214°] (Skinner

a. Ruhemann, C. J. 55, 239).—5. SO₂, alcohol, and nitrous acid give rise to the sulphazide $C_{14}H_{16}N_2SO_2$ [142°] (Limpricht, B. 20, 1241).— 6. On nitration of o-toluidine or its acetyl derivative NO₂ enters the p- and o- positions, but in presence of H₂SO₂ (10 pts.) the compound C_cH₃Me(NO₂)(NH₂) [1:4:6] [107°] is formed (Nölting a. Collin, B. 17, 265), together with the (1,2,6)-isomeride (Green a. Lawson, C.J. 59, 1018). -7. Benzoic aldehyde forms C.H.CH:NC,H, an oil (316°) (Étard, Bl. [2] 39, 530; Pictet, B. 19,1063).—8. Hydrobenzamide or warming forms the same body (310°) (Lachovitch, M. 9, 695).— 9. Toluidine heated with toluidine hydrochloride and benzoic aldehyde in sealed tubes at 120° forms C₆H₂.CH(C₆H₃Me.NH₂)₂, a crystalline powder turning blue in the air (Ullmann, *J. pr.* [2] 36, 251).—10. The methyl derivative of p-cay-ben-zoic aldehyde at 120° produces the compound C₆H₄(OMe).CH:N.C.H₇ [32°] (Steinhart, A. 241, 340).-11. Bromo-propiolic acid reacts, forming 5401.—11. Stome-proports acts teats, forming C₁,H₁₈N₂O₂ [185°] (Mabery a. Krause, B. 22, 3308).—12. αβ-Di-bromo-acrylic acid produces pale-yellow needles [115°] (M. a. K.).—13. The hydrochloride heated with MeOH at 250°-300° gives rise to di-methyl-toluidine, xylidine C₅H₃Me₂(NH₂) [4:3:1] and mesidine (Limpach, B. 21, 640).—14. Cyanogen passed into an alcoholic solution forms crystalline $C_{16}H_{18}N_{\bullet}$ which yields $B''H_{2}Cl_{2}$ and $B''2HNO_{3}$ (Bladin, Bl. [2] 41, 128).—15. On heating with S if yields C₁₄H₁₂N₂S [120°] crystallising from alcohol in yellow plates and yielding a dibromide $C_{14}H_{12}Br_2N_2S$ [190°] (Gattermann, B. 22, 425).—16. ClSO₃H in CHCl₃ forms C₁H₁NH₂SO₄H, which yields BaA'2 2aq. Both the acid and its salts reproduce o-toluidine when treated with hot water (Traube, B. 23, 1656).

Salts (Beilstein a. Kuhlberg, A. 156, 66; Rosenstiehl).—B'HClaq. S. 37.4 at 15.5°. S. (alcohol) 100. — B'2H_PtCl₈. Decomposed by boiling water (difference from p-toluidine) (De Coninck, Bl. [2] 45, 131).—B'2H_2ZnCl₄. White tables (Bibanoff, Mont. scient. [3] 4, 925).—B'_2ZnCl₂ 2aq (Lachovitch, M. 9, 513).—B'2HgCl₈. [115°]. Prepared by shaking an alcoholic solution of the base with HgCl₂ (Klein, B. 11, 743). Crystalline powder, sol. alcohol and ether.—B'2MnCl₂.—B'3H₃CuCl₂. Yellow crystalline pp. got by adding CuCl₂ to a solution of the hydrochloride (Pomey, C. R. 104, 365).—B'HBr. Trimetric prisms.—B'2HgBr₂. [104°]. Leaflets (Klein, B. 13, 835).—B'2ZnBr₂ (Leeds, J. 1882, 503).—B'HI. Trimetric prisms (Städel, B. 16, 28).—B'2HgL₂. White needles.—B'2ZnL₂—B'2HgCy₂.—B'4HgCy₂.—B'4H₂FeCy₄: crystals.—B'2H₂Cu₂(SO₃)₂. Formed by adding aqueous cupric sulphate and NaHSO₃ to a solution of the base in acetic acid (Denigès, C. R. 112, 870).—B'HGL₂(SO₃)₂ aq.—B'H₂SO₄. Crystals (Wellington a. Tollens, B. 18, 3313).—B'2H₃SO₄. S. 78 at 22° S. (89 p.c. alcohol) 1.6° at 21.5°.—B'H₃PO₄. S. 19, 1717). The salts B'2H₂PO₄ and B'3H₂PO₄ could not be prepared. Aniline sets free o-toluidine from B'H₂PO₄. p-Toluidine slso sets free o-toluidine from its phosphate.—B'HSCyCr(SCy)₃2NH₄. Red plates, v. sol. hot water (Christensen, J. pr. [2] 45, 362).—Oxalate: small plates. S. 238 at 21°. S. (84 p.e.

alcohol) 2.68 at 21°. S. (ether) .65 at 21°.-Chloroacetate B'C₂H₂ClO₂ [95°] (Bischoff, B. 21, 1257).—(β?)-naphtholate [81°] (Dyson, C. J. 48, 470).

Formyl derivative C, H, NH.CHO. [58°]. (288°). Plates (Ladenburg, B. 10, 1129; Tobias, B.15, 2446). The crystalline polymeride [211°] is v. sol. hot HOAc, nearly insol. ether and cold alcohol.

Thioformyl derivative C,H,.NH.CSH: [96°]; yellow needles; bitter taste; sol. alcohol and ether, insol. water. Formed by heating the formyl derivative with P2S3. It dissolves unaltered in cold aqueous NaOH, and is reprecipitated by an acid. By distillation in vacuo it

forms di-tolyl-formamidine $C_7H_7.NH > CH$ with

evolution of H₂S and CS₂. By heating for 7 hours at 190° under pressure a crystalline compound C₁₆H₁₆N₂S [160°] is formed with evolution of H₂S (Senier, B. 18, 2292; C. J. 47, 762).

Acetyl derivative C.H. NHAc. (Bedson a. King, C. J. 37, 753). (296°). S. ·85 at 19°. Formed from the base and HOAc (Beilstein a. Kuhlberg, A. 156, 77; Alt, A. 252, 318), or by shaking it with water and Ac2O (Hinsberg, B. 23, 2962). Converted by PCl, into the compound C,H,N:CClMe, which when carefully heated gives a base C,H,N:CMe.CH, CCl:NC,H, [53°], and when treated with excess of o-toluidine gives di-tolyl-acetamidine [69°]. Br in HOAc forms $C_6H_3MeBr(NHAc)$ [1:6:2] [157°].

Chloro-acetyl derivative C,H,NH.CO.CH2Cl. [112°]. Formed by the action of chloro-acetyl chloride on o-toluidine dissolved in benzene (Widman, J. pr. [2] 38, 299). Needles (from dilute alcohol). Converted by alcoholic potash into a pyrazine derivative

[160°].

Di-chloro-acetyl derivative C,H,NH.CO.CHCl2. Needles, volatile with steam

(Rugheimer a. Hoffmann, B. 18, 2987).

Tri-chloro-acetyl derivative C,H,NH.CO.CCl₁. [67°]. Formed from hexa-chloro-acetone and o-toluidine (Cloez, A. Ch. [6] 9, 215). Needles, sl. sol. cold alcohol.

Bromo-acetyl derivative C,H,.NH.CO.CH2Br. [118°]. Formed o-toluidine and bromo-acetyl bromide (Widman, J. pr. [2] 38, 298). Needles. Converted by alcoholic potash into $C_{1e}H_{1e}N_2O_2$ [160°].

Thio acetyl derivative.

C.H., NH.CS.CH. [68] (Wallach, B. 13, 529;
16, 147). Converted by NaOEt and EtBr into

oily C,H,N:CMe.SEt (262°).

Propionyl derivative C,H,NH.COEt. [87°]. (299° at 730 mm.). Formed from o-tolui-dine and propionic acid (Pictet a. Duparc, B. 20, 3421). Needles, sl. sol. hot water. Oxidised by KMnO, to propionyl-o-amido-benzoic acid [1170].

a. Bromo-propionyl derivative C.H., NH.CO.CHBr.CH., [131°]. Neadles, insol. ligroin (Tigerstedt, B. 25, 2920).

Bensoyl derivative C,H, NHBz. [131°] (Gudeman, B. 21, 2553); [148°] (Brückner, A. 205, 130). Long broad needles. Yields benzoylo-amido-toluio acid on oxidation by KMnO.
PCl, forms C.H.,N:CCl.C.H. (Just, B. 19, 982).
Oxalyl derivative C.O.(NH.C.H.).

[131°]. Plates (from alcohol) (Bladin, Bl. [2] 41, 130).

Phthalyl derivative C,H,N:C2O2:C6H4. [182°]. (near 360°). Needles, sol. hot HOAc (Fröhlich, B. 17, 2679).

Phenacyl derivative

C,H,NH.CH,CO.C,H, [89°]. Formed from the base in alcohol and w-bromo-acetophenone (Bischler, B. 25, 2865). Needles. Yields B'HCl and an acetyl derivative [92°].

m-Toluidine C_eH₄Me.NH₂ [1:8]. (197°) (Beilstein a. Kühlberg, A. 156, 83); (201°) (Buchka a. Schachtebeck, B. 22, 840); (205°) (Lorenz, A. 172, 180; Merz a. Müller, B. 20, 548). S.G. 25 H.C.v. 964,600. S.N. 128·1. H.C.p. 965,600. H.F. 2,900 (Petit, A. Ch. [6] 17, 155). Formed by reduction of m-nitro-toluene. Obtained also from p-toluidine by successive acetylation, nitration, saponification, diazotisation, boiling with alcohol, and reduction (Lorenz, B. 7, 448). Formed also by heating m-cresol with ZnBr₂, ammonia, and NH₄Br at 300° (M. a. M.). Prepared from benzoic aldehyde by nitration, conversion into m-nitro-benzylidene chloride and reduction of this body by zinc and HClAq in the cold (Vienne a. Steiner, Bl. [2] 35, 428; Widman, B. 13, 676; Bl. [2] 36, 216; Ehrlich, B. 15, 2011; Harz, B. 18, 3398). m-Toluidine can be shown to exist in commercial toluidine by means of the nitroso- derivative of di-methylm-toluidine, since o- and p- di-methyl-toluidines do not give nitroso- derivatives (Wurster a. Riedel, B. 12, 1796).

Properties .- Oil. Its solution in H2SO4H2O is coloured yellowish-brown by CrO, and blood-red by HNO₃. Its ethereal solution, shaken with water and a little bleaching-powder, is coloured reddish, the aqueous layer being brownish yellow. HNO, added to its solution in a mixture of H₂SO, and HOAc below 0° forms nitro-toluidine [134°] and some of its isomerides. Cyanogen passed into its alcoholic solution gives the compound (C,H,NH₂)₂C₂N₂ [200°] which forms B'₂HCl, B'₂2HNO₃, and B'₂2H₂SO₄ (Clève a. Bladin, Bl. [2] 41, 129). Yields (4,2,1)-xylidine when its hydrochloride is heated with MeOH at 200°-300° (Limpach, B. 21, 646).

Salts.-B'ĤCl. S. 96.3 at 12°. S. (94 p.c. alcohol) 61.9 at 9°.—B'₂H₂PtCl₄.—B'HNO₃. § 20.1 at 23.5°. S. (94 p.c. alcohol) 46 at 15°.—B'HBiI₃ (Kraut, A. 210, 323).—B'₂H₂SO₄. § 6.25 at 14°. S. (94 p.c. alcohol) 41 at 15°.—B'HBiI₃ (Kraut, A. 210, 323).—B'₂H₂SO₄. § 6.25 at 14°. S. (94 p.c. alcohol) 41 at 15°.—B'HBiI₃ (Kraut, A. 210, 323).—B'₂H₂SO₄. § 6.25 at 14°. S. (94 p.c. alcohol) 41 at 15°.—B'HBII₃ (Kraut, A. 210, 323).—B'₂H₂SO₄. § 6.25 at 14°. S. (94 p.c. alcohol) 41 at 15°.—B'HBII₃ (Kraut, A. 210, 323).—B'₂H₂SO₄. § 6.25 at 14°. S. (94 p.c. alcohol) 41 at 15°.—B'HBII₃ (Kraut, A. 210, 323).—B'₂H₂SO₄. § 6.25 at 14°. S. (94 p.c. alcohol) 45°. $B'_4H_4FeCy_6$ (Eisenberg, A. 205, 270).— $B'H_2C_2O_4$. Silky needles. S. 2.65 at 13°; S. (94 p.c. alcohol) 1.77 at 15°; S. (ether) 13 at 15°.—
B'₂2H₂C₂O₄. White plates, S. 1.45 at 15°; S. (94 p.c. alcohol) '96; S. (ether) '128 at 15.5°. —B'₂H₂C₂O₄ aq. Very unstable.

Formyl derivative C,H,NH.CHO. (278° at 724 mm.) (Niementowski, B. 20, 1892). Converted by long boiling into C,H,N:CH.NHC,H, [128°].

Acetyl derivative C,H,.NHAc. [65°].

(303°). S. .44 at 18°.

Di-chloro-acetyl derivative C.H.,NH.CO.CHCl. [100°]. Silky plates (Rügheimer a. Hoffmann, B. 18, 2988).

Bensoyl derivative C,H,.NHBz. (Just, B. 19, 983). [125°]

Phthalyl derivative C,H,N:C,O,:C,H,

[153°]. Needles, v. sol. HOAc (Fröhlich, B. 17, 2679).

p-Toluidine C.H.Me(NH.)[1:4]. [45°] (Städeler); [42·8°] (Mills). (198°) (M. a. H.); (206°) (Städeler, J. pr. 96, 65). S.G. 1·046. S. 35 at 11·5°. H.C.p. 958,800. H.C.v. 957,860. H.F. 9,700 (Petit, C. R. 107, 266). S.V. 128,700. Formed by reduction of p-nitrotoluene (Muspratt a. Hofmann, A. 54, 1; 66, 144; Noad, A. 63, 305: Hugo Müller, Z. 1864, 161; Sell, C. J. 16, 186; v. also o-toluidine). Formed also by heating p-cresol with ammoniacal ZnCl₂ (Buch, B. 17, 2687; Merz a. Müller, B. 20, 545) and by treating methylaniline hydrochloride at 350° (Hofmann, B. 5, 720). On sulphonating a mixture of o- and p-toluidine the o-toluidine is first attacked and the p-toluidine can then be distilled off.

Properties.—Plates (from ligroin or dilute alcohol). Its solution in H₂SO₄H₂O is turned yellow by GrO₂ and blue by HNO₃, in the latter case changing to red and finally brown (Rosenstiehl, Bl. 10, 200). Bleaching-powder does not colour its solution in water or ether. PbO₂ and HOAc diluted with water or alcohol give a bright red colour (Lauth, C. R. 111, 975). When boiled with water, crystals of an efflorescent hydrate separate on cooling (Lewy, B. 19, 2728).

Reactions.—1. Oxidised by KMnO, alkaline K₂FeCy₆, or chromic acid to C₂H₂₁N₃ or C₆H₂Me(NC₇H₇)₂(NH₂) [1:2:5:4], which may be readily reduced to C₆H₂Me(NHC₇H₇)₂(NH₂) crystallising from alcohol in colourless plates [166°], readily re-oxidised to C₂₁H₂₁N, (Barsilowsky, A. 207, 102, 118; J. R. 1887, 132; Perkin, C. J. 35, 728; 37, 546; Klinger a. Pitschke, B. 17, 2439; Green, C. J. 63, 1395). The compound $C_nH_2N_3$ orystallises from xylene in dark-red plates [227°], dissolving in acids, forming maroon-red liquids, quickly decomposing with separation of p-toluidine. Conc. H_2SO_4 dissolves $C_{21}H_{21}N_3$, forming a greenish-blue liquid which on warming becomes winered with production of p-toluidine.—2. Oxidised by CrO, and HOAc to toluene-azo-toluene C,H,N,C,H,. — 3. Nitrous acid forms C,H,N,NHC,H,. When NaNO, is added to a solution of the hydrochlorides of o- and ptoluidine, the o-toluidine is first attacked, yielding C,H,N,2,C,H,NH,2 and can thus be removed when mixed with p-toluidine (Wülfing, D. P. J. 263, 260).—4. Substituents usually enter positions 3 and 5 [Me=1] but when p-toluidine or its acetyl derivative is dissolved in H₂SO₄ (20 pts.) and nitrated the compound C₆H₃Me(NO₂)(NH₂) [1:2:4] is formed instead of the (1,3,4)- isomeride (Nölting a. Collin, B. 17, 263).—5. Bromine in conc. HClAq at 0° forms C₀H₂Br₂Me(NH₂) [5:3:1:4]; but C₀H₃BrMe(NH₂) [2:1:4] is formed in presence of conc. H₂SO₄ (Hafner, B. 22, 2538, 2902).—6. ICl forms $C_8H_2I_2Me(NH_2)$ [5:3:1:4] (Michael a. Norton, B. 11, 107).—7. Chlorine passed into p-toluidine dissolved in H₂SO, at 0° forms C_sH₂MeCl(NH₂) [1:3:4] and larger quantities of its (1,2,4)- isomeride. In HClAq (40 p.c.) the chief product is C_sH₂MoCl₂(NH₂)[1:3:5:4] (Hafner).—8. Cyanogen passed into an alcoholic solution forms C16H18N4 or (C,H,N)₂C₂N₂ crystallising in pearly scales [222°-230°] and yielding the crystalline salts B'₂2HCl, B'₂2HNO₂, B'₂H₂SO₄6aq, and B'₂H₂C₂O₄

(Hofmann, C. J. 1, 170; A. 66, 144; Clève a. Bladin, Bl. [2] 41, 125). The mother-liquor contains $C_{24}H_{23}N_s$ [182°] which yields the salts B'HCl 3aq (a red powder) and B'₂H₂PtCl₅. The compound $C_{16}H_{18}N_4$ is also formed when p-toluidine is heated with oxalimido-ether at 100° (Vorländer, B. 24, 805).—9. Cyamuric chloride forms $C_5N_8(NHC_7H_7)_8$ [288°] (Klason, J. pr. [2] 33, 294).—10. Chloro-acetic acid (h. mol.) forms C.H.NH.CH., CO.H. while shloro-(½ mol.) forms C,H,NH.CH2.CO2H, while chloroacetamide forms C,H,NH.CH, CO.NH, (P. F Meyer, B. 8, 1158). An alcoholic solution of chloro-acetic acid and HCyS yields C₁₀H₁₂N₂SO₂ [176°-182°] (Nencki, J. pr. [2] 16, 1).—11. Dichloro-acetic acid forms tolylamido-methyloxindole. - 12. Chloral reacts, with formation of CCl₂.CH(NHC,H₂)₂ [115°] (Wallach, A. 173, 278). When chloral is warmed with an alcoholic solution of p-toluidine there is also formed CCl₃.CH(OEt).NHC,H, [77°]. — 13. Aldehyde forms crystalline CH3.CH(NHC,H3)2 [c. 60°].—14. Furfuraldehyde added to a hot alcoholic solution of p-toluidine and its hydrochloride forms brown amorphous C10H22N2O2, which yields B'HCl and B'HNOs both crystallising in purple needles (Stenhouse, A. 156, 203). 15. Nascent formic aldehyde produces a base C₁₇H₁₈N₂ [134°] which is converted by nitrous acid with evolution of CO₂ into C₁₆H₁₆N₁O₃ [260°-264°] (Tollens, J. pr. [2] 36, 225).— 16. Benzoic aldehyde at 100° reacts, forming C_sH_s.CH(NHC,H_s)₂[100°] (326° i.V. at 723 mm.), which changes at 160° into an isomeride [120°-125°] and is converted by Br in CS₂ into C₁,H₁,Br₂N [160°-165°] (Schiff, Z. 1865, 400; Mazzara, J. 1880, 566; Pictet, B. 19, 1063). In the action of p-toluidine and its hydrochloride on benzoic aldehyde at 120° there is produced C₂₁H₆N₂ [178°] and C₆H₅.CH(C₆H₅Me.NH₂)₂ [186°] (c. 430°), which yield a di-acetyl derivative [218°] and a di-benzoyl derivative [196°] (Ullmann, J. pr. [2] 36, 267).—17. p-Methoxy-benzoic aldehyde forms C_cH₄(OMe).CH:(NC,H₂) [92°] (Steinhart, A. 241, 332).—18. p-Methoxy-benzophenone chloride yields the compound O.H. (OMe).OPh:NC,H. [96°] Hantzsch a. Kraft, B. 24, 3520).—19. An ethereal solution of $C_{c}H_{b}$.CCl:NSO₂Ph, obtained from benzene sulphonic anilide and PCl, forms the compound C₆H₂.C(NHC₇H₇):N.SO₂Ph, which separates in monoclinic crystals; a:b:c = ·650:1: 533; β= 69° 16', insol. water (Wallach a. Gossmann, A. 214, 216).—20. Alcohol, SO₂ and nitrous acid form the sulphazide C₁₄H₁₆N₂SO₂ [140°] (Limpricht, B. 20, 1241).—21. Heated with persulphocyanic acid it gives p-tolyl-thiobiuret (B. 17, 584).—22. On heating with sulphur it yields dehydrothiotoluidine and primuline; v. PRIMULINE.—23. Diazobenzene chloride yields C.H..N.2.NHC,H. [85°] which is identical with the product of the action of p-diazotoluene chloride on aniline (Nölting a. Binder, B. 20, 2005). 8005).—24. Bromo-propiolic acid forms di-tolyl-acetamidine C₁₆H₁₄N₂ [120°] (Mabery a. Krause, B. 22, 3307), which is also formed by the action of HOAc and PCl₂ on p-toluidine (Hofmann, Pr. 15, 55).—25. Itaconic acid in boiling aqueous solution forms C,H,.N < CH₂.QH.CO₂H CO.OH₂ [185°] (Scharfenberg, A. 254, 150).—26. Acetylcitric anhydride dissolved in chloroform forms

C.H.O(CO.H)(CO.NHC,H.)2 [189°] Klingemann, B. 22, 987).—27. Malic acid at 160° yields C.H.O(CO.NHC,H.)2 [208°] (Bischoff, B. 23, 2045).—28. The hydrochloride heated with MeOH yields C.H.Me.,(NH2) [1:3:4] (Limpach, B. 21, 640).—29. Ethylene oxide forms oxyethyl-p-toluidine (Demole, B. 7, 635).—30. SOCl2 forms thionyl-toluidine (q. v.).—31. POCl3 forms PO(NHC,H.)2 [192°], which yields PO(NHC,H.BP)3 [180°] and PO(NHC,H.2,NO.)2 [247°] (Rudert, B. 26, 569). Another product of the action of POCl3 is POCl(NHC,H.)2, whence water forms HO.PO(NHC,H.)3 [185°].

Salts.—B'HCl. S. 22.9 at 11°; S. (89 p.c.

(alcohol) 25 at 17°.—B'HAuCl,.—B',H,PtCl,. Not decomposed by boiling water (De Coninck, Bl. [2] 45, 131).—B'_PtCl_2 (Gordon, B. 3, 177; Cochin, Bl. [2] 31, 499).—B'_H_PtCy_4 (Scholz, M. 1, 905).—B'PtCl_Et_pO_5 (Saillard, Bl. [2] 18, 111). — B'₂PtCl₂M₂PO₂. — B'PtCl₂2Et₃PO₂ (Cochin, J. 1878, 315).—B'₂ZnCl₂ 3aq (Lachovitch, M. 9, 513).—B'₂H₂ZnCl₄ (Gräfinghoff, Z. 1865, 599). — B'₂H₂Cl₂. [125°]. Thick needles, sol. alcohol and ether (Klein, B. 11, 743).—B'₂NiCl₂2EtOH. Green pp. (Lippmann a. Vortmann, B. 12, 81).—B'₂CoCl₂. Blue mesdles.—B'₂H₂SnCl₈. Monoclinic crystals (Hjortdahl, J. 1882, 535).—B'₂H₂CuCl₄ (Pomey, C. R. 104, 300).—B'₂MnCl₂ (Leeds, J. 1882, 503).—B'₂UrO₂Cl₂.—B'HBr. Plates (Städel, B. 16, 28).—B'₂HgBr₂. [121°]. Plates (Klein, B. 13, 835).—B'₂ZnBr.—B'₂CdPr.—B'₃CdPr.—B'₄CdPr 28).—B'₂HgBr₂. [121°]. Plates (Klein, B. 13, 835).—B'₂ZnBr₂.—B'₂CdBr₂.—B'HI.—B'₂HgI₃. [81°].—B'HBiI₄ (Kraut, A. 210, 324).—B'₆(H₂SO₄),₈H₃ (Jörgensen, J. pr. [2] 14, 386).—B'₂ZnI₂.—B'₂CdI₂.—B'HNO₃. S. 17·7 at 23·5°; S. (89 p.c. alcohol) 42 at 20°.—B'₂AgNO₃. [101°] (Mixter, Am. 1, 239).—B'₂Cd(NO₃)₂.—B'₂H₂SO₄. S. 5·06 at 22°; S. (84 p.c. alcohol) 1·3 at 23°.—B'H₃SO₄. Crystals, converted into B'₂H₂SO₄ by water (Tollens, B. 18, 3312).—B'₄Ag₃SO₄ 2aq (Mixter).—B'H₃PO₄. Thick needles, v. sl. sol. cold water (Lewy, B. 19, 1717).—B'₄(NH₂Ph)H₄CoCy, 1₄aq (Weselsky, 19, 1717).—B'₂(NH₂Ph)H₃CoCy₆1½aq (Weselsky, J. 1869, 314).—B'₃H₃CoCy₆2aq.—B'₂HgCy₂.—B'HCr(SCy)₄2NH₃. Plates (Christensen, J. pr. [2] 45, 362).—B'₂Cu₂H₂(SO₃)₂. Yellowish-white hexagonal plates.—B'HgH₂(SO₃)₂ aq (Denigès, hexagonal plates.— B'HgH.(SU,)₂ aq (Deniges, C.R.112,870).—Oxalate B'H₂C₂O₄ gaq (Bornemann, B. 22, 2710). S. 87 at 14°; S. (84 p.c. alcohol) '48 at 22°. Insol. ether.—Chloroacetates B'C₂H₃ClO₂ [102°] (Baralis, J. 1884, 698).—B'C₂H₂Cl₂O₂. [141°]. (B.) [136°] (Duisberg, B. 18, 194).—B'C₂HCl₃O₂. [137°].—Trichloro-lactate B'C₃H₃Cl₂O₄. [135°].—Mucate B'C₄H₂O₄. [135°].—Mucate B'C₄H₂O₄. [135°].—Mucate B'C₄H₂O₄. [135°].—Mucate B'C₄H₂O₄. [135°].—Mucate B'C₄H₂O₄. [135°].—Support of the state of the stat B'₂C₈H₁₀O₈. Yellowish crystals, sol. hot water.
—Picrate B'C₆H₂N₂O₇. [169] (Smolka, M. 6, 923). S. 54 at 18.5°; S. (95 p.c. alcohol) 4.29 at 18°.—Phenate [31·1°]. Got by heating p-toluidine with phenol (Dyson, C. J. 43, 468).

p-toluidine with phenoi (Dyson, C. J. 43, 408).

Needles (from ligroin).—Compound with picramide B'C_eH_d(NO_{2)e}NH₂. Black needles with green reflex (Mertens, B. 11, 843).

Formyl derivative C,H,NK.CHO. [45°] (Hübner a. Rudolph, A. 209, 371); [52°] (Tobias, B. 15, 2446); [53°] (Bamberger a. Wulz, B. 24, 2080). Formed by boiling p-toluidine with formic acid or, together with the oxalyl derivative C₂O₂(NHO,H,₁)₂ [263°], by heating p-toluidine oxalate. Prisms. Converted by heating with P₂S₂ into C.H.,NH.CHS [174°]

(Senier, B. 18, 2294; C. J. 47, 766). Yields C,H,.NNa.CHO aq.

Acetyl derivative C,H,.NHAc. [147°] (Hübner a. Wallach, A. 154, 302; Kelbe, C. J. 44, 915). (807°) (Beilstein a. Kuhlberg, A. 156, 74). S. '089 at 22°. Dimorphous, crystallising in monoclinic forms; a:b:c=1·216:1: '788; β=106° 7', and also in trimetric crystals; a:b:c=651:1: '329 (Panebianco, G. 9, 362). Oxidised by KMnO, to acetyl-p-amido-benzoic acid (Hofmann, B. 9, 1299). Oxidised in like manner in the animal body (Jaffe a. Hilbert, H. 12, 295). PCl₃ forms a liquid product which solidifies when strongly cooled, the crystals doubtless being C,H,NH.CCl₂.CH₃. This readily splits up into HCl and C,H₂.N:CCl.CH₃. This is a colourless solid which is converted by Aq into acetyl-p-toluidine; by toluidine into di-tolyl-acetamidine. The chloro-imide C,H,N:CCl.CH₃ at 100° forms a base C₁₈H₁₈N₂Cl [72°], which may be crystallised from alcohol. It is probably C,H,N:CMe.CH₂.CCl:NC,H, It is decomposed by moist air or water with production of MeC(NC,H₁)NC₂H₂H, HOAc and MeCONHC,H₄, and by heat with production of an amorphous base C₁₈H₁₈N₂ (Wallach a. Fassbender, A. 214, 202). Nitrous acid converts acetyl-p-toluidine into C,H₂.NAc.NO [80°] (O. Fischer, B. 10, 959).

Chloro-acetyl derivative C,H,NH.CO.CH₂Cl. [162°]. Needles (Tommasi, Bl. [2] 19, 400; Meyer, B. 8, 1154).

Di-chloro-acetyl derivative C,H,NH.CO.CHCl₂. [153°] (Rügheimer a. Hoffmann, B. 18, 2980).

Tri-chloro-acetyl derivative C,H,NH.CO.CCl_s. [80°] (C.); [102°] (J.). S. (alcohol) 12·5 at 14°. Formed from p-toluidine and (CCl_s)₂CO (Cloez, A. Ch. [6] 9, 145), and also from p-toluidine and CCl_s.COCl (Judson, B. 3, 784).

Bromo-acetyl derivative C.H.NH.CO.CH.Br. [164°]. Needles (Abenius, J. pr. [2] 40, 433) converted by alcoholic potash into an azine and C.H.NH.CO.CH₂.OEt [32°] which is v. e. sol. alcohol.

Thio-acetyl derivative
C,H,NH.CS.CH₂. [131°]. Formed from the
acetyl derivative and P₂S₅ (Jacobsen a. Ney, B.
22, 906). Got also by heating tolyl-acetamidine
with CS₂ at 100°, or di-tolyl-acetamidine with
H₂S (Bernthsen a. Trompetter, B. 11, 1759;
Wallach a. Pannes, B. 13, 529). Converted by
NaOEt and EtBr into oily C,H,N:CMe.SEt
(272°) (Wallach a. Wüsten, B. 16, 147).

a-Bromo-propionyl derivative C,H,NH.CO.CHBr.CH₃. [125°] (Tigerstedt, B. 25, 2921). Plates, v. sol. CHCl₃.

Benzoyl derivative C,H,NHBz. [158°]. (232°). Formed from p-toluidine and BzCl (Jaillard, Z. 1865, 440; Kelbe, B. 8, 875; Hübner, A. 208, 310; Wallach, A. 214, 217; Gudeman, B. 21, 2553; Müller, B. 22, 2404). Needles, v. sol. alcohol. Converted by PCl, into C,H,N:CCl.C₆H₅ [52°] (Just, B. 19, 980) crystallising from ligroin in prisms.

p-Nitro-benzoyl derivative C.H.(NO₂).CO.NHC.H., [197°]. Formed from p-toluidine and p-nitro-benzoyl chloride. Yellow needles (Gattermann, B. 25, 1082). Converted by PCl₃ into C.H.(NO₂).CCl₂.NHC.H., [119°] crystallising in golden prisms.

Phenacyl derivative C,H,NH.CH₂.CO.C₆H₃. [127°]. Formed from w-bromo-acetophenone and p-toluidine in alcohol (Bischler, B. 25, 2866). Yellow prisms, yielding B'HCl and an acetyl derivative [89°] crystallising in white needles (from alcohol).

Phthalyl derivative C,H,N:C2O2:C6H4.

[205°] (Fröhlich, B. 17, 2679).

References.—Bromo-, Dibromo-1000-, Bromo-NITRO-, CHLORO-, TRI-CHLORO-NITRO-, IODO-, NITRO-, and NITROSO- TOLUIDINES.

TOLUIDINE SULPHONIC ACID v. AMIDO-FOLUENE SULPHONIC ACID and also Nitro- and IODO- TOLUIDINE SULPHONIC ACID

TOLUIDINE THIOSULPHONIC ACID v.

AMIDO-TOLUENE THIOSULPHONIC ACID.

TOLUIDO- v. TOLYL-AMIDO-

TOLUIDYL-CYANURAMIDE v. CYANIC ACIDS. TOLUÏDYL-MELAMINE v. CYANIC ACIDS.

TOLUISATIN v. DI-TOLYL-OXINDOLE. **TOLU-(αβ)-NAPHTHAZINE** $C_{17}H_{12}N_2$ $C_{e}H_{4} < CH:CH.C.N > C_{e}H_{2}Me$

[180°]. Formed by oxidation of a mixture of tolylene-odiamine (4 mol.) and (3)-naphthol (1 mol.) with 4 K₅FeCy₆. Formed also, together with the isomeride [169°], by the action of (3)-naphthoquin-

one on tolylene-o-diamine (Witt, B. 20, 577). Dissolves in conc. H₂SO₄ with violet-red colour. Forms with its isomeride a molecular compound

[142°].
Tolu- $(\alpha\beta)$ -naphthazine $C_{17}H_{12}N_2$ i.e. $C_0H_4 \underbrace{\begin{array}{c} CH:CH.CN\\ CN \end{array}}_{CN} C_0H_3Me \quad \begin{bmatrix} 4\\ 3 1 \end{bmatrix}.$ [169°].

Formed as above, and also, together with amido-benzene p-sulphonic acid, by boiling sulphobenzene-azo- (β) -naphthyl-p-tolyl-amine with dilute H_SO₄, and by boiling 'wool black' C_eH₄(SO₂H).N₂.C_eH₄(SO₄H).N₂.C₁₀H₆.NHC,H, with dilute H₂SO₄ (Witt, B. 20, 577). Yellow crystals, v. sl. sol. slcohol. Conc. H₂SO₄ forms a brownish-red solution.

TOLUNITRANILIC ACID v. NITRO-DI-OXY-

TOLUQUINONE

TOLUOIN v. DI-METHYL-BENZOÏN.

TOLUOL v. Toluene.

TOLUPHOSPHINIC ACID v. PHOSPHOTOLUIC

TOLUQUINOL v. HYDROTOLUQUINONE. TOLUQUINOLINE v. METHYL-QUINOLINE. TOLUQUINONE C, H, O, i.e.

C.H.MeO₂ [2:4:1]. [69°].

Formation.—1. By oxidation of tolylenep-diamine or of o-toluidine (Nietzki, B. 10, 832, 1934; A. 215, 158; Ladenburg, B. 10, 1128).-2. By boiling crude creosol with MnO₂ and dilute H₂SO₄ (Carstanjen, J. pr. [2] 23, 425). Separated from accompanying xyloquinone by reduction with SO2, crystallising the hydrotoluquinone from benzene, and re-oxidising by fuming HNO.-3. By oxidation of m-xylidine (Nölting a. Baumann, B. 18, 1151).

Preparation.-20 g. of o-toluidine are dissolved in 600 c.c. of water containing 160 g. of H2SO4; 20 g. of finely powdered potassium bichromate are added by degrees, with continual agitation, keeping the temperature at 10°-15°. After standing over night, 83 g. more K2Cr2O, are added; the quinone is then extracted with ether, dried over CaCl., and the ether removed by distillation. The yield is about 86 p.c. of the theoretical quantity (Schniter, B. 20, 2283).

Properties.—Golden-yellow plates smelling like quinone. Very volatile. M. sol. hot water, v. sol. alcohol and ether. Its aqueous solution is turned brown by alkalis. SO2 reduces it to hydrotoluquinone. Dilute (50 p.c.) H_2 SO_4 changes it to isotoluquinone $(\text{C}_7\text{H}_6\text{O}_2)_x$, a powder not melted at 300° (Spica, G. 12, 225). An aqueous solution of toluquinone mixed with one of hydrotoluquinone deposits steel-blue needles of the quinhydrone [52°], v. e. sol. water. o-Nitro-aniline forms [2:1:4:6] C₆H₂MeO₂·NH.C₆H₄NO₂ crystallising from alcohol in red crystals decomposing at 200° (Leicester, B. 23, 2796). Nitrop-toluidine C,H,Me(NO2).NH,[1:3:4] forms a homologous body.

Chlorimide $C_eH_uMe < {0 \atop NCI} \left[12 \atop 2\right]$. [88°] (Hirsch, B. 18, 1514). Formed by the action of a solution of bleaching-powder on the hydrochloride of amido-o-cresol. Yellow prisms (from alcohol) volatile with steam. Decomposes vio-lently at 100°-115°. By digestion with NaHSO₂ for several hours it is converted into amidocresol sulphonic acid.

Chlorimide $C_6H_3Me < \stackrel{\text{NCl}}{\dot{O}} [1_2^5]$.

Made in like manner from amido-m-cresol (Staedel, A. 259, 218). Yellow prisms (from alcohol), volatile with steam.

Oxim C₅H₅MeO(NOH)[2:1:4]. Nitroso-ocresol. [135°]. Formed by the action of nitrous acid on o-cresol (Nölting a. Kohn, B. 17, 370). Formed also by the action of hydroxylamine hydrochloride on toluquinone (Goldschmidt a. Schmid, B. 17, 2063). Needles (from water), decomposed by fusion. V. sol. alcohol and ether. Yields nitro-cresol [95°] on oxidation.—NaA'3aq.

-KA': brown needles, v. sol. water.

Oxim C₆H,MeO(NOH) [2:4:1]. Nitroso-mcresol. Formed by boiling nitroso-di-methyl-m-toluidine with dilute (10 p.c.) NaOHAq (Wurster a. Riedel, B. 12, 1799). Formed also by the action of NO. SO. H on m-cresol and water at 0° (Bertoni, G. 12, 303). Small needles (from water). Decomposes at 145°-150°. Sl. sol. hot water. Gives Liebermann's reaction. Forms an acetyl derivative C₆H₃MeO(NOAc) [92°] crystallising in large prisms.

Di-oxim $C_6H_3Me(NOH)_2[2:1:4]$. Formed by the action of hydroxylamine on either monooxim (Nietzki a. Guitermann, B. 21, 430) or by boiling nitroso-o- or m-toluidine with hydroxylamine hydrochloride (Mehne, B. 21, 729). low needles, exploding at 220°-234°, v. sl. sol. benzene, sol. hot water, alcohol, and ether. Reduced by tin and HClAq to tolylene-p-diamine. Alkaline K₂FeCy₆ forms di-nitroso-toluene. Ac₂O forms C₆H₂Me(NOAc)₂ [120°], crystallising from alcohol in colourless needles.

Ditolylimide C₆H₂Me(NC₇H₇)₂. [146°]. Formed by oxidation of tolylene-di-tolyl-diamine in alkaline solution (Green, C. J. 63, 1409). Red prisms, v. sol. alcohol, v. e. sol. benzene.

References .- DI-BROMO-, CHLORO-, IODO-, and OXY- TOLUQUINONE.

TOLUQUINOXALINE

METHYL - QUIN-OXALINE. o-TOLURIC ACID C, HINO, i.e.

C,H,.OO.NH.CH2.CO2H." Toluyl - glycocoll. [162.5°]. Formed from toluyl chloride, amidoacetic asid, and NaOH (Gleditsch a. Möller, A. 250, 876). Found in the urine of dogs that have taken o-toluic acid. Thick plates, sol. water.

m-Toluric acid. [139°]. Formed as above, and also when m-xylene or m-toluic acid is administered to a dog (G. a. M.; cf. Schultzen a. Naunyn, Z. 1868, 29). M. sol. water.—ZnA', 4aq.—CuA', 6aq.

p-Toluric acid. [161°]. Formed from

p-toluyl chloride and glycocoll (G. a. M.) and also by passage of p-xylene or p-toluic acid through the animal body (Kraut, A. 98, 360). Plates (from water), sl. sol. cold water, v. sol.

alcohol.—CaA'₂ 3aq.—BaA'₂ 5aq.—AgA'.

TOLUYL. The radiole CH₂.C₆H₄.CO. The same name has been given to the radicle CH₂.C₆H₄.OH₂, which is isomeric with xylyl, and has also been called tolyl, but may be more conveniently named methyl-benzyl. Tolyl

is CH, C,H,

o-TOLUYL-ACETIC ACID

Ethyl ether C6H4Me.CO.CH2.CO2Et. Oil, got by boiling the nitrile with alcohol and HCl.

Gives a violet colour with FeCl_s.

Nitrile C,H,.CO.CH,.CN. [74.4°]. Formed by boiling o-toluyl-cyano-acetic ether with water (Haller, C. R. 108, 1117). Prisms, sol. alcohol and alkalis. Diazobenzene chloride and NaOH form C_7H_7 .CO.CHCy.N₂.C₆H₅ [125°]. Cold alcoholic HCl forms the hydrochloride of C,H,.CO.CH2.C(NH).OEt [116°].

TOLUYL. ACRYLIC ACID C11H10Os i.e. C₆H₄(CH₃).CO.CH:CH.CO₂H. [138°]. Formed by the action of AlCl₃ on a mixture of maleic anhydride and toluene (Pechmann, B. 15, 888).

TOLUYL ALCOHOL v. Tolyl-carbinol. TOLUYLAMIDE v. Amide of TOLUIC ACID. p-TOLUYL-o-BENZOIC ACID C15H12Os i.e. [1:4] C₆H₄Me.CO.C₆H₄.CO₂H [1:2]. Phenyl tolyl ketone carboxylic acid. Mol. w. 240. [146°]. Formed by the action of phthalic anhydride on toluene in presence of AlCl_s (Friedel a. Crafts, A. Ch. [6] 14, 447; C. R. 92, 833). Prisms (containing aq) (from alcohol-toluene), v. sl. sol. hot water. Its dilute aqueous solution tastes sweet. Potash-fusion yields benzoic and p-toluic acids. Warm conc. H_2SO_4 forms (β) -methylanthraquinone. — BaA', 4aq. — CdA', 1aq. CuA'2 4aq.

Methyl ether MeA'. [53°].

Ethyl ether EtA'. [69°].
Toluyl-benzoic acid C,H,.CO.C,H,.CO2H. [222°]. Formed by oxidation of di-tolyl-methane and of u-di-tolyl-ethane (Weiler, B. 7, 1184; Ador a. Crafts, C. R. 85, 1163; O. Fischer, B. 7, 1195; Anschütz, A. 235, 316). Slender needles (from MeOH), sl. sol. hot water.—KA': tufts of long needles.—AgA'.

Reference. - DI-CHLORO-TOLYL-BENZOIC ACID.

TOLUYLENE- v. TOLYLENE-

o-TOLUYL-PROPIONIC ACID C11 h12O, i.e. C.H., CO.CH., CH., CO.H. [127°] (C.); [120°] (B.). Formed by the action of succinyl chloride or succinic anhydride on toluene and AlCl, (Claus, B. 20, 1378; Burcker, Bl. [2] 49, 448). or needles (from water), v. sol. ether.—BaA', 4aq: sol. hot water.—AgA': small needles, insol. Aq. w-TOLUYL-TOLUIC ACID. Lacton.

Lactone C_eH₄ CH:C.C,H, [98°]. Got from nitro-tolyl-

methylene-phthalide, HIAq, and P (Heilmann, B. 23, 3166). Thin prisms. Converted by alcoholic NH, into C,H, CO.NH [196].

oo-DITOLYL C_{1,}H_{1,} i.e. [1:2] C₄H₄Me.C₅H₄Me [1:2]. Mol. w. 182. (272°). S.G. 105 9945. Formed from o-bromo-toluene and Na (Fittig, A. 139, 178). Liquid.

mm-ditolyl [1:3] C₆H₄Me.C₆H₄Me[1:3]. (280°) (Schultz, B. 17, 468); (288°) (Perrier, C. R. 114, 484); (289°) (Stolle, B. 21, 1096). S.G. 1993. Formed by diazotising di-amidoo-ditolyl and boiling the product with alcohol. Formed also by the action of Na on m-bromotoluene (P.), and by reduction of di-oxy-m-ditolyl and di-chloro-m-ditolyl. Oil; yields Oil; yields diphenyl di-m-carboxylic acid and isophthalic acid on oxidation.

pp-ditolyl [1:4] $C_6H_4Me.C_6H_4Me$ [1:4]. [121°]. S.V. 1980. Formed by adding Na to a solution of p-bromo-toluene in toluene (Zincke, B. 4, 396; Louguinine, B. 4, 514). Monoclinic prisms (from ether). Yields on oxidation p-tolyl-benzoic acid [244°] and diphenyl di-p-carboxylic acid (Carnelley, C. J. 32, 653). SbCl₅ at 360° forms CCl, and per-chloro-diphenyl.

om-Ditolyl [1:2] C.H.Me.C.H.Me [1:3]. (270°). Formed from [2:4:1] $C_8H_3Me(NH_2)$. $C_8H_2Me(NH_2)$ [1:3:4] by elimination of amidogen by the diazoreaction (Schultz, B. 17, 471). Yields isophthalic

acid on oxidation by CrO,

op-Ditolyl [1:2] $C_6H_4Me.C_6H_4Me$ [1:4]. (272°-280°). Formed from crude bromo-toluene and Na. It is also one of the products of the passage of a mixture of benzene and toluene through a red-hot tube (Carnelley, C. J. 37, 707; 47, 587). Oxidised by CrO₃ and HOAc to o-tolylp-benzoic acid [180°] and terephthalic acid.

Ditolyls have also been obtained by Barbier $(C.\ R.\ 78,\ 1769)$ and Varet a. Vienne, $Bl.\ [2]\ 47$,

919).

References .- DI-AMIDO-, DI-BROMO-, DI-IODO-, NITRO-AMIDO-, OXY-AMIDO-, and OXY-DITOLYL.

p-TOLYL-ACETAMIDINE $C_0H_{12}N_2$ *i.e.* [96°]. Formed CH_3 .C(NH). NHC_6H_4Me [1:4]. from p-toluidine hydrochloride and acetonitrile (Bernthsen a. Trompetter, B. 11, 1757). Prisms. Alkaline in reaction.—B"2H2PtCl.—B"H2C2O. Prisms, v. sol. water and alcohol.

Di-o-tolyl-acetamidine $[1 \cdot 2]$ C₆H₄Me.NH.CMe:NC₆H₄Me [1:2]. [136°] (W.); [140°] (L.). Formed by the action of PCl. on o-toluidine and HOAc (Ladenburg, B. 10, 1262), and also by the action of o-toluidine on CMe(SEt):NC,H, or on CMeCl:NC,H, (Wallach a. Wüsten, B. 16, 148; A. 214, 208). Needles.

Di-p-tolyl-acetamidine [1:4] $C_6H_4Me.NH.CMe:N.C_6H_4Me$ [1:4]. tolyl-imid-tolyl-amide. [120°]. Formed from p-toluidine and CMeCl:NC,H,, which is the product of the action of PCl, on acetyl-p-toluidine Wallach a. Fassbender, A. 214, 203; B. 9, 1214; 16, 148). Formed also from p-toluidine and bromopropiolic acid (Mabery a. Krause, B. 22, 8307). Prisms (from benzene). Formed also from p-toluidine, HOAc, and PCl, -B',H2PtCl, -B'HCl. [200°].

op-Di-tolyl-acetamidine [142]. C.H.Me.NH.CMe:NC.H.Me. Formed from o-toluidine and the product of the action of PCl, on acetyl-p-toluidine, and also from p-toluidine and the product of the action of PCl, on acetyl-o-toluidine (Wallach a. Pannes, A. 214, 209). Formed also from p-toluidine and [1:2] C.H.MeN:CMe.SEt, and from o-toluidine and [1:4] C.H.MeN: CMeSEt; the compounds obtained, [143°] and [140°] respectively, being perhaps isomeric and not identical (Wallach, B. 16, 148)

o-TOLYL-ACETIC ACID C.H 10 O2 i.e. [89°]. Formed by [1:2] $C_6H_4Me.CH_2.CO_2H$. saponification of its nitrile (Radziszewski a. Wispeck, B. 15, 1747; 18, 1281). Needles, sl. sol. cold water. Yields phthalic acid on oxidation.—CaA', 4aq.—AgA'. Plates, sol. hot water Amide [161°]. Plates, sl. sol. ether.

Nitrile C.H.N. (244°). S.G. 22 1.0156. Formed from \(\omega\)-bromo-o-xylene and KCy. m-Tolyl-acetic acid [1:3] C₆H₄Me.CH₂.CO₂H. [61°]. Formed by saponifying its nitrile, which is obtained from ω-bromo-m-xylene (R. a. W.). Needles, sol. hot water.—CaA', 3aq.—AgA'.

Amide [141°]. Sl. sol. cold water. Nitrile (241°). S.G. 22 1.0022. Oil.

Methylether MeA'. (228°). S.G. 175 1.044.

Liquid (Senkowski, M. 9, 854).

Ethyl ether Eth. (238°). S.G. 175 1.018. p-Tolyl-acetic acid [1:4] C_0H_4 Me. CH_2 . CO_2H_4 . [946]. Formed in like manner from w-bromo-pxylene (R. a. W.; Strassmann, B. 22, 1230). Got also by reducing p-tolyl-glyoxylic acid with HI (Claus a. Kroseberg, B. 20, 2051). Needles, sl. sol. cold water. Phthalic anhydride and NaOAo yield $C_6H_4 < \frac{CO}{C(CH.C_6H_4Me)} > O$ [151°], whence HNO2 at 0° forms the compound $C_6H_4 < C(NO_2).CH(NO_2)C_7H_7 > 0$ [140°], which is converted by treatment with boiling alcohol into $C_6H_4 < CO \over C(CNO_2)C_7H_7 > O$ [207°], and is reduced by HI and P to $C_0H_4 < \stackrel{CH:C.C,H_7}{CO.O}$ [116°] (Ruhe-

Ethyl ether EtA'. (240°). Amide C.H., NO. [185°]. Plates. Formed from the acid, and also by the action of yellowammonium sulphide on p-tolyl methyl ketone

mann, B. 24, 3965).—NaA': needles, sol. water.-

CaA', 3aq.—AgA': needles, sol. hot water.

(Willgerodt, B. 21, 534). Nitrile C.H. CN. [18°]. (243°). S.G.

·9922.

Reference. - Nitro- and Oxy-Tolyl-ACETIC ACID.

m-TOLYL-ACETYLENE [1:3] C.H.Me.C:CH. (164°). Formed by distilling Methyl-styrene. barium m-tolyl-propiolate with lime (Müller, B. 20, 1215). Formed also by saturating a solution of m-tolyl-acrylic acid with HBr at 0° and treating the product with NaOHAq. Oil, becoming solid on standing. Gives an explosive red pp. with ammoniacal Cu2Cl2. Br forms C,H,.CHBr.CH₂Br [45°].

C,H,.C:C.C,H,. Di-tolyl-acetylene Formed by heating C,H,.CHBr.CHBr.C,H , with alcoholic potash at 140° (Goldschmiedt a. Hepp, B. 6, 1505). Needles (from alcohol).

DI-TOLYL-ACIPIPERAZINE OXY-DI-

TOLYL-PYRAZINE.

Di-tolyl-di-aci-piperazine DI-OXY-DI-TOLYL-PYRAZINE.

m-TOLYL-ACRYLIC ACID C10H10O2 i.e. [1:3] C₄H₄Me.CH:CH.CO₂H. Methyl-cinnamic acid. [111°]. Formed by heating m-toluic aldehyde with Ac, O and NaOAc (Bornemann, B. 17, 1474; 20,1382; Von Miller, B. 20,1213; 23,1899). Silky needles, v. sol. alcohol, sl. sol. ligroin. FeCl, gives a bright-yellow pp. in neutral solutions. Reduced by sodium-amalgam to m-tolyl-propionic acid [43°]. KMnO, oxidises it to m-toluic acid. Br forms C,H,.C,H,Br,.CO,H [167°].-BaA'2—AgA'. Crystalline powder.

p-Tolyl-acrylic acid [1:4] C₆H₄Me.CH:CH.CO₂H. [197°]. Formed from p-toluic aldehyde, NaOAc, and Ac₂O (Von Miller, B. 23, 1897; cf. Kröber, B. 23, 1033). Needles (from benzene), sl. sol. hot water. Reduced by HI and P to p-tolyl-propionic acid. Br forms C_sH₄Me.CHBr.CHBr.CO_sH.

TOLYL ALCOHOL v. Tolyl-Carbinol.

p-TOLYL-ALLYL-HYDRAZINE C₁₀H₁₄N₂ i.s. C,H,N(C,H,).NH₂. (160°-170° at 90 mm.). Got from tolyl-hydrazine and allyl bromide (Michaelis, B. 26, 2178). Oil. Yields B'HCl. Benzoic aldehyde forms the compound C,H,.N(C,H,).N:OHPh [61°], while cinnamic aldehyde forms the corresponding hydrazide [118°]. Ferric chloride gives the tetrazone $(C_1H_1(C_2H_3)N)_2N_2$ [104°],

o - TOLYL-ALLYL-THIO-SEMI-CARBAZIDE C,H,NH.NH.CS.NHC₈H₅. [105°]. Formed by warming o-tolyl-hydrazine with allyl thiocarbimide (Avenarius, B. 24, 268). Needles, insol. water, v. sol. alcohol and CS2

p - Tolyl - allyl - thio - semi - carbazide [128°].

Formed in like manner (A.). Needles.

o-TOLYL-ALLYL-THIO-UREA C11H14N2S i.e. C,H,NH.CS.NHC,H, Tolyl-thiosinamine. [98°]. Formed from o-toluidine and oil of mustard (Jaillard, Z. 1865, 441; Maly, Z. [2] 5, 258; Weith, B.8, 1528; Prager, B. 22, 2998). Pearly crystals, m. sol. ether, v. sol. HOAc. According to Dixon (C. J. 55, 622), it melts at 76°. Converted by heating with HClAq into tolyl-propylene-ψ-thio-urea Cyanogen passed into its alcoholic [126°]. solution forms C18H14N4S, which on warming with dilute H,SO, and alcohol gives a pp. of $CS < NC_3H_5 > C_2O_3$ oxalyl-tolyl-allyl-thio-urea [157°]

Reference. -- OXY-TOLYL-ALLYL-THIO-UREA.

O-TOLYL-AMIDO-ACETIC ACID C.H.INO. i.e. C.H.Me.NH.CH. CO.H. Tolyl - glycocoll. [150°]. Prepared by the action of chloro-acetic acid and aqueous NaOAc on o-toluidine (Staats, B. 13, 137; Cosack, B. 13, 1091; Ehrlich, B. 16, 204; Bischoff a. Nastvogel, B. 22, 1787).-CuA', 2aq. Small needles.—CaA', 3aq: needles, m. sol. water (Mauthner, M. 11, 377). Ethyl ether EtA'. (281°). S.G. 30 1.058.

(281°). S.G. 30 1.058. Ethyl ether EtA'. Oil (Bischoff, B. 25, 2270).

Acetyl derivative C,N,NAc.CH2.CO2H. 2°]. Tables (from dilute alcohol).

Chloro-acetyl derivative C,H,N(CO.CH,Cl).CH,.CO,H. [117]. Formed from the acid and chloro-acetyl chloride in ether (Widman, J. pr. [2] 38, 304). Rectangular plates, v. sol. hot alcohol, sl. sol. water.

Bromo-acetyl derivative C,H,N(CO.CH,Br).CH,CO,H. [124°]. Rectan.

gular plates.

Glycolyl derivative C,H,N(CO.CH₂.OH).CH₂.CO₂H. [144°]. Formed by boiling the chloro-acetyl derivative with conc. Na_2CO_3Aq (Abenius, J. pr. [2] 40, 503). V. e. sol. water. Yields KA' aq, BaA'₂7aq, and AgA', and is converted by heating at 160° into $C_rH_rN<_{CH_rCO}^{CO.CH_2}>0$ [109°], whence alcoholic NH₂ produces C₂H₂N(CO.CH₂·OH).CH₂.CO.NH₂ [152°], crystallising in plates (from alcohol).

o - Toluide C,H,.NH.CH,.CO.NHC,H,. [92°]. Formed by boiling chloro-acetic ether with o-toluidine (Ehrlich, B. 16, 205). Needles,

insol. water and HClAq.

m-Tolyl-amido-acetic acid [1:3]C₆H₄Me.NH.CH₂.CO₂H. Formed from chloro-acetic acid and m-toluidine in ether

(Ehrlich, B. 15, 2011). Solid mass.—CuA'₂ 2aq. Ethyl ether EtA'. [68]. Formed from chloro-acetic ether and m-toluidine. Six-sided plates, v. sol. alcohol and ether, sl. sol. Aq.

p-Tolyl-amido-acetic acid [1:4]C₆H₄Me.NH.CH₂.CO₂H. [119°]. Formed, together with a compound [168°], from p-toluidine chloro-acetic acid and NaOAc (dissolved in a little water) at 100° (P. J. Meyer, B. 8, 1158; 14, 1328; Staats, B. 13, 137; Schwebel, B. 10, 2047; Bischoff, B. 23, 1997; 25, 2281). Got also by saponifying its ether. Needles (from water). When fused with potash it yields a product which, when dissolved in water, is oxidised by air to an indigo-derivative (Heumann, B. 24, 1346). When fused with p-chloro-acet-toluide it yields methyl-indigo (Eckenroth, B. 24, 693). The compound [168°], formerly supposed to be p-tolyl-amido-acetic acid, ought, according to Bischoff a. Hausdörfer (B. 25, 2281) to be represented as CO₂H.CH₂.N(C₇H₇).CH₂.CO.ONH₃C₇H₇.

Ethyl ether Eth'. [49°]. Formed from chloro-acetic ether and p-toluidine. Monoclinic

plates, v. sl. sol. hot water.

[163°]. Formed by Amide C₉H₁₂N₂O. heating p-toluidine with chloro - acetamide. Plates, v. sl. sol. cold water.

Anilide C,H,NH.CH,CO.NHPh. Formed by heating chloro-acetic acid with

Needles (from hot water).

p-Toluide C, H, NH.CH2.CO.NC, H,. [136°]. Formed in like manner, and also by heating toluidine with the compound of glyoxal with NaHSO_s (Hinsberg, B. 21, 112). Plates, v. sl. sol. hot water. Yields a chloro-acetyl derivative [158°

Nitrile C,H,NH.CH2.CN. [126°]. A pro-

duct of the action of heat on the amide.

Acetyl derivative C,H,.NAc.CH,.CO,H. [175°]. Formed from chloro-aceuc etner and C,H, NAcNa, the product being saponified with potash (Paal a. Otten, B. 23, 2596). Plates (from water).—NaA' 3aq: plates, v. sol. water.
I mide (C,H,NH.CH,CO),NH. [210°

[210°]. Got by heating the acid at 200° and adding NH, Aq to the product. Needles, sl. sol. water

and NH, Aq (Bischoff).

Di-o-tolyl-di-amido-acetic acid $(C_7H_7NH)_2CH_1CO_2H_2$ [240°]. Formed heating di-chloro acetic acid with o-toluidine (P. Meyer, B. 16, 925). Needles, insol. water, sl. sol. alcohol, v. sol. aqueous acids and alkalis.

Reference. -- NITRO-TOLYL-AMIDO-AGETIC ACID.

p-TOLYL-AMIDO-ACETOPHENONE

[1:4]C_sH_sMe.NH.CH₂.CO.C_sH_s. Phenacyl-p toluidine. [134°]. Formed from p-toluidine and bromo-acetophenone in alcohol (Lellmann a. Donner, B. 23, 167). Tables, v. sol. benzene, m. sol. alcohol. Yields, on nitration, [1:3:4]C₀H₂Me(NO₂).NH.CH₂Bz [165°] and a dinitro-derivative [156°]. Forms B'HNO₂ and B'HCl.

p-TOLYL-AMIDO-ACRYLIC ETHER

[1:4]C₆H₄Me.NH.CH:CH.CO₂Et. [116°]. Formed from p-toluidine and formyl-acetic ether (Von Pechmann, B. 25, 1053). Yellow plates, insolwater and ligroin. Alcoholic potash yields water and ligroin. Alcoholic potash yields plates [144°]. Ac O forms acetyl-toluidine. TRI-p-TOLYL-TRI-AMIDO-BENZENE

C₂,H₂,N₃ i.e. C₅H₃(NHC,H₁)₃. [187°]. Formed by heating phloroglucin with *p*-toluidine at 210° (Minunni, *B*. 21, 1984). Needles, v. sl. sol. cold alcohol. Conc. H₂SO₄ forms a bluish-green solution, turning black on heating.—B"HCl: amorphous; decomposed by water.—B"'2HCl: small needles.-B"2H2PtCls.

Tri-acetyl derivative [193°]. Plates. Tri-benzoyl derivative [282°]. Minute

p-TOLYL-AMIDO-BENZOYL-(a)-NAPHTHO-C₆H₅.CO.C₆H₃ CO.C.NHC₆H₄Me. QUINONE

Formed by boiling p-toluidine with benzoyl-(a)-naphthoquinone in alcoholic solution for 15 minutes (Kegel, A. 247, 185). Red needles (from HOAc), v. sl. sol. alcohol

p-TOLYL-o-AMIDO-BENZYL-AMINE [4:1] $C_6H_4Me.NH.CH_2.C_6H_4.NH_2$ [1:2]. [80·5°]. Got by reducing p-tolyl-o-nitro-benzyl-amine (Söderbaum a. Widman, B. 23, 2187). Thin leaflets (from alcohol). Ac₂O yields a di-acetyl derivative [186°]. The mono-acetyl derivative C,H,NAc.CH₂,C,H,NH₂ [99°] may be got by reducing C,H,NAc.CH₂,C,H,NO₂.

o-TOLYL-α-AMIDO-n-BUTYRIC ACID

C,H,NH.CHEt.CO,H. [84°]. Long prisms. Forms an acetyl derivative [116°] and yields propyl-o-toluidine on distillation.

Ethyl ether EtA'. (278°). S.G. 20 1.019. Got by heating o-toluidine with a-bromo-butyric ether at 100° (Bischoff a. Mintz, B. 25, 2314).

o-Tolyl-a-amido-isobutyric acid C₇H₇NH.CMe₂.CO₂H. [62°]. Formed together with the β - isomeride [110°] by the action of KOH and a current of steam on the ether [57°], which is got by heating o-toluidine with a-bromoisobutyric ether (Bischoff a. Mintz, B. 25, 2834). The acid [110°] yields an acetyl derivative [219°], sl. sol. ether. On distillation it forms a lactone [95°].

p-Tolyl- α -amido-n-butyric acid

C.H.NH.CHEt.CO.H. [153°-156°]. Got from the ether EtA' (279°) S.G. 28 1.011, which is produced when p-toluidine is heated with a-bromo-n-butyric acid at 105°. Leaflets. Yields propyl-p-toluidine on distillation.

Acetyl derivative C,H,NAc.CHEt.CO,H.

[149°].

p-Tolyl-a-amido-isobutyric acid [1:4] C₆H₄Me.NH.CMe₂.CO₂H. [150°]. Formed, together with the 8- isomeride, by the action of hot potash in a current of steam on the ether EtA' [36°], which is got by heating p-toluidine with a-bromo-isobutyric acid. Yields an acetyl

derivative [146°]. Forms isopropyl-p-toluidine on distillation.

p-Tolyl-eta-amido-isobutyric acid

C,H,NH.CH2.CHMe.CO2H. [196°]. Formed as above. Gives an acetyl derivative [206°] and decomposes on distillation into p-toluidine and

a lactone $C_7H_7N < \stackrel{CH_2\cdot CHMe}{CO} \stackrel{C}{\cdot CMe_2\cdot O} > CO$ (?) [170°].

o-TOLYL-AMIDO-CROTÓNIC ACID [1:2] C₆H₄Me.NH.CMe:CH.CO₂H.

Methyl ether MeA'. [31°]. Formed by the action of o-toluidine on methyl aceto-acetate (Conrad a. Limpach, B. 21, 523). Prisms (from alcohol).

Ethyl ether EtA'. Yellowish oil. p-Tolyl-amido-crotonic acid.

Methyl ether MeA'. [60°]. Prisms. Ethyl ether EtA'. [29°]. Formed from

acetoacetic ether and p-toluidine.
TOLYL-AMIDO-CYANURIC

ACID Cyanuric acid, described under CYANIC ACIDS. p-TOLYL-AMIDO-METHENYL-BENZAMID-

OXIM CPh $\stackrel{\text{N.O}}{\sim}$ C.NH.C.H.Me[1:4.]. [135°].

Formed by heating p-tolyl-thiocarbimide with benzamidoxim (Koch, B. 24, 398). Plates (from alcohol), v. sol. ether.

 $\mathbf{DI} \cdot p \cdot \mathbf{TOLYL} \cdot \mathbf{AMIDO} \cdot \mathbf{METHYLENE} \cdot o \cdot \mathbf{PHE}$ NYLENE DIAMINE C21H22N4 i.e.

 $C_eH_4 < NH > 0 < NH.C_7H_7$. [186°]. (above 400°). Formed by heating C(NC,H,)2 with o-phenylenediamine at 135° (Moore, B. 22, 1639; 3188). Tables (from alcohol), v. sl. sol. boiling water.
—Salts: B'28HCl. Needles.—B',3H2PtCl₆8aq. -B'H₂SO₄. Needles, insol. ether, v. sol. hot Aq.

Nitrosamine $C_6H_1 < N(NO) > C(NC_7H_7,NO)_2$. [130°]. Yellow needles, v. sol. alcohol.

Di-acetyl derivative [143°]. Needles. Di-benzoyl derivative [185°]. P. Tetra-benzoyl derivative [274°]. Prisms.

Tetra-p-tolyl-tetra-amido - di - methylene - o- $C_6H_4:N_2(C(NHC_7H_7)_2)_2.$ phenylene-diamine [116°]. Formed by heating C(NC,H,)2 with ophenylene-diamine at 200° (Moore, B. 22, 3195). Needles, v. sol. alcohol. Decomposed by dilute HClAq at 150° into di-p-tolyl-urea and ophenylene-diamine.—B'2H₂SO₄. Needles, sl. sol. cold water. -B',3HCl. -B',3H,PtCl, 15aq.

Tetra-acetyl derivative [115°

Tetra-bensoyl derivative [146°]. DI-p-TOLYL-AMIDO - METHYLENE-o-PHE-NYLENE-PHENYL-GUANIDINE

 $C:NPh C(NHC_1H_1)_x$ [187°]. Formed

by boiling $C(NC_7H_7)_2$ with o-phenylene-phenylguanidine (Keller, B. 24, 2508). Needles (from

boiling alcohol), sl. sol. ether. DI - p - TOLYL - AMIDO - METHYLENE - o -

PHENYLENE-p-TOLYL-GUANIDINE
C₆H₄:N₂(CNC,H₇):C(NHC,H₇)₂. [188°]. Formed
by heating o-phenylene-p-tolyl-guanidine with
C(NC,H₇)₂ at 220° (Keller, B. 24, 2513). Prisms
(from benzeme). (from benzene).

DI - p - TOLYL - AMIDO - METHYLENE - o -TOLYLENE-p-TOLYL-GUANIDINE C,H,:N₂(C:NČ,H₁):O(NHC,H₁)₂ [210°]. Formed by heating tolylene-tolyl-guanidine together with C(NC,H,)₂ at 220° (Keller, B. 24, 2521). Needles, sl. sol. alcohol.

o-TOLYL- AMIDO - METHYL - MALONAMIC ETHER CO₂Et.CMe(NHC,H₁).CO.NH₂. Formed by dissolving a-cyano-tolyl-a-amido-propionic ether in conc. H₂SO₄ (Gerson, B. 19, 2966). Long needles, sol. hot water. Converted by boiling NaOHAq into tolyl-amido-propionic acid.

p-TOLYL-AMIDO-p-METHYL-OXINDOLE

 $C_{16}H_{16}N_2O$ i.e. $C_6H_9Me < \frac{NH}{CH(NHC,H_2)} > CO$.

[167°]. Formed by heating di-chloro- (or dibromo-) acetic acid with p-toluidine (Duisberg, B. 18, 190). Needles, v. sol. hot alcohol. Reduces ammoniacal AgNOs, forming a mirror. Readily absorbs oxygen from the air, forming $C_eH_sMe < NH_{C(NC,H_s)} > CO$. Alcoholic KOH forms a blood-red solution.—B'HCl: crystalline

powder.

Di-acetyl derivative C₁₆H₁₄Ac₂N₂O. [147°].

Nitrosamine C₁₆H₁₃N₃O₂. [c. 220°]. p-TOLYL-AMIDO-METHYL-THIAZOLE

S<CH=_____CMe. [125°]. Formed oxy-methyl-thiazole and p-toluidine (Hantzsch a. Weber, B. 20, 3130).

TRI-p-TOLYL-TRI-AMIDO-NAPHTHALENE $C_6H_4 < \stackrel{C(NHC,H_1):C(NHC,H_2)}{C(NHC,H_2):CH}$. [160°]. Formed by heating nitroso-ethyl-(a)-naphthylamine with p-toluidine at 150° (Fischer a. Hepp, A. 256, 244). Needles (from alcohol). Readily oxidised

to di-methyl-phenyl-rosinduline o-TOLYL-AMIDO-(a)-NAPHTHOQUINONE $C_{17}H_{13}NO_2$ i.e. $C_{10}H_5O_2(NHC_7H_7)$. [142°]. Formed from o-toluidine and (a)-naphthoquinone (Elsbach, B. 15, 689). Red needles, sol. NaOHAq.

p-Tolyl-amido-(a)-naphthoquinone C₁₀H₂O₂(NHC₇H₇). [202°]. Formed from (a)-naphthoquinone and p-toluidine in hot alcoholic solution (Plimpton, C. J. 37, 638). Formed also by the action of p-toluidine on (β) -oxy- (α) naphthoquinone, and by heating p-tolyl-amido-(B)-naphthoquinone with HOAc at 150° (Elsbach, B. 15, 687). Red needles. Forms a crimson solution in conc. H2SO4. Insol. NaOHAq.

 $C_6H_4 < C(NC_7H_7):C.NHC_7H_7$ Ditoluide [147°]. Formed by oxidation of tri-p-tolyl-amidonaphthalene (Fischer a. Hepp, A. 256, 246).

Orange plates. o-Tolyl-amido-(β)-naphthoquinone

 $C_{10}H_5O_2(NHC_7H_7)$ or $C_{10}H_5(OH)O(NC_7H_7)$. [240°]. Formed from (β) -naphthoquinone and o-toluidine (Elsbach, B. 15, 689). Red needles, sol. dilute NaOHAq. Not affected by heating with HOAc at 150°. On heating with dilute HClAq it yields (β)-oxy-(α)-naphthoguinone and o-toluidine.

p - Tolyl - amido - (β) - naphthoguinone [246°]. Formed from (8)-naphthoquinone and p-toluidine (Elsbach, B. 15, 686). Red needles, sol. hot alcohol and NaOHAq. On heating with HClAq at 130° it is split up into p-toluidine and (B)naphthoquinone.

Methyl ether MeA'. [150°]. Red crystals

(Zincke a. Brauns, B. 15, 1969). Ethyl ether Eth'. [137°] Isopropyl ether PrA'. [139°].

p-Toluide C₁₀H₃(NHC,H₇)O(NC,H₇). [183° cor.]. Formed by heating di-bromo-(a)-naphthol with p-toluidine (Meldola, C. J. 45, 159). Formed also from (β) -naphthoquinone oxim, p-toluidine, and HOAc (Bromme, B. 21, 391). Orange crystals with green lustre.

DI-o-TOLYL-DI-AMIDO-OIAZTHIOLE

 $\mathbf{C}_{10}\mathbf{H}_{10}\mathbf{N}_{4}\mathbf{S}$ i.e. $\mathbf{S} < \overset{\mathbf{C}(\mathbf{NHC},\mathbf{H}_{7}):\dot{\mathbf{N}}}{\mathbf{C}(\mathbf{NHC},\mathbf{H}_{9}):\dot{\mathbf{N}}}$. [185°]. Formed by boiling an alcoholic solution of o-tolyl-thioby boiling an alcoholic solution of o-tolyl-thiourea with dilute HClAq and H₂O₂ (Hector, B.
23, 368). V. sol. alcohol. Cyanogen passed into
its alcoholic solution forms C_{1e}H_{1e}N₄SCy₂[89°].—
B'HClaq. [219°].—B'₂H₂PtCl_e. [192°].—
B'AgNO₃. White pp.—B'₂H₂PtCl_e. [232°].
Needles.—Picrate: B'C₆H₁₆N₄O₇. [201°].

Acetyl derivative C_{1e}H₁₅AcN₄S. [221°].

Bensoyl derivative C_{1e}H₁₆BzN₄S. [214°].

Nitrosamine C_{1e}H₁₅(NO)N₄S. [135°].
Di-p-tolyl-di-amido-oiagthiole C_{1e}H_{1e}N₅S.
[127°]. Formed in like manner from p-tolylthio-urea (Hector, B. 23, 364). Thick vellowish

thio-urea (Hector, B. 23, 364). Thick yellowish prisms, v. sol. alcohol. Cyanogen forms B'Cy₂ [190°].—B'HCl. [233°]. White needles, becoming reddish in air.—B'₂H₂PtCl₃ l¹₂aq. [109°].—B'AgNO₃ l¹₂aq. [177°].—B'₂H₂HgCl₄. [114°—119°].—Picrate [238°]. Yellow prisms, sl. sol. alcohol.

Acetyl derivative C₁₈H₁₃AcN₄S. [166°].

Benzoyl derivative C₁₆H₁₃BzN₄S. [186°].

Nitrosamine C₁₈H₁₅(NO)N₄S. [247°].

o-TOLYL-m-AMIDO-PHENOL C₁₈H₁₃NO i.e.
C₇H₁NH₂C₄H₂OH. (c. 373° cor.). Formed from resorcin, o-toluidine, and CaCl at 270° (Philip, J. pr. [2] 34, 70). Oil. Reduced by hot zincdust to hydro-acridine and acridine.

Formyl derivative. [169°]. Tables. C,H,NH.C,H,OH. o-Tolyl-p-amido-phenol [90°]. (367° cor.). Formed from hydroquinone, o-toluidine and CaCl₂ at 240° (Philip, J. pr. [2] White laminæ (from light petroleum). Reduced by distilling with zinc to hydro-acridine and acridine. Converted by heating with conc. HCl into hydroquinone and o-toluidine.

Salt .- B'HCl : crystalline powder. Mono-formyl derivative [136.5°].

Di-acetyl derivative [106°]: needles.

Di-benzoyl derivative. [171°].

p-tolyl-m-amido-phenol C₁₈H₁₈NO

I.NH.C₆H₄OH. [92°]. (350° cor.). Fr

C,H,NH.C,H,OH. [92°]. resorcin, p-toluidine and CaCl, at 260° for 8 hours (Hatschek a. Zega, J. pr. [2] 33, 209). Prisms or needles (from benzene mixed with petroleum ether). Distilled over zinc-dust, it gives C,H,NHPh.

Salts.—B'HCl; decomposed by water.

Formyl derivative. C,H,.N(CHO).C,H,OH. [146°]. V. sol. ether.

Di-benzoyl derivative. C,H,NBzC,H,OBz. [c. 105°]. Gives with fuming

HNO, a di-nitro-derivative, [110].
Nitrosamine.—C,H,N(NO).C,H,OH.Oxy-[10c°]. phenyl - tolyl - nitrosamine. Yellow

needles (from dil. alcohol).

p-Tolyl-p-amido-phenol [122°]. (850° 860°) C,H,.NH.C,H,.OH. p-101y1-p-amino-pienor of the first state of the fi

Di-bensoyl derivative. [169°]. Nitroso-derivative C,H,.N(NO).C,H,.OH [180°]; needles. Very unstable.

o-TOLYL-a-AMIDO-PHENYL-ACETIC ACID NHC,H,.CHPh.CO,H. [143°]. Formed from abromo-phenyl-acetic acid and o-toluidine (Stöckenius, J. 1878, 781). Plates, almost insol. water.

p-Tolyl-a-amido-phenyl-acetic acid. Formed in like manner, using p-toluidine. Insol. water. Ethyl ether EtA'. [90°]. Yellowish prisms.

o-TOLYL - AMIDO-TRI-PHENYL-METHANE CPh₃.NHC,H₇. [142°]. Formed from CPh₃Br and o-toluidine (Wittich, B. 17, 705). Prisms (from ether).

p-Tolyl-amido-tri-phenyl-methane [177°].

Crystals (from ether).

[145°-Nitrosamine CPh₃.NC₇H₄.NO. 148°]. p-TOLYL-AMIDO-DI-PHENYL-NAPHTHO-

QUINOXALINE. Phenylo-chloride $C_6H_4 < \frac{C(NH.C_6H_4Me):CH.C.N=CPh}{C(NH.C_6H_4Me):CH.C.NPhCl:CPh.}$

Got by heating C₁₀H₈(OEt) N=CPh with p-NPhCl:CPh toluidine at 120° (Witt a. Schmidt, B. 25, 2005). Bronzed plates, v. sol. alcohol, forming a crimson solution. Conc. H₂SO₄ forms a slate-blue solution, becoming first blue and then violet on dilution.

TRI-p - TOLYL - TRI - AMIDO - DI-PHENYL-TOLYL-CARBINOL C41 H30 N3O. Tri-tolyl-rosaniline. The salts are formed by heating rosaniline salts with p-toluidine (Hofmann, A. 132, 290).— C₄₁H₃₈N₃Cl. Small blue crystals (from alcohol), insol. water.

TOLYL-AMIDO-PIPITZAHOIC ACID Toluido-PIPITZAHOIC ACID.

o-TOLYL-a-AMIDO-PROPIONIC ACID

CH, CH(NHC, H,).CO,H. CH. CH(NHC, H,).CO.H. [116°]. Formed by saponification of the nitrile (Tiemann a. Stephan, B. 15, 2038; C. C. 1886, 470). Got also by boiling o-tolyl-amido-methyl-malonic acid with NaOHAq (Gerson, B. 19, 2963). White crystals. Ethyl ether EtA' (278°). S.G. 20 1.047.

Formed from a-bromo-propionic ether and o-toluidine (Bischoff a. Hausdörfer, B. 25, 2298). Acetyl derivative C,H,NAc.CHMeCO,H.

[177°]. Leaflets.

Amide. [125°]. Minute needles. Nitrile CH_a.CH(NHC,H₁).CN. [73°]. Formed by digesting the cyanhydrin of acetic aldehyde (lactonitrile) with o-toluidine.

p - Tolyl - α -amido - propionic acid. Formed in like manner. Plates, sl. sol. water and ether. May be sublimed.

Ethyl ether EtA'. [35°]. Tables.

Acetyl derivative [166°].

Amide. [145°]. Needles, sol. hot Aq. Nitrile. [82°]. Plates, sl. sol. hot Aq.

p-Tolyl-β-amido-propionic acid

C.H.Me.NH.CH2.CH2.CO2H. [86°]. Formed from β-iodo propionic acid and p-toluidine (Bischoff a. Mintz, B. 25, 2352). Pearly scales, v. e. sol.

fication of o-tolyl-amido-pyrotartramic ether, which is formed by the action of H2SO, upon the a-cyano-a-o-toluido-butyric ether (Schiller-Wechler, B. 18, 1050). White needles. V. sol. alcohol and benzene, sl. sol. ether and hot water, insol. cold water and ligroin. Has both acid

and basic properties.

DI-p-TÔLYL-DI-AMIDO-SUCCINIC ACID CO₂H.CH(NHC,H_.).CH(NHC,H_.).CO₂H. [c. 200°]. Formed, together with 'di-tolil succin' C₁₈H₁₆N₂O₂ [228°], by boiling di-bromo-succinic acid with p-toluidine (Junghahn a. Reissert, B. 26, 1766). V. sol. alcohol, insol. water. Yields a di-nitroso- derivative [125°], and a di-acetyl derivative C₂₂H₂₁N₂O₈ aq, which yields CaA" aq, BaA" aq, and anhydrides C₂₂H₂₂N₂O₈ [232°], and C₂₂H₂₀N₂O₄ [222°].—Na₂A".—CaA".—CuA".

Ethyl ether EtA". [169°]. Needles.

DI-p-TOLYL-TRI-AMIDO-TOLUENE $C_6H_2Me(NHC_7H_7)_2(NH_2)$ [1:2:5:4]. [166°]. Formed by reducing amido-toluquinone ditoluide with zinc-dust and HOAc (Green, C. J. 63, 1403; cf. Barsilowsky, B. 6, 1209). Plates (from dilute HOAc). A very feeble base.

DI-p-TOLYL-DI-AMIDO-TOLUQUINONE $C_8HMe\tilde{O}_2(NHC_7H_7)_2$ [1:2:5:4:6]. [178°]. Formed from azotoline, MeOH, and conc. H_2SO_4 (Fischer a. Hepp, A. 262, 251). Brownish-red needles.

Di-p-tolyl-di-amido-toluquinone $C_6HMeO_2(NHC_7H_7)_2$ [1:2:5:3:6]. [241°]. Formed by boiling toluquinone with p-toluidine and al-cohol (Fischer a. Hepp, A. 256, 259). Yellow

Di-p-tolyl-di-amido-toluquinone p-toluide $C_{28}H_{27}N_3O$. [191°]. Claret-red plates (Fischer a. Hepp, B. 21, 676).

TOLYLAMINE is TOLUIDINE (q.v.).

Di-o-tolyl-amine NH(C6H4Me[1:2])2 i.V.) at 728 mm. (Graebe, A. 238, 363). Obtained, together with diphenylamine by heating o-toluidine with aniline hydrochloride at 280° (Girard a. Willm, Bl. [2] 25, 248). Formed also by heating o-cresol with ZnBr₂, ammonia, and NH₂Cl at 335° (Merz a. Müller, B. 20, 547). Oil.

Di-m-tolyl-amine NH(C,H₁)₂. (322°). Formed in like manner (Cosack, B. 13, 1091; Merz a. Müller, B. 20, 549). Oil, volatile with steam.

Sl. sol. acids, v. sol. alcohol.

Acetyl derivative NAc(C,H,)2.

Nitrosamine (C,H,)₂N.NO. [103°].
Di-p-tolyl-amine NH(O,H₂)₂. [79°]. (831°
i.V.) (Graebe, A. 238, 363). Formed by heating p-toluidine with its hydrochloride (Girard, A. 140, 346; Gerber, B. 6, 446). Formed also by heating p-cresol with ZnCl2, ammonia, and NH, Cl at 335° (Merz a. Müller, B. 20, 546). White needles. Its salts are decomposed by

 $(C,H_1)_2N.NO.$ [103°]. Nitrosamine Yellow trimetric crystals (Lehne, B. 19, 1544;

Cosack).

Acetyl derivative (C,H,)2NAc. [85°]. Bensoyl derivative (C,H,)2NBz. [125°]. References .- TETRA-BROMO- and NITEO- DI-TOLYL-AMINE

DI-p-TOLYL-ISOAMYL-AMINE C1. H2.N i.e. (C,H,)2NC,H11: (290°-300° at 15 mm.). Formed by heating di-p-tolyl-amine with isoamyl alcohol and HCl in sealed tubes (Girard, Bl. [2] 24,

TOLYL-ANILINE v. PHENYL-TOLUIDINE. Vol. IV.

TOLYL-ARSINE v. ARSENIC.

TOLYL-AURAMINE C24H27N, 6.6. C.H.Me.N:C(C.H.NMe.)2. Formed from auramine hydrochloride and p-toluidine (Fehrmann, B. 20, 2853). Decomposed by H_2S in alcoholic solution at 50° into $CS(C_0H_4NMe_2)_2$ and p-toluidine.— $B'_2H_2PtCl_4$. Red flakes, v. sl. sol. water.

TOLYL-AZIMIDO-TOLUENE

COMPOUNDS

DI - p- TOLYL - TETRAZINE C16H16N4 i.e. $C_eH_iMe.N < N = CH > N.C_eH_iMe.$ [185°]. Mol. w. 269 (obs.). Formed by the action of chloroform and alcoholic potash on p-tolyl-hydrazine (Ruhemann, C. J. 55, 248; 57, 51). plates (from hot alcohol). Yields Yields a nitroderivative $C_{16}H_{15}(NO_2)N_4$ [144°].

TOLYL - DISAZO - compounds v. DISAZO-

TOLYL BENZAMIDE v. Benzoyl derivative of TOLUIDINE.

p - TOLYL - BENZAMIDINE $C_{14}H_{14}N_2$ C₆H₃.C(NC₇H₇).NH₂. [99°]. Formed from benzonitrile and toluidine hydrochloride (Bernthsen, B. 8, 1319; 9, 429). Forms a crystalline nitrate and oxalate.

Di-p-tolyl-benzamidine

 C_eH_s . $C(NC_rH_s)$. NHC, H_r . [131°]. Formed by the action of p-toluidine on benzonitrile, and on ω chloro-benzylidene-p-toluidine C,H,N:CCl.C,H, (Bernthsen; Just, B. 19, 981). May be sublimed. Reference. - AMIDO-TOLYL-BENZAMIDINE.

p-TOLYL-BENZAMIDOXIM C₁H₁N₂O i.e. C_cH_c.(NOH).NHC,H_c. [176°]. Formed by heating hydroxylamine hydrochloride with C₄H_c.CS.NHC,H_c and Na₂CO₂ at 100° (H. Müller, P. 20.2406). White needles, sl. sol. hot water .-B. 22, 2406). B'HCl: white needles, sl. sol. water. ClCO2Lt converts it into $C_{\bullet}H_{\bullet}.C < \stackrel{\text{N.O.}}{\underset{\text{NC,H}}{\sim}} > \text{CO.}$ [163°].

TOLYL-BENZENE v. METHYL-DIPHENYL

Tri-p-tolyl-benzene $C_{27}H_{24}$ i.e. $C_{e}H_{s}(C_{7}H_{7})_{s}$. [171°]. Formed by passing dry HCl into p-tolyl methyl ketone for some days (Claus, J. pr. [2] 41, 402). Needles. Oxidised by CrO₂ to C₂H₂(CO₂H)₃. Yields C₂,H₂Br₂ [212°].

p-TOLYL-BENZIMIDO PHENYL ETHER

[120°]. $C_6H_5.C(NC_7H_7).OPh.$ Formed from C.H. CCl:NC, H, and NaOPh (Hantzsch, B.

26, 927).

om-TOLYL-BENZOIC ACID

[1:3] C,H,Me.C,H,CO,H[1':3']. [204°]. Formed by oxidation of [1:3] C,H,Me.C,H,Me [1':8'] with dilute HNO, (Perrier, C. R. 114, 484). Silky needles, sol. alcohol, m. sol. Aq.

pp-Tolyl-benzoic acid [1:4] C.H.Me.C.H., CO.H. [1':4']. [244°]. Formed by oxidation of pp-ditolyl by CrO, and HOAc (Carnelley, J. 1877, 384). Sl. sol. hot water.— ÀgA'.

op-Tolyl-benzoic acid. [180°]. Formed by oxidation of [1:2] C.H.Me.C.H.Me [1':4'] (Carnelley, C. J. 37, 707). Insol. water, sol. hot alcohol.

TOLYL-BENZYL-AMINE v. BENZYL-TOLU-

p-TOLYL-BENZYL-ISOBUTYL-UREA

C₁₀H₂₀N₂O. [41°]. Got from p-tolyl-benzyl-carbamic chloride and isobutylamine (Hammerich, B. 25, 1824). Crystals (from ligroin).

p-TOLYL-BENZYL-CARBAMIC CHLORIDE | $(C_aH_4Me)N(CH_2Ph).COCl.$ Formed from p-tolylbenzyl-amine and COCl₂ (Hammerich, B. 25,

1822). Oil.

p-TOLYL-BENZYL-CARBINOL C₁,H₁,O i.e. C_eH₄Me.CH(OH).CH₂Ph. [66°]. (above 360°). Formed by reducing p-tolyl benzyl ketone in alcoholic solution by Na (Mann, B. 14, 1646). Small concentric needles, sol. alcohol and ether.

p-TOLYL-BENZYLENE-ACETAMIDINE

 $C_{16}H_{16}N_2$ i.e. $C_6H_4 < N_2 = \dot{C}Me^{1}$. [79°]. Formed by reduction of the acetyl derivative of o-nitrobenzyl-p-toluidine in acetic acid solution with tin and HCl (Lellmann a. Stickel, B. 19, 1610). Small glistening plates. V. sol. alcohol, ether, and chloroform.

p-TOLYL BENZYL KETONE C15H14O i.e. $C_0\hat{H}_4$ Me.CO.CH₂Ph. [109°]. (above 360°). Formed by the action of AlCl, on a mixture of phenylacetyl chloride and toluene (Mann, B. 14, 1646; Strassmann, B. 22, 1229). Plates, sol. alcohol and ether. Na added to its alcoholic solution forms the carbinol and an acid C₁₉H₂₀C₂ [93°].

Oxim [131]. Plates, insol. water. o-TOLYL BENZYL OXIDE C. H.O. i.e. [1:2] C₆H₄Me.O.CH₂Ph. (285°-290°). Oil (Staedel, B. 14, 899). Converted by HNO, (S.G. 1.5) into a tri-nitro- derivative [145°].

m-Tolyl benzyl oxide. [43°]. (300°-305°). Tables (from alcohol), v. sol. ether.

p-Tolyl benzyl oxide [41°]. Formed from C.H.Me.OK and CH, PhCl (Frische, A. 224, 154). Crystals. Converted by HNO, (S.G. 1.5) into

di-nitro-p-cresol [84°].
TOLYL-BENZYL-ISOPHOSPHINE C₁₄H₁₅P or $C_{27}H_{28}P_{2}$. [187°]. An indifferent body formed by the action of zinc on a mixture of benzyl chloride and C.H.MePCl, (Michaelis a. Gleichmann, B. 15, 1963)

p-TOLYL BENZYL SULPHONE

C₆H₄Me.SO₂.CH₂Ph. [145°]. Formed by the action of benzyl chloride on sodium toluene p-sulphinate (Otto, B. 13, 1278). Needles, sol. hot alcohol.

p-TOLYL-BENZYL-THIO-SEMICARBAZIDE C₁₈H₁₇N₂S. [121°]. Formed from benzyl thio-carbimide and p-tolyl-hydrazine (Dixon, C. J. 61, 1022). Prisms, sl. sol. cold alcohol.

C15H16N2S o-TOLYL-BENZYL-THIO-UREA i.e. C₈H₄MeNH.CS.NHCH₂Ph. [139°]. Formed by boiling o-tolyl-thiocarbimide with benzylamine in alcohol (Dixon, C. J. 59, 556). Tufts of prisms, m. sol. hot alcohol.

m-Tolyl-benzyl-thio-urea. [114°]. Formed from benzyl-thiocarbimide and m-toluidine.

Vitreous crystals, v. sl. sol. hot Aq.

p-Tolyl-benzyl-thio-urea [121°]. Crystals. p-TOLYL-BENZYL-UREA

CH,Ph.NH.CO.NHC,H,Me. [181°]. Formed from p-tolyl cyanate and benzylamine (Kühn a. Henschel, B. 21, 505).

p-Tolyl-di-benzyl-ures C₂₂H₄₂N₂O i.e. (CH₂Ph)₂N.CO.NHC₆H₄Me. [169°]. Got from C,H,),N.COCl and p-toluidine (Hammerich, B. 25, 1820). Needles (from alcohol), insol. ether.

p-Tolyl-di-benzyl-urea C22H22N2O i.e. CH₂PhN(C₆H₄Me).CO.NH.CH₂Ph. Formed from tolyl-benzyl-carbamic chloride and benzylamine (H.). Prisms, v. sol. alcohol.

Dip-tolyl-bensyl-urea C_H_N_O. [115].

Formed from p-tolyl-benzyl-carbamic chloride and p-toluidine. Needles, v. sol. hot alcohol. Di-p-tolyl-di-benzyl-urea

[93°]. $(CH_2PhN(C_sH_sMe))_2CO.$ Got from ptolyl-benzyl-carbamic chloride and p-tolylbenzyl-amine. Crystalline, sol. alcohol.

DI-p-TOLYL-BIURET (C₈H,MeNH.CO)₂NH. [c. 220°]. Formed from p-tolyl-urea and p-tolyl cyanate at 160° (Kühn a. Henschel, B. 21, 506).

White needles (from alcohol).

Tri-p-tolyl-biuret $(C_6H_4MeNH.CO)_2NC_6H_4Me.$ [156°]. Formed from di-p-tolyl-urea and p-tolyl cyanate.

p-TOLYL-BORIC ACID C₆H₄Me.B(OH)₂. [240°]. Formed by the action of water on boronp-tolyl-dichloride (v. vol. i. p. 531). Needles, sol. hot water. HgCl, gives a white pp. of C,H,HgCl.

TOLYL-BROMO-ETHYLENE v. BROMO-TOLYL-

ETHYLENE

TOLYL BROMO-METHYL KETONE

C₆H₄Me.CO.CH₂Br. [55°]. Formed by the action of hot water on di-exo-bromo-eso-methylatrolactic acid (Böttinger, B. 14, 1598). Crystals, volatile with steam, sol. water and ether.

p-TOLYL BROMO-METHYL SULPLIONE C.H.Me.SO.CH.Br. $[92^{\circ}].$ Formed by the action of bromine on an aqueous solution of C,H,SO₂,CH₂,CO₂H (Otto, J. pr. [2] 40, 544). Formed also from C,H,SO₂Na and CH₂Br₂. Thin monoclinic needles, sol. hot alcohol.

p-Tolyl di-bromo-methyl sulphone C,H,Me.SO2.CHBr2. [117°]. Formed from tolyl-sulphono-acetic acid and Br (O.). Mono-

clinic prisms, v. sol. hot alcohol.

TOLYL-BUTANE v. BUTYL-TOLUENE.

m-TOLYL-BUTYLENE C.H.Me.CH2.C3H5. (195°). Formed from C₀H₄Me.CH₂Cl, allyl iodide, and Na (Aronheim, B. 9, 1790). Oil. Yields a liquid dibromide.

p-TOLYL-p-BUTYL-PHENYL-THIO-UREA C.H.Me.NH.CS.NH.C.H., C.H., [137°]. Formed from isobutyl-phenyl-amine and p-tolyl-thiocarbimide (Mainzer, B. 16, 2024). Plates, sol. alcohol and ether.

DI-p-TOLYL-ISOBUTYL-UREA C₁₉H₂₄N₂O. [119°]. Formed from (C₂H₂)₂N.COCl and isobutylamine (Hammerich, B. 25, 1822). Prisms,

v. sol. alcohol and ether.

m-TOLYL-ISOBUTYRIC ACID

C,H,Me.CH2.CHMe.CO2H. [92°]. Formed by oxidation of m-isobutyl-toluene by dilute HNO. (Kelbe, B. 16, 620). Needles (from ligroïn).-ÀgA': pp.

Reference. - NITRO-TOLYL-ISOBUTYRIC ACID. o-TOLYL-CARBAMIC ACID C.H.NO. i.s.

C.H.Me.NH.CO.H.

Ethyl ether EtA'. [42°] (C.); [46°] (L.). Formed from o-toluidine and ClCO2Et (Cosack, B. 12, 1450; Nevile a. Winther, B. 12, 2324). Formed also from C.H.Me.CCl:NCl and alcoholic potash (Lachmann, B. 12, 1349). Tables, sol. Volatile with steam. alcohol and benzene.

Yields o-tolyl cyanate on distilling with P₂O₃.

Isobutyl ether C₄H₂A'. (275°-280°). Oil
(Mylius, B. 5, 974). Partially decomposed on distillation into tolyl cyanate and isobutyl

alcohol.

Bensyl ether CH₂PhA'. [84°]. Formed from o-tolyl cyanate and benzyl alcohol (Gattermann a. Cantzler, B. 25, 1087). Needles

Phenyl ether PhA'. [92°]. Formed by boiling di-o-tolyl-ures with di-phenyl carbonate for some hours (Eckenroth a. Rückel, B. 23, 699). Crystals (from ligroïn).

m-Phenylene ether C.H.A'2. [154°]. Formed by heating o-tolyl cyanate with resorcin

at 120° (G. a. C.). Colourless plates.

p-Phenylene ether C.H.A'2. [206·5°]. Formed by heating o-tolyl cyanate with hydroquinone at 150°.

o-Tolyl ether C,H,A'. [126°]. Formed from o-tolyl cyanate and o-cresol (G. a. C.). Needles.

 (β) -Naphthyl ether $C_{10}H_7A'$. [149°]. Formed by heating o-tolyl cyanate with (3)naphthol at 160° (G. a. C.). Plates.

Di-phenyl-ethylene ether C.H.Ph.A' [234°]. Formed by heating o-tolyl cyanate with hydrobenzoïn at 100° (Auwers, B. 24, 1778). Needles. An isomeride [163°] is obtained from isohydrobenzoïn.

m-Tolyl-carbamic acid [1:3]C,H,Me.NH.CO,H.

Ethyl ether C₁₀H₁₃NO₂. Prepared from m-toluidine and chloro-formic ether (Cosack, B. 13, 1090) Oil, v. sol. alcohol and other.

p-Tolyl-carbamic acid $[1:4]C_6H_1Me.NH.CO_2H.$

Ethyl ether EtA'. [52°]. Formed from p-toluidine and ClCO2Et (Hofmann, Pr. 19, 108; B. 3, 656). Monoclinic prisms, v. sol. alcohol (Levin, J. 1882, 384).

Phenyl ether PhA'. [115°]. Formed by distilling di-p-tolyl-urea with di-phenyl carbonate (Eckenroth a. Rückel, B. 23, 698). (from hot ligroin).

Di-p-tolyl-carbamic acid (C,H,)₂N.CO₂H.

Ethyl ether EtA'. [62°]. Formed from
(C,H₁)₂N.COCl and NaOEt (Hammerich, B. 25, 1824). Large plates, v. sol. alcohol and ether.

Chloride $(C_6H_4Me)_2N.COCI.$ Formed from COCl₂, a solution of di-p-tolylamine in benzene, and conc. NaOHAq (Hammerich, B. 25, 1821). Needles (from alcohol), v. e. sol. ether. AgCy forms the compound (C,H,),N.CO.CN(AgCy), crystallising in needles which begin to melt at 103°.

o-TOLYL-CARBAMINE [1:2]C,H,Me.N:C. (184°). S.G. 24 .968. Formed by the action of alcoholic potash on o-toluidine and chloroform (Nef, A. 270, 309). Oil, volatile with steam. When heated at 245° for 3 hours it changes to the isomeric o-toluic nitrile. Heated with S in CS2 at 130° it yields o-tolyl thiocarbimide. H2S at 100° forms C,H,Me.NH.CHS [101°].

Chloride C.H.Me.N.CCl. (2150). Formed by passing chlorine into a chloroform solution

of the carbamine.

p-Tolyl-carbamine C.H.Me.N:C. (99° at 36 mm.). Formed in like manner from p-toluidine (Nef). Oil.

Chloride C₆H₄Me.N:CCl₂. (226°). Con-

verted by p-toluidine into tri-p-tolyl-guanidine. o-TOLYL-SEMI-CARBAZIDE C₈H₁₁N₂O i.e. NH₂.CO.N₂H₂C,H₇. [160°]. Formed by heating o-tolyl-hydrazine with urea (Pinner, B. 21, 1219). Flat needles, m. sol. water.

[158°]. p-Tolyl-semi-carbazide $C_8H_{11}N_3O$. Sl. sol. cold water. Reduces Fehling's solution. Di-p-tolyl-carbaside CO(NH.NHC,H,) [201°]. Formed from p-tolyl-hydrazine and

carbamic ether (Preund, B. 24, 4197). Tables, insol. water, v. sol. hot alcohol.

TOLYL-CARBIMIDE v. TOLYL ISOCYANATE, vol. ii. p. 316.

o-TOLYL-CARBINOL C.H.O i.s.

[1:2]C₈H₁Me.CH₂OH. ω -Oxy-o-xylene. Mol.w. 122. [34°]. (223°i.V.). S.G. 42 1·02. S.1 in the cold; 1·5 at 100°. Formed by boiling ω -bromo-oxylene with KOHAq (Colson, Bl. [2] 43, 8; A. Ch. [6] 6,116). Formed also by reducing the aldehyde C_eH₄Me.CHO (Raymann, Bl. [2] 27, 498), by the action of HNO₂ on the amine C₆H₄Me.CH₂.NH₂ (Kröber, B. 23, 1028), and by reducing o-toluic amide in acid solution by sodium-amalgam (Hutchinson, B. 24, 174). Needles, v. sol. alcohol and ether, sol. hot water. Oxidised by alkaline KMnO, to o-toluic acid [103°].

m-Tolyl-carbinol [1:3]C₈H, Me.CH₂.OH. (217°). S.G. $^{\circ}$ 1.036 (C.); 17 916 (R. a. W.). S. 5 in the cold. Formed by boiling the bromide C.H.Me.CH.Br with alkalis (C.). Liquid, v. sol. alcohol and ether, sl. sol. cold water. Yields m-toluic acid [108°] on oxidation.

Ethyl ether C,H,.CH,.OEt. (202°) at 740 mm. S.G. 17 .93. Oil.

Acctyl derivative C,H,.CH,.OAc. (226°). Formed from C,H,.CH,Br and KOAc (Radziszewski a. Wispek, B. 15, 1747). Oil.

p-Tolyl-carbinol [1:4] $C_aH_aMe.CH_aOH.$ [59°]. (217°). Formed from p-toluic aldehyde and alcoholic potash (Cannizzaro, C. R. 54, 1225; A. 124, 255). Needles, sl. sol. cold water.

Ethyl ether C_sH_s.OEt. (203°) at 740 mm. S.G. ¹⁷ ·93. Formed from ω-bromo-pxylene and alcoholic potash (Radziszewski a. $m \check{W}ispek,$ B. 15, 1745). $\bar{\ }$ Oil.

Di-tolyl-carbinol (C,H,Me),CH.OH. W.); [62°] (Ador a. Crafts, B. 10, 2175). Formed by reducing di-tolyl ketone with sodiumamalgam (Weller, B. 7, 1184). Needles (from alcohol), insol. water.

Reference.—OXY-TOLYL-CARBINOL.

TOLYL-CARBINYL-AMINE METHYL-BENZYL-AMINE.

DI-p-TOLYL-CARBINYL AMINE

 $(\mathbf{C}_{7}\mathbf{H}_{7})_{2}\mathbf{C}\mathbf{H}.\mathbf{N}\mathbf{H}_{2}.$ [93°]. Formed by reducing the oxim of di-p-tolyl ketone in alcoholic solution with sodium-amalgam and HOAc at 60° (Goldschmidt a. Stöcker, B. 24, 2798). Tables. -B'HCl. [235°]. White needles.

Acetyl derivative [159°]. Needles. TOLYL-CARBINYL-UREA METHYLv. BENZYL-UREA.

Di-p-tolyl-carbinyl-urea

NH. CO.NH.CH(C,H,)2. [152°]. Formed from the hydrochloride of the amine and KCyO (Goldschmidt, B. 24, 2799). Needles.

DI-p-TOLYL-CARBONATE (C,H,O),CO. Formed, together with di-ethyl carbonate, by heating p-tolyl ethyl carbonate at 300° (Bender, B. 19, 2268). V. sl. sol. cold alcohol.

DI-TOLYL DICARBOXYLIC ACID

[3:4:1]C₄H₃Me(CO₂H).C₄H₃Me(CO₂H)[1:3:4] [over 300°]. Formed by heating its nitrile with dilute H₂SO₄ (Löwenherz, B. 25, 1036). Plates, insol. water, sl. sol. hot alcohol.

Nitrile C14H12(CN)2. [190°]. Formed from di-amido-ditolyl by Sandmeyer's diazo- reaction. Needles, m. sol. cold alcohol.

Reference.—OXY-DITOLYL DICARBOXYLIC ACID. TOLYL CHLORIDE v. a-CHLORO-XYLENE.

p-TOLYL CHLORO-BENZYL SULPHONE C.H.Me.SO.CHCl.C.H. [203°]. Formed by heating sodium toluene p-sulphinate with benzylidene chloride (Otto, J. pr. [2] 40, 519).

Needles (from HOAc).

p-TOLYL CHLORO-ETHYL SULPHONE [1:4]C_sH₄Me.SO_s.CHCl.CH₅. [48°]. Formed by CH₃.CCl₂.CO₂Na boiling boiling CH_3 . CCl_2 . CO_2 Na (1 mol.) with C_7H_7 . SO_2 Na (2 mols.) in aqueous solution. Formed also by heating ethylidene chloride (1 mol.) with (2.5 g.) with sodium toluene p-sulphinate (4.5 g.)and alcohol at 150° (Otto, J. pr. [2] 40, 515,534). Trimetric tables, v. sol. hot alcohol. Does not act upon C,H,.SO,Na even at 200°.

p-Tolyl chloro-ethyl sulphone C.H.Me.SO., CH., CH.Cl. [79°]. Formed from C.H.Me.SO., CH., CH., OH and PCl., (Otto, J. pr. [2] 30, 357). Needles or plates, v. sol. hot

alcohol.

p-TOLYL CHLORO-METHYL SULPHONE C₆H₄Me.SO₂.CH₂Cl. [84°]. Formed by heating an aqueous solution of CHCl₂.CO₂Na (1 mol.) with C,H,.SO2Na (2 mols.) on the water-bath (Otto, J. pr. [2] 40, 528). Tables (from benzene), v. sol. hot alcohol.

p-Tolyl di-chloro-methyl sulphone C₆H₄Me.SO₂.CHCl₂. [114°]. Formed by passing chlorine into a warm aqueous solution of C₇H₂,SO₂,CH₂,CO₂H (Otto, J. pr. [2] 40, 544). Crystals, insol. water, v. sol. hot alcohol.

TOLYL-CRESOL C.H.MC.CH..C.H.OH. (250°-255° at 10 mm.). Formed by heating ω chloro-xylene with phenol and zinc filings (Mazzara, G. 9, 421). Oil, sol. alcohol, ether, and alkalis. Not coloured by FeCl₃.

Acetyl derivative (250° at Decomposed in moist air into HOAc and tolyl-

cresol

p-TOLYL-CUMINYL-AMINE C₁₇H₂₁N C₆H₄Me.NH.CH₂.C₆H₄Pr. [36°]. (above 200°). Formed by the reduction by sodium-amalgam of the product of the action of cuminol on p-toluidine (Uebel, A. 245, 293). Plates or prisms.—B'HCl. Plates or needles, sl. sol. hot

Nitrosamine C₁₂H₂₂N.NO. [67°]. Prisms. p-TOLYL-CUMINYL-UBEA C₁₂H₂₂N₂O i.e. C₁₆H₁₃.NH.CO.NHC₆H₄Me. [150°]. Formed from cuminylamine and p-tolyl cyanate (Goldschmidt

a. Gessner, B. 22, 928). Needles, v. sol. alcohol. p-TOLYL-CUMYLIDENE-AMINE

C.H.Me.N:CH.C.H.Pr. [51]. Formed from cuminic aldehyde (cuminol) and p-toluidine in alcoholic solution (Uebel, A. 245, 292). Yellow-ish plates (from alcohol). Decomposed into the parent substances by heating with acids and alkalis.

p-tolyl- ψ -cumyl-urea

C.H.Me.NH.CO.NH.C.H.Mes. [218°]. Formed from ψ -cumidine and p-tolyl cyanate (Goldschmidt a. Bardach, B. 25, 1361). Needles.

C.H.Me.NH.CN. o-TOLYL-CYANAMIDE [77°]. Formed by heating C,H,NH.CS.NH.OH or by the action of hydroxylamine on o-tolyl thiocarbimide (Tiemann, B. 22, 1940; Voltmer, B. 24, 381). Tables, v. sol. alcohol and KOHAq, m. sol. water.

DI-o-TOLYL-CYANAMIDE C₁₅H₁₄N₈ C(NC,H.), Carboditolylimide. (above 500).
Formed by heating the alkyl derivatives of dio-(above 800°). tolyl-thio-urea (Will a. Bielschowski, B. 15, 1317). Amorphous, v. sol. benzene. Converted by dilute HClAq into di-o-tolyl-urea.

Di-p-tolyl-cyanamide C_{1e}H_{1,}N₂SO or C(NC,H₂)₂. [60°]. (above 230°). Formed by the action of HgO on a benzene solution of ditolyl-thio-urea (Will, B. 14, 1488). prisms, sol. benzene and ether. On heating with aniline it gives phenyl-di-tolyl-guanidine. Acids and alkalis convert it into di-p-tolyl-urea. On distillation it yields compounds melting at 49° to 60° and at 149° (Schall a. Paschkowetzky, B. 25, 2892). Phenyl-hydrazine at 190° forms C₁₆H₁₆N₆ [163°], which crystallises from alcohol in plates, and yields B'₃4HCl and B'₂2H₂PtCl₆. On heating with phenyl-hydrazine at 165° there is formed N₂PhH:C(NC,H₇)₂ [138°], orystallising in pale-red needles, yielding B'₂H₂PtCl₆ (Wessel, B. 21, 2274). Tolylene-o-diamine at 140° forms $C_{22}H_{24}N_4$ [196°], which yields B'₂3HUl [143°] (Dahm a. Gasiorowski, B. 19, 3057).
TOLYL CYANATE v. CYANIC ACID.

TOLYL CYANIDE v. Nitrile of TOLUIC ACID. TOLYL-CYANURIC ACID v. CYANIC ACIDS. TOLYLENE ALCOHOL v. DI-OXY-XYLENE.
TOLYLENE-DI-ALLYL-DI-THIO-DI-UREA

[1:2:3] C₆H₅Me(NH.CS.NHC₅H₅)₂, ϵ [152°]. Formed from c-tolylene-o-diamine and oil of mustard (Lellmann, A. 228, 246). Needles (from alcohol), decomposed on fusing into tolylenethio-urea and di-allyl-thio-urea.

Tolylene-di-allyl-di-thio-di-urea [1:3:4] C₄H₄Me(NH.CS.NHC₅H₅)₂. [150°]. Formed from *i*-tolylene-*o*-diamine and oil of mustard (Lellmann, A. 221, 24). Decomposed by heat like the preceding isomeride; melting a second time at 210°-230°

Tolylene-di-allyl-di-thio-di-urea [1:2:5] C₆H₃Me(NH.CS.NHC₅H₅)₂. [175·5°]. Formed from tolylene-p-diamine and allyl thiocarbimide (Lellmann a. Würthner, A. 228, 209).

Plates or prisms, nearly insol. ether. Tolylene-di-allyl-di-thio-di-urea. Formed from tolylene-m?-di-amine and allyl thiocarbimide. Prisms (from alcohol), insol. ether, v. sol. HOAc. Melts without decomposi-

TOLYLENE-DI-AMIDO-DI-ACETIC ETHER [1:2:4] $C_0H_0Me(NH.CH_0.CO_2Et)_2$. [70°]. Formed from tolylene-m-diamine and chloro-acetic ether (Zimmermann a. Knyrim, B. 16, 516). Needles (from water).

c-TOLYLENE - o-DIAMINE $C_1H_{10}N_2$ i.e. $H_2Me(NH_2)_2$ [1:2:3]. Mol. w. 122. [62°]. $C_6H_2Me(NH_2)_2$ [1:2:3]. Mol. w. 122. [62°]. (255°). Formed from $C_6H_2Me(NH_2)(NO_2)$ [1:2:3] by reducing with Sn and HCl (Lellmann, A. 228, 243). Reddish crystals, smelling of acetamide. -B"2HCl. Sol. water, ppd. by HCl.

c-Tolylene-m-diamine C.H.Me(NH2)2 [1:2:6]. [104°]. Formed by reduction of nitro-toluidine [92°], or of liquid di-nitro-toluene (Ullmann, B. 17, 1960). Prisms, sol. hot water. Gives a brown colour with nitrous acid. CrO, and FeCl, give a brown colour.—B'HCl: crystals, v. e. sol.

water.

i-Tolylene-m-diamine C.H.Me(NH2)2 [1:2:4]. [99°]. (c. 280°). Formed by reducing di-nitrotoluene (Hofmann, Pr. 11, 518) by reducing (4,1,2)-nitro-toluidine (Nölting a. Collin, B. 17, 268), and by the action of SnCl, on amido-toluene-azo-amido-cresol (Graeff, A. 229, 348). Needles (from water), v. sol. alcohol, ether, and

hot water. Chloroformic ether, ClCO2Et, forms [1:2:4] C₆H₂Me(NH₂)(NH₂CO₂Et) [91°], chiefly C₆H₂Me(NH₂CO₂Et) [137°] (S 4. 268, 314). Phenyl-thio-carbimide (Schiff. yields CO₂Et.NH.C₃H₃Me.NH.CS.NHPh [155°]. Hep-toic aldehyde in the cold forms C,H₃N₂(C₇H₁₄)₂, but when heated it yields an orange mass which forms a highly fluorescent solution (Schiff a. Vanni, A. 253,319). Benzoic aldehyde forms C, H_s(N:CHPh)₂ [122°-128°], while cinnamic aldehyde gives $C_7H_4(NC_9H_8)_2$ [162°] (Schiff, A. 140, 98; 239, Cl.CO.CO₂Et forms C₆H₂Me<NH.CO NH.CO [c. 260°] and [1:2:4] $C_6H_3Me(NH_2).NH.CO.CO_2Et$ [170°], which yields an acetyl derivative [192°]. Phenyl-thiocarbimide forms the compound NHPh.CS.NH.C, H, Me.NH.CO.CO, Et [155°] and C_aH₃Me NH.CS NPh [198°] (Schiff, A. 268, Tolylene-m-diamine heated with citric acid at 125° forms C₁₃H₁₂N₂O₄, crystallising from boiling alcohol in minute octahedra, decomposing at about 187° (Schneider, B. 21, 665). Acetyl-acetone at 100° forms a syrup, which when headed with H₂SO₄ yields C₁₂H₁₄N₂ [191^o] (Combes, C. R. 108, 1252).—B"HCl.—B"2HCl.—B"2HBr.—B"H₂SO₄ 2aq: prisms. S. 5-6 at 19-5 (Beilstein a. Kuhlberg, A. 158, 351).— B"H, PtCl. B"2HCyS. Prisms (Lussy, B. 7, 1265).

o-Acetyl derivative C_cH₃Me(NHAc)NH₂[1:2:4]. [140°]. Formed from C_cH₃Me(NH₂)(NO₂) [107°] by acetylation and reduction (Wallach, A. 234, 350). Needles.

-B'₂H₂PtCl_s. p-Acetyl derivative

 $C_0H_2Me(NH_2)(NHAc)$ [1:2:4]. [160°]. Formed by acetylating the diamine, and got also by reducing C₄H₂Me(NO₂)(NHAc) [1:2:4] (Tiemann, B. 3, 221; Wallach, B. 15, 2826, 2831). Long white needles.

Di-acetyl derivative C.H.Me(NHAc). [224°]. Got by boiling the base with Ac₂O or HOAc (Koch, A. 153, 132; Tiemann, B. 3, 8;

Ladenburg, B. 8, 1211). Needles.

Benzoyl derivative C.H.Me(NO2)(NHBz) [1:2:4]. [142°]. Got by reducing C.H.Me(NO.)(NHBz) (Bell, B. 7, 1505).

Di-benzoyl derivative [224°]. sl. sol. alcohol (Ruhemann, B. 14, 2656).

Thiobensoyl derivative C.H.Me(NH.)NH.CSPh. [197°] (Bernthsen a. Trompetter, B. 11, 1760).

Phthalyl derivative

C.H.Me
NH.CO
C.H. [192°]. Formed from the base and phthalic anhydride (Biedermann, B. 10, 1161). Needles, split up by dilute HCl into phthalic acid and a base CarHatNaO4.

Di-phthalyl derivative C.H.Me(NC,H,O,)2. [233°]. Crystals, insol.

water and alcohol.

s-Tolylene-m-diamine C,H,Me(NH2), [1:3:5]. (284°). Formed by reducing s-di-nitro-toluene Syrup.-B"H2SO4. -(Staedel, A. 217, 202). B"H.SnCl.: crystals, v. sol. water.

Tolylene-p-diamine C.H.Me(NH2): [1:2:5]. [64°]. (274°). Formed by reducing (5.1,2)- and (2,1,5)- nitro-toluidines (Beilstein a. Kuhlberg, A. 158, 852; Fileti a. Cross, G. 18, 806), and C.H.Me.N. C.H.Me.NH. [2:1:5] or [5:1:2] (Nietzki, B. 10, 832, 1158). Plates (from benzene), v. sol. water, alcohol, and ether. Forms toluquinone on oxidation. FeCl, added to a solution of tolylene-p-diamine hydrochloride mixed with o-toluidine gives an intense green colour.—B"2HCl. Plates.—B"H₂SO₄. S. 84 at

Di-acetyl derivative C.H.Me(NHAc). [220°]. Prisms (from dilute alcohol) (Nietzki,

B. 10, 1157; 12, 2237).

Tolylene-o-diamine $C_{s}H_{3}Me(NH_{2})_{2}$ [1:3:4]. [88.5°]. (265°). Formed by reducing (31,4)nitro-p-toluidine (Beilstein a. Kuhlberg, A. 158, 351; Graeff, A. 229, 343). Plates, m. sol. cold water. Its aqueous solution quickly blackens when exposed to air. FeCl, forms by oxidation C₁₄H₁₈N₃O, crystallising from wood spirit in brownish-red plates [247°] (O. Fischer a. Sieder, B. 23, 3802).

Reactions .- 1. Benzoic aldehyde at 140° forms $C_{21}H_{18}N_2$ or $C_7H_6 < N(C_7H_7) > CPh$ [195.5°]

(Ladenburg, B. 11, 591, 1656; Hinsberg, B. 19, 2026; 20, 1585), which yields B'MeI [209°], B'EtI [181°], and B'EtI₃ [125°], and is oxidised by KMnO₄ to an acid C_2 H₁₈N₂O₂ [254°].—2. Furfuraldehyde, acting on the hydrochloride, forms C₁₇H₁₄N₂O₂ [128·5°], which yields B'MeI [195·5°], B'MeI, [128°], and B'MeI, [109°]. — 3. Acetic aldehyde (2 mols.) added to a cold solution of the base (1 mol.) in glacial acetic acid forms

 $C_rH_e < NEt_N > CMe$ (Hinsberg).—4. Anisic alde-

hyde, added to a dilute aqueous solution of tolylene-o-diamine hydrochloride containing a little alcohol, forms C23H22N2O2 [152°-156°], which separates on addition of ammonia (L.).-5. Salicylic aldehyde at 110° forms C24H22N4O2 [106° 110°], and at 135° it forms azurine C₈₅H_{s2}N₄O_s [250.5°], which exhibits blue fluorescence in alkaline solutions .- 6. Glucose (2 mols.) added to an alcoholic solution of tolylene-o-diamine forms C₂H₆(N:C₆H₁₂O₅)₂, crystallising in satiny needles, v. sol. water [c. 160⁵], which gives a red colour Glucose converts tolylene-o-diwith FeCl₃. amine acetate into amorphous 'glycodiamidotoluene ' C₁₃H₁₈N₂O₅, m. sol. water (Hinsberg, B. 20, 495; Griess a. Harrow, B. 20, 2209). Glucoson C₆H₁₀O₆ acts in aqueous solution on phenylene-o-diamine, forming C₁₂H₁₆N₂O₄ [0. 180°] (E. Fischer, B. 22, 93).—7. Arabinose forms C₁₂H₁₆N₂O₄ [238°] (G. a. H.).—8. Acrolein in boiling aqueous solution forms tolylene-o-diamine acroson [185°] (Fischer a. Tafel, B. 22, 99).—9. Formic aldehyde forms C₁₈H₂₉N₄ [222°] (Fischer a. Wrezinski, B. 25, 2713).—10. Butyric aldehyde forms tolylene-butenyl-diamine

 $C_rH_{\bullet} < NH > CPr$ [158°], and the compound $C_{13}H_{22}N_2$ or $C_7H_6 < N(C_4H_9) > CPr$ (Hinsberg, B. 20, 1590).-11. Nitrous acid or, better, amyl nitrite acting on a salt of tolylene-o-diamine forms azimido-toluene C,H,N, [84°] (823°), which is v. sol. alcohol and hot toluene (Zincke a. Lawson, A. 240, 115; cf. Ladenburg, B. 9, 220; Bössneck, B. 19, 1759; Nölting a. Abt, B. 20, 3001). Azimido-toluene forms the salts B',H,PtCl, NaC,H,N,, Hg(C,H,N,), [285], and AgC, H.N. AcCl converts azimido-toluene into

the (3)-acetyl derivative $C_7H_6 < \stackrel{N}{N} > NAc$ [94°], while the isomeric (a)-acetyl derivative $C_7H_6 < N^{AC} > N$ [132°] is formed by the action of nitrous acid on the acetyl derivative of tolyleneo-diamine. -12. Boiling chloro-acetic ether forms oxytoluquinoxaline dihydride and a compound O₁₈H₁₆N₂O₂(?) [147°], whence HNO₂ forms a compound [248°] (Hinsberg, A. 237, 365).—13. Isatin yields C₁₅H₁₁N₃ [290°], sl. sol. alcohol (Hinsberg, A. 237, 344).—14. Equal mols. of tolylene-o-diamine and acetoacetic ether when gently warmed together eliminate H.O, and form a condensation product $C_{19}H_{18}N_2\hat{O}_2[82^\circ]$; if this is heated at 100°-120° acetic ether is evolved, and a tolyleneethenyl-o-diamine $C_0H_2Me < NH_N > C.CH_2$ is pro-

duced (Ladenburg a. Rügheimer, B. 12, 953; Witt, B. 19, 2977, 3299).—15. Chloro-acetoacetic ether forms C,H6(NH)2CMe.CHCl.CO2Et [110°], crystallising in needles (Autenrieth a. Hinsberg, B. 25, 606). -16. Oxydehydracetic acid forms C₈H₈O₅(C₆H₂Me(NH₂)₂) [147°] (Feist, B. 25, 325).—17. o-Aldehydo-benzoic acid in hot

water forms $C_0H_4Me < NH > C.C_0H_4.CO_2H [258°]$ (Bistrzycki, B. 23, 1043).—18. Di-oxy-quinone

forms di-oxy-methyl-phenazine $C_{19}H_{10}N_2O_2$ [c. 265°], which yields a di-acetyl derivative [160°]. 19. Opianic acid gives, in alcoholic solution at 0°, the acid C₆H₂Me:(N₂H):C.C₆H₂(OMe)₂.CO₂H, crystallising in needles, decomposing at 234° (Bistrzycki, B. 24, 627).—20. Di-bromo-pyruvic acid (1 mol.) forms, on boiling, C,H, N;C.OH

[235°]. Salts.—B"2HCl. Needles (Hübner, A. 209, 364).-B"H2SO, 1 aq. Scales. S. 9.29 at 19.5°.

 $-B''^2H_2C_2O_4$ aq (Hinsberg, B. 16, 1532). Compound with pyrocatechin B"C6H6O2. [78°]. Needles (from ligroin) (Merz,

B. 19, 726).

Compound with cyanogen B"Cy2 aq. [c. 244°]. Formed by passing cyanogen into an alcoholic solution of the base (Bladin, B. 18, 666). Crystals, m. sol. alcohol and ether. Converted by heating with water into NH, and two isomeric compounds C.H.N.O, one melting at 290°, the other beginning to decompose at 230°-240°. B"Cy, yields the salts B"Cy,HCl 1;aq, B"Cy,2HCl, B"Cy,H,PtCl, 2aq, minute needles, (B"Cy,)2H,PtCl, 2aq, and B"Cy,H,SO, aq: minute tables.

p-Acetyl derivative C_cH_aMe(NH₂)(NHAc) [1:3:4]. [131°]. Formed by reducing C_cH_aMe(NO_c)(NHAc) with iron and dilute HOAc (Boessneck, B. 19, 1757). Plates (from water). Converted by nitrous acid into acetyl-azimido-toluene. On distillation it yields tolylene-acetamidine $C_0H_3Me < \frac{N}{NH}$ CMe [203°]

(Niementowski, B. 25, 861).

Di-acetyl derivative C.H.Me(NAcH). [210°]. Formed by boiling the base with Ac.O (Bistrzycki, B. 23, 1878). Thin prisms (from water). Yields HOAc and tolylene-acetamidine

manner (B.). Yields tolylene-propenyl-amine [166°] on distillation.

Benzoyl derivative C_cH_sMe(NH₂)(NHBz)[1:3:4]. [194°]. Formed by reduction of C_cH₃Me(NO₂)(NHBz) with tin and HOlAq (Hübner, A. 208, 314). Crystals, split up by distillation into water and tolylenebenzamidine [240°].

Di-bensoyl derivative C.H.Me(NHBz)2. [264°]. Got by shaking tolylene-o-diamine with BzCl and NaOHAq (Hinsberg, A. 254, 255), by the action of Bz₂O on the diamine (Bistrzycki, B. 23, 1879; 24, 631), and by the action of BzCl on the benzoyl derivative (Hübner, A. 208, 314).

Needles (from HOAc).

o-Chloro-benzoyl derivative C₆H₃Me(NH₂).NH.CO.C₆H₄Cl. [153°] (Schreib, B. 13, 467). Converted by benzoyl chloride into $C_aH_aMe(NHBz).NH.CO.C_aH_aCl$ [178°].

Di-cinnamyl derivative

 $C_6H_3Me(NH.CO.C_8H_7)_2$. [206°]. Formed from tolylene-o-diamine and cinnamic anhydride (B.). Groups of small needles (from dilute alcohol).

Oxalyl derivative C₂O₂(NH.C,H_e.NH₂)₂.

[above 300°]. Obtained by the reduction of C₂O₂(NH.C,H_e.NO₂)₂ (Hinsberg, B. 15, 2691).

Small needles (from alcohol-ligroïn). On fusion it splits up into Aq and C_{1e}H₁₄N₄ [193°]_e—

Salts.—B"H₂Cl₁aq.—B"H₂PtCl_{1e}—

B"H SO, 5ca Calculator readles B"H₂SO₄5aq. Colourless needles. Phthalyl derivatives.

The compounds $C_eH_aMeN_aH_a(C_eH_aO_a)$ [104°] and $C_eH_aMe(NC_eH_aO_a)_a$ [272°] are known (Biedermann, B. 10, 1165; Ladenburg, B. 10, 1125).

Bensene-sulphonyl derivative C_sH,Me(NH₂).NH.SO₂Ph. The hydrochloride, B'HCl, is formed by the action of benzene sulphonic chloride on tolylene-o-diamine in benzene (Bistrzycki a. Cybulski, B. 24, 633; cf. Lellmann, A. 221, 18)

Phenyl-acetyl derivative C_sH_sMe(NH₂).NH.CO.CH₂Ph. [195°]. Formed from tolylene-o-diamine and phenylacetyl chloride (B. a. C.). Needles, v. sol. hot alcohol. Di-phenylacetyl derivative

C_eH₃Me(NH.CO.CH₂Ph)₂. [176°]. Needles, in-

sol. ether, v. sol. hot. alcohol. References.—Bromo-, Chloro-, and Nitro-TOLYLENE-DIAMINE.

TOLYLENE-DIAMINE SULPHINIC ACID v. DI-AMIDO-TOLUENE SULPHINIC ACID.

TOLYLENE-o-DIAMINE SULPHONIC ACID $C_0H_2Me(NH_2)_2.8O_0H$ [1:2:3:5]. Formed by reducing nitro-toluidine sulphonic acid (Nietzki a. Pollini, B. 23, 139). Needles. Forms azines with o-diketones.

c-Tolylene-m-diamine sulphonic acid

 $C_1H_{10}N_2SO_3$ i.e. $C_0H_2Me(NH_2)_2SO_3H$ [1:2:6:4]. S. 07 at 14°. Formed from toluene p-sulphonic acid by nitration and reduction (Schwanert, A. 186, 360; Marckwald, A. 274, 349). Prisms, not melted at 280°.—BaA', 4aq.—HA'HCl 2aq.— HA'HBr 2aq.—HA'HNO, aq: needles, sl. sol. alcohol.—(HA')2H2SO4 aq: plates.

Tolylene-m-diamine sulphonic acid C₄H₂Me(NH₂)₂.SO₃H [1:2:4:5]. S. ·1054 at 10°. Got by reduction of CaH2Me(NO2)(NH2).SO2H on distillation. [1:2:4:5] with SnCl_s (Limpricht a. Foth, B. 18, Di-propionyl derivative 2185; A. 230, 309). Small prisms.—KA' aq.—C₄H₂Me(NH.CO.Et)₂. [138°]. Formed in like BaA'₂Staq.—HA'HCl aq.—HA'HBr aq: prisms. Tolylene-m diamine sulphonic acid

 $C_8H_2Me(NH_2)_2.SO_3H[1:2:4:x]$. Formed tolylene-m-diamine and fuming H2SO4 (Wiesinger, B. 7, 464). Small prisms.—NaA' 4aq. KA' aq. — MgA', 5aq. — CaA', 6\frac{1}{2}aq. — SrA', 7aq. —
BaA', 6\frac{1}{2}aq. — MnA', 3aq.
TOLYLENE - DIAMINE THIOSULPHONIC

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TOLYLENE-BENZAMIDINE v. BENZENYL-TOLYLENE-DIAMINE

TOLYLENE-BENZENYL-DIAMINE v. BEN-

ZENYL-TOLYLENE-DIAMINE.

p-TOI.YLENE-BENZYL-DI-AMINE

[1:2:5] C₆H₃Me(NH₂).NHCH₂Ph. Got by reducing p-nitroso-benzyl-o-toluidine by alcoholic ammonium sulphide (Boeddinghaus, A. 263, 309).—B"2HCl. Needles, sl. sol. alcohol.

TOLYLENE BLUE C₁₃H₁₈N₄HCl. Formed by

mixing solutions of nitroso-dimethyl-aniline hydrochloride and (1,2,4)-tolylene-m-diamine (Witt, C. J. 35, 358; B. 12, 931). Prisms (containing aq) with coppery lustre, forming blue solutions in water and alcohol. Acids turn the solutions reddish-brown. Alkalis pp. the base as a tarry mass. Tin and HClAq reduce it to the leuco-base, which forms a deliquescent hydrochloride and crystalline (C₁₅H₂₀N₄)HSnCl₃. On boiling tolylene-blue with water for some time part is reduced to leuco-tolylene blue and part oxidised to tolylene red C₁₅H₁₆N₁. Tolylene blue heated in aqueous solution containing HOAc for 12 hours at 40° forms tolylene violet C₁₄H₁₄N₄, characterised by an insoluble sulphate. Hydrated tolylene-violet C14H14N4 aq is a scarlet powder which dissolves in alcohol and in ether, forming solutions with orange fluorescence.

Tolylene red $\begin{bmatrix} 1_4^3 \end{bmatrix} C_6 H_3 (NMe_2) < N \\ N > C_6 H_2 Me (NH_2) \begin{bmatrix} 3 \\ 4 \\ \vdots \\ 6 \\ 1 \end{bmatrix}$

(Bernthsen a. Schweitzer, A.236, 332; Andresen, B. 19, 2217). Orange-red needles (containing 4aq). Forms a red fluorescent solution in alcohol. Its neutral salts are red, its acid salts are blue. By elimination of NH, by the diazoreaction it is converted into di-methyl-amidomethyl-phenazine.

Isomeride of tolylene red

 $\begin{bmatrix} 1_4^3 \end{bmatrix} C_0 H_3 (NMe_2) < \stackrel{N}{\dot{N}} > C_6 H_2 Me (NH_2) \begin{bmatrix} 3 \\ 4^5 : 1 \end{bmatrix}.$

Formed from chloro-di-nitro-toluene and dimethyl-p-phenylene-diamine and reduction of the product (Witt, B. 25, 3008). Coppery needles, sol. warm water.

TOLYLENE-BUTENYL-DIAMINE v. Bu-

TENYL-TOLYLENE-DIAMINE.

TOLYLENE-DICARBAMIC ETHER v. i-

TOLYLENE-m-DIAMINE.

TOLYLENE DIVIANGE: [94°]. Formed from tolylene m-diamine and COCl₂ (Snape, C. J. 49, Social Sol. ether. When heated with phenol at 130° it yields C.H.Me(NH.CO.Ph)2 [147.5°] crystallising in needles.

TOLYLENE-ETHENYL-DIAMINE C.H. 10 N.2 i.e. C_eH₃Me NH CMe. [203°] (Niementowski, B. 25, 861). (350°) (Nölting a. Witt, B. 17, 81). Formed by boiling (1,3,4)-tolylene-o-diamine with HOAc (Ladenburg, B. 8, 677) by reducing the acetyl derivative of nitro-p-toluidine (Hobrecker, B. 5, 920), and by adding aldehyde to a solution of tolylene-o-diamine in very dilute HOAc (Hinsberg, B. 20, 1589). Tables (from water). Forms with acetone and SO, the compound $(C_sH_{10}N_2C_3H_6O)SO_4$ (Boessneck, B. 21, 1909). $-B'_2H_2PtCl_s$.—B'HNO₄.

TOLYLENE-ETHENYL-ETHYL-DIAMINE

C₁₁H₁₄N₂ i.s. CMe:CH.C.N = CMe. CH.C.N Et CMe. [166°]. Formed from [1:3:4] C.H.Me(NH2).NHEt and Ac.O (O. Fischer, B. 26, 200).

Tolylene-ethenyl-ethyl-diamine

CMe:CH.C.NEt CMe. [93°]. Got by ethylation of tolylene-ethenyl-diamine (Hübner, A. 210, 351). Formed also by the action of aldehyde (2 mols.) on tolyene-o-diamine (1 mol.) (Hinsberg, B. 20, 1585) and by heating $C_{\rm e}H_{\rm 3}{\rm Me(NO_2)NEtAc}$ with zinc-dust and HOAc (Niementowski, B. 20, 1884). Needles (containing 3aq). Combines with ethyl iodide, forming B'EtI and B'EtI, [111°], which yield B'EtOH and BEt₂PtCl_a. Salts.—KHI aq [143°].—B'HNO, aq. [99°]. Melts at 95° when anhydrous.—B'C₆H₂N₂O₅. Crystals, sl. sol. alcohol.

TOLYLENE-ETHYL-DIAMINE

CMe:CH.C.NH, [55°]. Formed by reduc-CH :CH.C.NHEt tion of nitro-ethyl-p-toluidine (Gattermann, B. 18, 1484; Fischer, B. 26, 199). White plates, quickly blackened by light and air. Rotates on

Reactions.-1. Bisulphide of carbon yields $C_6H_3Me < NE_t > C.SH$ [139°].—2. Ac₂O forms tolylene-ethenyl-ethyl-diamine [166°].—8. Benzoic aldehyde yields $C_6H_2Me < N_{NEt} > CPh$; while o- and p-nitro- and o-oxy- benzoic aldehydes from corresponding bodies [170°], [176°], and [78°].

Salts. - B'HCl. [176°]. - B'2H2C2O4. [151°].

Tolylene-ethyl-diamine

C₆H₂Me(NHEt).NH₂ [1:2:5]. (264° cor.). Formed by reducing nitroso-ethyl-o-toluidine (Koch, A. 243, 307). Oil, v. sol. ether.—B"2HCl. [124°]. Crystals, insol. ether.

Tolylene-ethyl-diamine

 $C_sH_sMe(NH_2)(NHEt)$ [1:2:4]. $(282^{\circ}$ uncor.). Formed by reduction of nitro-ethyl-p-toluidine (Nölting a. Stricker, B. 19, 549). Oil. Gives the reactions characteristic of m-diamines.

Tolylene-di-ethyl-diamine

C.H.Me(NHEt), [1:3:4]. (Hinsberg, A. 265, 191). (265° uncor.). Oil Turns black in air.

Tolylene-di-ethyl-diamine

 $C_{e}H_{s}Me(NEt_{z})(NH_{z})$ [1:2:5]. (240° i.V.). Formed by reducing nitro-di-ethyl-o-toluidine (Bernthsen, B. 25, 3138). Oil. FeCl, slowly gives a purple colour in an aqueous solution of the sulphate. Na₂S₂O₃ and K₂Cr₂O₇ added to its solution in HOAC slowly ppt. SO,H.S.C,H.Me(NEt,)(NH,) [210°-215°].—B'H.SO,L Large colourless tables. DI-TOLYLENE-ETHYLENE-TETRA-AMINE

v. Ethylene-di-tolylene-tetra-amine. TOLYLENE-DI-ETHYL-DI-THIO-DI-UREA [1:3:4] MeC₆H₂(NH.CS.NHEt)₂. [149°]. Got from ethyl mustard oil (5.5 g.) and tolylene-diamine (3 g.) in alcohol (15 c.c.) (Lellmann, A. 221, 23). V. sol. alcohol, more sol. HOAc, sl. sol. water. Decomposes when melted into tolylenethio-urea, Me.C.H. (NH) CS, and CS(NHEt)

(1.2.4)-Isomeride [225°]. Formed from tolylene-di-thio-di-urea and EtI at 105° (Lussy, B. 8, 668). Crystals.

TOLYLENE-DI-ETHYL-DI-UREA

[1:2:4]C₆H₃Me(NH.CO.NHEt)₂. [175°]. Formed by heating tolylene-urea with Etl at 110° (Lussy, B. 8, 292). Crystals, v. sol. alcohol and ether.

TOLYLENE HYDRATE v. PHENYL-BENZYL-

DI-TOLYLENE KETONE OXIDE

 $\begin{bmatrix} 1_4^3 \end{bmatrix} C_s H_s Me < \begin{matrix} 0 \\ CO \end{matrix} > C_s H_s Me \begin{bmatrix} 3 \\ 4 \end{bmatrix}$. ୮166°7.

Formed by heating oxytoluic acid with Ac2O (Weber, B. 25, 1745). White flakes, sol. alcohol and ether.

MERCAPTAN C₆H₈Me(SH)₂ 7°]. (263°) TOLYLENE [37°]. Dithiocresorcin. [1:2:4]. (Klason, B. 20, 355).

Di-tolylene mercaptan

HS.C.H.Me.C.H.Me.SH. [113°]. Formed from the diazo- compound of di-amido-o-ditolyl by combination with potassium xanthate at 70°-75° and saponification of the resulting oil with alcoholic potash (Leuckart, J. pr. [2] 41, 214). Yellowish plates, v. sol. alcohol and ether.

Methyl ether [118°]. Needles. TOLYLENE METHENYL-AMIDINE C.H.N.

i.e. $C_eH_2Me < NH > CH$. [114°]. Formed by boiling from (1,3,4)-tolylene-diamine with formic acid (Ladenburg, B. 10, 1123; O. Fischer, B. 22, 644).-B'2H2PtCl. Yellow prisms.

TOLYLENE-METHENYL-DÎAMINE v. METHENYL-TOLYLENE-DIAMINE and the preceding

TOLYL-METHENYL - METHYL - DIAMINE

C₀H₁₀N₂ i.e. CMe:CH.C.N CH:CH.C.NMe CH. [94°].

Formed by heating C₇H₈(NH₂).NHMe with formic acid (O. Fischer, B. 26, 195).

Isomeride CMe:CH.C.NMe CH. Formed by heating tolylene-methenyl-amidine with MeI (O. Fischer, B. 22, 644). Formed also from (1,3,4)-tolylene-diamine and formic aldehyde (Fischer a. Wreszinski, B. 25, 2711). Oil. -B'HI aq.—B'HCl. Prisms (from HClAq).
TOLYLENE-METHYL-DIAMINE

[1:3:4] C_eH₃Me(NH₂).NHMe. [44°]. Formed by reducing nitro-methyl-p-toluidine (Gattermann, B. 18, 1487; O. Fischer, B. 26, 194). Formed also by heating methyl-p-amido-toluene-azobenzene sulphonic acid with SnCl₂ and HClAq (Bamberger a. Wulz, B. 24, 2082). Four-sided tables.

Reactions. - 1. Acetic anhydride yields $C_1H_1 < \frac{NH}{NMe} > CMe [142^\circ].$ —2. CS_2 reacts, forming C,He NH CS [194°].—3. Benzoic aldehyde forms C,H, NMe CPh [127°], while o-nitroand o-oxy-benzoic aldehyde form corresponding bodies [158°] and [180°]. Salts.—B'HCl. [175°-180°]. Plates (from

alcohol).-B'H2C2O4. [124°].—B'C.H.N.O..

[164°], Tolylene-di-methyl-diamine [28]. (270°). $C_0H_2Me(NMe_2)(NH_2)$ [1:5:2]. nitroso-di-methyl-m-Formed by reducing tolnidine (Wurster a. Riedel, B. 12, 1801; 18,

126). Needles or prisms, v. sol. water, alcohol, and ether. Yields toluquinone on oxidation.

Acetyl derivative [158°].—

B'2H2PtCl, 4aq.

Tolylene-di-methyl-diamine $C_eH_2Me(NMe_2)(NH_2)$ [1:2:5]. [47°]. (240° i.V.). Formed by reducing nitro-di-methyl-o-toluidine (Bernthsen, B. 25, 3134). FeCl, gives an intense bluish-red colour in neutral solutions. Na₂S₂O₂ and K2Cr2O, added to the acetic acid solution ppt. $C_{e}H_{2}Me(NMe_{2})(NH_{2})S.SO_{e}H$ [c. 240°]. Salt.-B'2H2SO4. Needles, v. e. sol. water.

Tolylene-tetra-methyl-diamine $C_8H_3Me(NMe_2)_2$ [1:2:5]. (c. 260°). Formed by heating the preceding body with MeOH and HCl at 180° (W. a. R.). Liquid. Coloured blue in

aqueous solution by FeCl.

Methylo-iodide B'MeI. [160°]. Needles.

Tolylene-tetra-methyl-diamine $C_6H_3Me(NMe_2)_2[1:3:4]$. (226°) at 717 mm. Got by methylation (Niementowski, B. 20, 1888). Liquid, v. sl. sol. water. FeCl, at 40° to 50° gives a reddish-brown colour. HNO, added to its solution in H_2SO_4 gives a red colouration.— B'₂ H_2 PtCl₆.—B'HHgCl₈.—B'C₆ H_2 N₂O_{7-C} Yellow tables.

Reference.-NITRO - TOLYLENE - TETRA-METHYL-DIAMINE.

TOLYLENE-METHYL-ETHENYL-AMIDINE $\begin{array}{c} \text{CMe:CH.C.N} \\ \text{CH:CH.C.NMe} \\ \end{array} \hspace{-0.5cm} \hspace{-0$ C₁₀H₁₂N₂ i.e. Formed, together with C₁₀H₁₂N₂O [163°], which crystallises with 2aq, by reducing the compound C₂H₂Me(NO₂).MeAc (Niementowski, B. 20, 1878), and by acting on tolylene-methyl-diamine with Ac₂O (O. Fischer, B. 26, 196). Needles (from hot water).—B'₂H₂PtCl₂ dried at 100°). $[234^{\circ}-244^{\circ}]$.—B'HCl $\frac{1}{2}$ aq.—B'MeL $[221^{\circ}]$. B'MeOH. [115°-135°].

TOLYLENE-TETRA-METHYL DI-PYRROLE TETRACARBOXYLIC ACID C₂₃H₂₂N₂O₈ i.e. [1:2:4] C_8H_3Me $(N < \frac{CMe:C.CO_2H}{CMe:C.CO_2H})_s$. [248°].

sol. hot alcohol and ether.

Ethyl ether Et, A'v. Formed by heating di-acetyl-succinic ether with tolylene-m-diamine and HÖAc at 150° (Knorr, A. 236, 314). TOLYLENE-METHYL-THIO-UREA

CMe:CH.C.NH CH:CH.C.NMe>CS. [194°]. Formed from $C_7H_6(NH_2)(NHMe)$ and CS_2 (Fischer, B. 26, 196). Needles (from alcohol).

TOLYLENE-DI-OXAMIC ACID C11H10N2O. i.e. C₆H₃Me(NH.CO.CO₂H)₂ [1:2:4]. Formed, together with the amide, by the action of alcoholic NH, on the ether. Crystalline. Very sweet. Decomposed by warm KOHAq into oxalic acid and tolylene-

diamine.—Ag.A".—PbA".—BaA" 2aq.
Di-amide C.H.Me(NH.CO.CONH2)2[1:2:4]. Formed from the ether and alcoholic ammonia (Schiff, A. 268, 3-3). White powder, m. sol.

alcohol.

Ethyl ether [1:2:4] C.H.Me(NH.CO.CO.Et), [130°]. Formed by boiling C.H.Me(NH.).NH.CO.CO.Et with alcohol and oxalic ether (Schiff a. Vanni, A. 268, 840). Small needles (from ether).

Amide - ether $C_tH_tMe < NH.CO.CO.NH_t$ [1: $\frac{2}{4}$] [210°]. Formed from C.H.Me(NH.).NH.CO.CO.NH, by boiling

with alcohol and oxalic ether (Schiff, A. 268, 841).

Amide ether

*C₆H₂Me NH.CO.CONH₂[1²] [c. 220°]. Got by fusing C₆H₂Me(NH₂).NH.CO.CO₂Et with oxamic ether.

TOLYLENE-OXAMIDE C.H.Me NH C.O.

[c. 260°]. Formed from (1,2,4)-tolylene-diamine and chloro-glyoxylic ether, and also by heating tolylene-diamine oxalate to 160° (Schiff a. Vanni, A. 268, 312).

TOLYLENE-PENTENYL-DIAMINE

[13] C₆H₃Me<N_{NH}>C.C₄H₉. [146°]. Formed by reducing the valeryl derivative of nitro-p-toludine (Friederici, B. 11, 1974; Hübner, A. 209, 365). Prisms (from ligroïn), v. e. sol. alcohol.

TOLYLENE - DI - PHÉNYL - DI - AMIDO-METHYLENE-DIAMINE C20H20N4 i.e.

 $C_8H_8Me < \frac{NH}{NH} > C(NHPh)_2$. [161°]. Formed from $C(NPh)_2$ and (1,3,4)-tolylene-diamine at 130°–149° (Dahm a. Gasiorowski, B. 19,3057). Needles (from benzene), v. sol. alcohol.— B'_2 3HCl. [174°].— $B'H_2SO_4$. Plates, v. sol. water and alcohol. TOLYLENE-PROPENYL-DIAMINE

C_uH_xMe NH CEt. [166°]. Formed by distilling the dipropionyl derivative of tolylene-odiamine (Bistrzycki, B. 23, 1879). Needles, m. sol. water, almost insol. ligroin.

TOLYLENE RED v. Tolylene blue. DI-TOLYLENE DISULPHIDE

CMe:CH.C.S.C.CH:CH.

CH:CH.C.S.C.CH:CMe. [116°]. Formed by the action of heat at 200°-250° on tolylene diazosulphide C₀H₃Me<\(\frac{N}{S}\)>N [43°], obtained from ethenyl-amido-tolyl mercaptan and HNO₂ (Jacobsen a. Ney, B. 22, 911). Conc. H₂SO₄ gives an intense blue colour.

TOLYLENE . DI - THIO - DI - CARBAMIC ETHER [1:3:4] C_oH₃Me(N:C(SH).OEt)₂ [120°]. Formed by boiling tolylene-di-thiocarbimide with alcohol (Billeter a. Steiner, B. 20, 230). Plates (from benzene). Yields a yellowish-white silver salt.

TOLYLENE - DI - THIOCARBIMIDE [1:2:4] C₄H₃Me(N.CS)₂. [56°]. (c. 300°). Formed from tolylene-m-diamine and CSCl₂, and also by heating tolylene-di-thio-di-urea with conc. HClAq (Billeter a. Steiner, B. 18, 3292; 20, 230). Needles. Converted by NH,Aq into tolylene-di-thio-di-urea [206°], by aniline into di-phenyltolylene-di-thio-di-urea [168°]. Copper at 250° produces C₄H₃MeCy₂ [141°].

Tolylene-di-thiocarbimide [1:3:4]

C₃H₂Me(N.CS)₂. [42°]. Formed from an aqueous solution of tolylene-o-diamine hydrochloride and CSCl₂ in CHCl₂ (B. a. S.). Plates. Converted by HCl at 200° into o-tolylene-thio-urea.

TOLYLENE-DI-THIO-DIGLYCOLLIC ACID C.H.Me(S.CH.CO.H.). [152°]. Formed from thio-orein °C.H.Me(SH), [35°], chloro-acetic acid, and NaOHAq (Gabriel, B. 12, 1640). Needles, m. sol. hot Aq.

TOLYLENE THIO-UREA C.H.Me NH CS.
Formed from C.H.Me(NH,HCl): [1:2:3] and

excess of NH₄SCN by boiling with water, evaporating, and heating to 120° (Lellmann, A. 228, 245). Small red crystals (from alcohol). Does not melt below 326°; may be sublimed. Insol. water, sl. sol. alcohol, dissolved by warm NaOH; re-ppd. by HCl. When boiled with lead acctate and excess of NaOH, no PbS is formed.

Tolylene - thio - urea C_sH₂Me NH CS. [284°]. Formed by heating (1,3,4)-tolylene-odiamine sulphocyanide at 180° (Lellmann, A. 221, 10]. Formed also by the action of alcoholic NH, on o-tolylene di-thiocarbimide (Billeter a. Steiner, B. 20, 231). Silvery plates, sol. alcohol, HOAc, and cold NaOHAq, sl. sol. water and CHCl₂.

m-Tolylene - thio - urea $C_sH_3Me < \stackrel{NH}{NH} > CS$. [149°]. Formed from (1,2,4)-tolylene-diamine and alcoholic CS_2 (Lussy, B. 8, 293). Orystalline powder, v. e. sol. alcohol.

m-Tolylene-di-thio-di-urea [1:2:4] C_bH₂Me(NH.CS.NH₂)₂. [218°] (L.); [206°] (B. a. S.). Formed by heating tolylene-m-diamine sulphocyanide (Lussy, B. 7, 1265; Gebhardt, B. 17, 3046; Billeter a. Steiner, B. 18, 3293; 20, 228). Crystalline powder, insol. water and ether, nearly insol. alcohol, m. sol. hot HOAc.

TOLYLENE-TOLENYL-AMIDINE $C_{16}H_{14}N_{2}$ i.e. $C_{6}H_{3}Me < {}^{NH}_{N} > C.C_{6}H_{4}Me$. Formed by reducing [1:4] $C_{6}H_{4}Me$. CO.NH. $C_{6}H_{3}(NO_{2})Me$ [4:3:1] (Hübner, A. 210, 331). Needles, sl. sol. water.—B'HNC₃.—B'₂H₂SO₄. Crystals, sl. sol. water.

TOLYLENE - DI - p - TOLYL - DI - AMIDO - METHYLENE-DIAMINE C22H24N4 i.e.

C₆H₃Me < NH < C(NH.C₆H₄Me)₂. [196°].

Formed by heating C(NC₈H₄Me)₂ with (1,3,4)-tolylene-diamine (Dahm a. Gasiorowski, B. 19, 3059). Needles (from alcohol).—B'₂3HCl. [143°]. Deliquescent needles, v. sol. water.

TOLYLENE-TOLYL-DIAMINE $C_{14}H_{16}N_{2}$ i.e. [1:3:4] $C_8H_3Me(NH_2)NHC_6H_4Me[1:4]$. Amido-dip-tolyl-amine. [109°]. Formed by reducing nitro-di-p-tolylamine (O. Fischer, B. 23, 8798; 26, 187). Prisms (from ligroin), turning brownish-red in air. H_2SO_4 forms a blue solution changing to green. FeCl₂ forms $C_{28}H_2N_3O$ [188°] crystallising in red plates, yielding $B_2^2H_2PtCl_6$. Boiling with acetic anhydride and sodium acetate produces $C_7H_6 < \frac{N(C_7H_7)}{N} > CMe$ [95°], which forms $B_2^2H_2PtCl_6$. Benzoic aldehyde forms $C_7H_6 < \frac{N(C_7H_7)}{NH} > CHPh$ [156°], which yields $(C_{21}H_{28}N_2)_2H_2PtCl_6$ and $C_{21}H_{20}N_2HAUCl_4$.

"Salts.—B'H,Co. Nearly insol. cold Aq.—B'C.H.N.O. Brownish-red crystals.

Acetyl derivative [126°]. Prisms.

Tolylene-p-tolyl-diamine [1:4:3] C_sH₂Me(NH₂).NHC_sH₄Me [1:4]. [1077]. Formed from toluene-azo-toluene or s-di-p-tolyl-hydrazine (hydrazo-toluene) in alcoholic solution by the action of SnCl₂ and HCl (Täuber, B. 25, 1022; cf. Melms, B. 3, 554; Goldschmidt, B. 11, 1626). Plates, v. e. sol. alcohol. NaNO₂ colours its solution in H₂SO₄ deep blue (?). Benzoic aldehyde reacts, forming

 $C_7H_6 < N(C_7H_7) > CPh$ [165°] (cf. Lellmann, B. 15, 832). Benzil in alcohol and HCl forms C28H24N2O [173°]. Ethyl nitrite and H2SO4 added to its alcoholic solution form the azimide $C_7H_6 < N^{(C_7H_7)} > N$ [93°]. On oxidation together with p-toluidine it forms the ditoluide of amidotoluquinone (Green, C. J. 63, 1408).

Tolylene-di-p-tolyl-diamine [1:2:5] $C_bH_bMe(NHC_1H_1)_2$. [118°]. Formed by heating hydrotoluquinone (40 g.) with p-toluidine (160 g.) and ZnCl₂ (80 g.) to 200°–280° for 5 hours (Green, C. J. 63, 1408). Plates, sol. HOAc, insol.

TOLYLENE-p-TOLYL-GUANIDINE

C_eH_sMe</br>
NH
C:NC₇H₇. [198°]. Formed from tolylene-o-diamine and C(NC,H,)2 (Keller, B. 24, 2518). Prisms (from alcohol), or tables (from benzene). Phenyl cyanate forms the compound $C_7H_6 < N > C:NC_7H_7CO > NPh$ [233°], crystallising from hot alcohol in needles.—B'HCl.-B'2H2PtCls.—B'2H2SO45aq. Needles, sl. sol. cold

Acetyl derivative C,H, < NAC > C:NC,H,. [149°]. Needles, v. sol. alcohol.

Di-benzoyl derivative [201°]. Needles. Nitrosamine C,H,:N,H(NO):C:NC,H,. [c. 140°]. Decomposed on fusion.

TOLYLEN E-p-TOLYL-THIO-UREA

CH :CH.C.N(C,H,) >CS. [270°]. Formed from tolylene-tolyl-diamine and CS₂ in alcohol (Fischer a. Sieder, B. 23, 3799). Prisms, insol. water, sl. sol. ligroin. Conc. H2SO, forms a green solution.

TOLYLENE-UREA C,H,N2O i.e.

 $[1:\frac{3}{4}]C_6H_3Me < NH > CO.$ [292°]. Formed by heating tolylene-o-diamine with urea and got also by the action of HClAq at 140° on $C_eH_aMe < NH > C.OEt [163°]$, which is formed by the action of HN:C(OEt)2 on tolylene-o-diamine hydrochloride (Sandmeyer, B. 19, 2651). Formed also by heating C,H,(NH2).NH.CO.NPh2 (Lellmann a. Bonhöffer, B. 20, 2124) or the compound C,H,(NH2).NH.CO.NHPh (Leuckart, J. pr. [2] 41, 324, who states that it melts above 300°), and by heating tolylene-o-diamine in benzene with COCl₂ in toluene in sealed tubes at 100° (Hartmann, B. 23, 1048). Needles, sl. sol. hot

water.—B'HCl: prisms.

TOLYLENE - DI - UREA C.H., N. O. i.e.
[1:3:4] C.H., Me(NH.CO.NH.). [282°]. Got from tolylene-diamine hydrochloride and potassium eyanate in aqueous solution (Lellmann, A. 221, 14). Needles, sl. sol. water and chlo oform, sol. alcohol and hot conc. HCl; v. sol. HOAc.

Tolylene-di-urea $C_9H_{12}N_4O_2$ i.e. [1:2:4] C₂H₂Me(NH.CO.NH₂)₂. [220°]. Formed from tolylene-m-diamine sulphate and KCyO (Strauss, A. 148, 157). Formed also from tolylene cyanate and ammonia (Lussy, B. 8, 291). Scales, sl. sol. alcohol and hot water.—B"2HCL

TOLYLENE-VIOLET v. Tolylene blue.

s-DI-TOLYL-ETHANE C.H.Me.CH. CH. CH. Me. (296°). Formed from C₆H₄Me.CH₂Cl and sodium (Vollrath, Z. 1866,

489). Oil.
u-Di-p-tolyl-ethane $CH_3.CH(C_6H_4Me)_2$. (295°). S.G. $\frac{20}{4}$ 974. Formed by shaking paraldehyde with toluene and H,SO, at 0° (O. Fischer, B. 7, 1191). Formed also by heating di-a-tolylpropionic acid with lime (Haiss, B. 15, 1476) and by the action of ethylidene chloride and AlCl₃ on toluene (Anschütz, B. 18, 664; A. 235, 315). Oil. Yields toluyl-benzoic acid and di-tolyl ketone on oxidation.

s-mp-Di-tolyl-ethane $C_2H_1(C_1H_1)_2$. (297°–300°). Formed from ethylene bromide, toluene, and AlCl₂ (Friedel a. Balsohn, Bl. [2] 35, 52; A. Ch. [6] 1, 487). Yields isophthalic and

terephthalic acids on oxidation.

References .- AMIDO-, CHLORO-, TRI-CHLORO-, DI-NITRO-, and OXY-TOLYL-, ETHANES.

TOLYL ETHER v. DI-TOLYL OXIDE. o-TOLYL-ETHYL ALCOHOL. Hexa

hydride $CH_2 < CH_2 \cdot CHMe > CH.CHMe.OH.$ (195°-200°). Formed, together with the tetrahydride CH₂ CH₂·CMe C.CHMe.OH (142° at 50 mm.) by reducing C,H,Me.CO.CH, in moist ethereal solution by sodium (Kipping a. Perkin, jun., C. J. 57, 22). Oil. Yields an acetyl derivative (204°-208°).

p - TOLYL - ETHYL - m - AMIDO - PHENOL. Ethylderivative C,H,.NEt.C,H,.OEt. Formed from C,H,.NH.C,H,.OH, potash, and EtI

(Hatschek a. Zega, J. pr. [2] 33, 217). Oil. p-Tolyl-ethyl-p-amido-phenol. Ethyl derivative C₁,H₂₁NO i.e. C,H₂,NEt.C₂H₄.OEt. (c. 340°). From C,H,.NH.C,H,.OH, potash, and EtI (H. a. Z.). Oil.

TOLYL-ETHYL-AMINE v. ETHYL-TOLUIDINE. Di-tolyl-ethyl-amine (C_uH₄Me)₂NEt. (255°-260° at 20 mm.). Formed by heating di-p-tolylamine with EtOH and HCl at 250°-280° (Girard, Bl. [2] 24, 120). Oil.

o-TOLYL ETHYL CARBONATE C, H, O, i.e. [1:2] C_eH₄Me.O.CO.OEt. (236°). Formed from sodium o-cresol and ClCO₂Et (Bender, B. 13, 700).

m-Tolyl ethyl carbonate. (246°). Formed in like manner from m-cresol. Oil.

p-Tolyl ethyl carbonate. (245°). Oil. By long heating at 300° it yields di-p-tolyl carbonate and di-ethyl carbonate (Bender, B. 19, 2268).

DI - p - TOLYL - ETHYLENE C18H18 C.H.Me.CH:CH.C.H.Me. Di-methyl-stilbene. [179°]. (above 300°). Formed by distilling (C,H,Me)2CH.CCl, with zinc-dust (Goldschmiedt a. Hepp, B. 6, 1504; Elbs a. Förster, J. pr. [2] 39, 300), and by heating di-p-tolyl fumarate, CO₂ being evolved (Anschütz, B. 18, 1948). Plates, v. sol. CHCl₂. Yields a dibromide [204°].

v. sol. CHCl₂. Yields a dibromide [204]. u-Di-tolyl-ethylene CH₂:C(C₈H₄Me)₂. (305°). Formed by the action of alcoholic potash on CH₂Cl.CH(C,H₇)₂ (Hepp, B. 7, 1413). Oil. Yields di-tolyl-ketone [94°] on oxidation.

Tetra-tolyl-ethylene C₃₀H_{2s} i.e. (C,H₁)₂C:C(C,H₁)₂. [215°]. A product of the action of chloroform and AlCl₂ on toluene (Schwarz, B. 14, 1529). Plates, with greenishyellow fluorescence, sol. benzene.

References .- Bromo- and DI-CHLORO- TOLYL-

ETHYLENE.

o-TOLYL-ETHYLENE-DIAMINE

C,H,NH.C₂H₄,NH₂. (c. 267°). Formed from o-tolylamido-ethyl-phthalimide and HCl (Newman, B. 24, 2194). Sol. water and alcohol.-B"2HCl. [168°-173°].—B"2C₆H₃N₃O₇. [148°]. Green needles.

Di-benzoyl derivative [164.5°]. Needles. p-Tolyl-ethylene-diamine. Formed in like Liquid. -B"2HCl. manner (N.). Liquid. -B"
B"H₂PtCl₆. Yellowish plates. [218°].-

Di-acetyl derivative [107°]. Needles. Di-bensoyl derivative [161°]. Cubes.

C16H20N2 i.e. Di-o-tolyl-ethylene-diamine $C_{1e}H_{20}N_2$ i.e. $C_7H_7NH.CH_2.CH_2.NHC_7H_7$. [71°]. S. 3 at 100°. S. (cold alcohol) 9. S. (ether) 14. Formed by the action of o-toluidine on ethylene bromide in presence of Na_2CO_3 (Bischoff, B. 23, 1982, 2031; 25, 3257; cf. Mauthner a. Suida, M. 7, 230; Colson, Bl. [2] 48, 799). Plates (from ligroin), v. sol. alcohol. Yields indole on heating with zinc-dust. CH2Cl.CO2H and NaOAc from ditolyl-pyrazine hexahydride [154°]. -B"2HCl.-B'H.PtCl. —B"H.SO, (dried at 100°).—B"2HBr. [222°]. S. 3 at 100°.

Acetyl derivative C₂H₄(NAc.C₇H₇)₂. [153°].

Bromo-acetyl derivative

 $C_2H_4(NC_7H_1.CO.CH_2Br)_2$. [205°]. Sl. sol. cold HOAc and ligroin.

Bromo-propionyl derivative C₂₂H₂₈N₂Br₂O₂. [181°]. Bromo-n-butyryl derivative

 $(C_{2i}H_{30}N_2Br_2O_2)C_eH_e$ (from benzene). [190°]. Bromo-isobutyryl derivatives

 $C_7H_7NH_7.CO.CB_7Me_2$ [137°]

C₂₁H₃₀N₂Br₂O₂ [173°].
Di-p-tolyl-ethylene-diamine [97·5°]. Formed by heating p-toluidine with ethylene bromide at 150° (Gretillat, M. S. [3] 3, 383; Bischoff, B. 25, 3260). Crystals, v. e. sol. alcohol. - Hydrobromide [255°].

Acetyl derivative C,H,(NAc.C,H,).

[139°].

Bromo-acetyl derivative C20H22N2Br2O2. [196°].

Bromo-propionyl derivative. [182°]. Bromo-n-butyryl derivative $\mathbf{C}_{2i}\mathbf{H}_{30}\mathbf{N}_{2}\mathbf{Br}_{2}\mathbf{O}_{2}$. [125°].

Bromo-isobutyryl derivative

 $C_{24}H_{30}N_2Br_2O_2$. [175°]. Di-o-tolyl-di-ethylene-diamine C16H22N2. [171°]. Formed, together with di-o-tolylethylene-diamine, by heating o-toluidine with C₂H₄Br₂ (M a. S.). Needles, sl. sol. alcohol. Forms indole when distilled with zinc-dust.

Di-p-tolyl-di-ethylene-diamine (360°). Formed by heating C.H.Cl.OH with p-toluidine at 220° (Wurtz, A. Suppl. 7, 94; Demole, A. 173, 138). Prisms, v. sl. sol. alco-

hol.-B"H,PtCl,

Tri-p-tolyl-tri-ethylene-triamine N₃(C,H_.)₃(C,H_.)₃. [186°]. Formed by heating p-toluidine with C₂H_.Br₂ (G.). Needles, v. sl. sol. alcohol. Is perhaps identical with the preceding body. Its hydrochloride melts at 189°.

Reference .- NITRO - DI - TOLYL - ETHYLENE - DI-

-TOLYL ETHYLENE ETHYL DIOXIDE C.H.Me.O.CH., CH., OEt. (244°). Formed from C.H.O.C.H.Br and alcoholic potash (Schreiber, B. 24, 195).

DI-p-TOLYL ETHYLENE DIKETONE

C₁₈H₁₈O

i.e. C₂H₄(CO.C₆H₄Me)₂. Di-toluyl-ethane. [159°]. Formed by the action of succinyl chloride and AlCl, on toluene (Hollemann, R. T. C. 6, 70; Claus, B. 20, 1877). Needles, v. sl. sol. cold alcohol, insol. alkalis. AcCl converts it into di-p-tolyl-furfurane C₁₈H₁₆O [164°]. Ammonium acetate and HOAc on boiling form di-p-tolyl-pyrrole [197°]. P2S, forms di-p-tolylthiophene [171°].

DI-p-TOLYL-ETHYLENE-DI-METHYL-DI-AMINÉ C₁₆H₂₁N₂ i.e. C₂H₄(NMe.C₄H₄Me)₂. [80°]. Formed by boiling its di-methylo-di-bromide with NH₃Aq (Hübner, A. 224, 337). Tables or prisms (from alcohol), v. sl. sol. water.— B"H₂HgCl₄. [190°].—B"H₂PtCl₈. Orange-

yellow powder. Methylo-iodide B"MeI. Needles. De-

composed at 100°. M. sol. hot water.

Methylo-bromide B"2MeBr. Formed by heating di-methyl-p-toluidine with C₂H₄Br₂ at 105° for some days. Yields B"Me₂Hg₂Cl₃ [159°-162°], B'Me,SnCl, B"Me,PtCl, B"2C, H2MeN2O2, [197°], which crystallises from alcohol

TOLYL ETHYLENE OXIDE v. ETHYLENE

ETHER OF CRESOL.

DI-p-TOLYL ETHYLENE DISULPHONE $(C_1H_1.\hat{S}O_2)_2C_2H_1$. [201°]. Made by boiling sodium toluene p-sulphinate with C₂H₄Br₂ and alcohol (Otto, J. pr. [2] 30, 354; 40, 534). Got also by boiling the same salt with CH₈.CCl₂.CO₂Na. Needles or plates. Dilute KOHAq converts it into toluene sulphinic acid and C, H, SO2, C, H, OH [55°], which yields C,H,.SO₂.C₂H,O [79°], C,H,.SO₂.C₂H,I [100°], C,H, SO₂.C₂H,OBz [176°] and (C₂H₂,SO₂.C₂H₄)₂S [150°-160°]. Ammonia forms (C,H,.SO₂,C₂H₄)₂NH, which yields a hydrochloride [201°].
DI-p-TOLYL-ETHYLENE-UREA

 $CH_2 N(C, H_1) > CO.$ [228°]. Formed from di-

p-tolyl-ethylene-diamine and COCl₂ (Michler a. Keller, B. 14, 2184). Needles.

 $p ext{-} ext{TOLYL-DI-E} ext{THYL-PHOSPHINE}$

C.H., PEt. (240°). Formed from C.H., PCl. and ZnEt. (Czimatis, B. 15, 2016). Liquid.—
B'Mel. [137°].—B'.Me., PtCl. Yellow plates.
p.TOLYL ETHYL SULPHONE

C,H, SO₂,C₂H₅. [56°]. Formed by oxidation of C,H,.SEt and by the action of EtBr on sodium toluene p-sulphinate (Otto, B. 13, 1276; 18, 161). Formed also by warming the acid C,H,.SO,.CHMe.CO,H with KOHAq. Trimetric plates; a:b:c = .526:1: .721. Sol. alcohol and

p-TOLYL ETHYL SULPHONE a-CARB-OXYLIC ACID C,H,.SO,.CHMe.CO,H. Tolyl-sulphono-propionic acid. [37°]. Formed by sulphono-propionic acid. heating C,H,,SO₂Na with CH,,CHBr.CO₂Et and alcohole at 150° and saponifying the product (Otto, J. pr. [2] 40, 555). Crystals (from alcohol), converted by Cl into CH, CHCl.SO, C,H,.

TOLYL ETHYL DISULPHOXIDE v. Ethyl

ether of Toluene Thiosulphonic acid.

TOLYL ETHYL THIOBIURET C, H, N, S, [134°]. Formed from tolyl-thiobiuret, alcohol, aqueous NH, and EtI (Tursini, B. 17, 585).

Needles (from alcohol). o-TOLYL-ETHYL-THIOSEMICARBAZIDE C.H.NH.NH.CS.NHEt. [130°]. Formed o-tolylhydrazine and an alcoholic solution of ethylthiocarbimide (Dixon, C. J. 57, 262). Needles, v. sl. sol. cold water. CuSO, colours its alco-

holic solution deep blue.
TOLYL ETHYL DI-THIO-CARBONATES

CS(OEt).S.C.H.Me. The o-, m-, and p- compounds formed by the action of potassium xanthate on cold solutions of o-, m-, and p-diazotoluene chloride are oils (Leuckart, J. pr. [2] 41, 188). They are converted into tolyl mercaptans by boiling with alcoholic potash. o-TOLYL-ETHYL-THIO-UREA

CS(NHEt).NHC_eH₄Me. [84°]. Formed from o-tolyl-thiocarbimide and ethylamine (Staats, B. 13, 136). Prisms, insol. water, sol. alcohol and

p-Tolyl-ethyl-thio-urea [96°]. Formed in like manner (Weith, B. 8, 1530). Tables, v. sol. hot water.

o-Tolyl-di-ethyl-thio-urea CS(NEt2).NHC,H, [102°]. Formed from o-tolyl-thiocarbimide and NHEt. (Gebhardt, B. 17, 3038). Needles or prisms

p-TOLYL-ETHYL-TOLUTRIAZINE DIHY- $C_{17}H_{19}^{'}N_{s} \quad \textit{i.e.} \quad C_{s}H_{s}Me < \stackrel{N.CHEt}{\stackrel{N.N.C_{s}H_{4}Me}{}}$ DRIDE

[168°]. Formed by heating toluene-azo-toluidine with propionic aldehyde at 140° (Goldschmidt a. Poltzer, B. 24, 1009). Needles (from hot benzene). — B'HCl. [96°]. - Platinochloride [221°]. Small yellow needles.

p-TOLYL-ETHYL-UREA $C_{10}H_{11}N_2O$ NHC,H,.CO.NHEt. Formed from p-toluidine and ethyl cyanate (Sell, A. 126, 162). Crystals, insol. water, v. e. sol. alcohol.

DI-o-TOLYL-FORMAMIDINE C15H16N2 i.e. NHC,H,.CH:NC,H,. [151°]. Formed by boiling the formyl derivative of o-toluidine for a long time, or by heating it with o-toluidine and PCl_s (Ladenburg, B. 10, 1260). Formed also by distilling the thioformyl derivative of o-toluidine in vacuo (Senier, C. J. 47, 762). Prisms (from alcohol), insol. dilute NaOHAq. Yields crystalline $C_{15}C_{16}N_2Br_2$ - $B'_2H_2PtCl_6$

Di-m-tolyl-formamidine NHC,H,.CH:NC,H,. [123°]. Formed by boiling m-toluidine with formic acid (Niementowski, B. 20, 1893). Needles or plates, insol. water. Br in CS₂ yields crys-

talline NHC,H,-CHBr.NBrC,H,-B'HCl. [214°].

B',H,PtCl,-B'C,H,NBrC,H,-B'Cllow needles.

Di-p-tolyl-formamidine [141°]. Formed by distilling C,H,NH.CHS in vacuo (Senier, C. J. 47, 767; B. 18, 2296). Prisms.—B',H,PtCl,

DI-p-TOLYL-FURFURANE C18H18O CH:C(\dot{C} ,H₁) O. [164°]. Formed by the action of Accl on $C_2H_2(CO.C_7H_7)_2$ (Holleman, R. T. C. 6, 72). Small plates, converted by P2S, into ditolyl-thiophene, and by ammonium acetate into di-tolyl-pyrrole.

p-TOLYL-GLYOXAL C.H.Me.CO.CHO. [102°]. Formed from the oxim by dissolving in aqueous NaHSO3, stirring the crystalline mass with alcohol and a little HOAc, filtering, and boiling with dilute H2SO4 (Müller a. Pechmann, B. 22, 2556). Needles (from hot water), v. sol. alcohol. Reduces cold ammoniacal AgNO, but not Fehling's solution. When shaken with benzene (containing thiophene) and H₂SO₄ it colours the benzene green. Yields p-tolyl-gly-oxylic acid and p-toluic acid on oxidation. Phenyl hydraside C,H,C(N,HPh).CH(N,HPh). [145°]. Yellow needles (from dilute alcohol).

Oxim C,H,.CO.CH:NOH. Tolyl nitrosomethyl ketone. [100°]. Formed from tolyl methyl ketone, amyl nitrite and NaOEt. Needles (from benzene).

Acetyl derivative of the syn-oxim C,H,CO.CH:NOAc [68°]. From the oxim and Ac,O (Söderbaum, B. 25, 3461). Tables (from MeOH). Cold NaOHAq splits it up into NaCy and p-toluic acid. Conc. H₂SO₄ acts in like manner. Ac₂O at 100° forms C₇H₇.CO.CN [52°], whence boiling NaOHAq forms p-toluic acid.

Acetyl derivative of the anti-oxim C,H,C(OH)2.CH:NOAc. [148°]. From the oxim and AcCl at 0°, followed by water. Cold NaOHAq forms C,H,CH(OH).CO,H [146°], v. sol. ether. Conc. H,SO, converts this acetyl anti-oxim into the oxim. Converted by KCy dissolved in dilute alcohol into p-toluyl-formoin C,H,.CO.CH(OH).CO.CO.C,H, [161°] (Söder. baum, B. 25, 3473)

p - TOLYL - GLYOXALINE C,oH,oN C,H,N < CH:CH (285°). Got by warming

 $C_rH_rN < CH = CH$ C(SH):Nwith dilute HNO, (Marckwald, B. 25, 2365). Pale-yellow crystals, sl. sol. water. Smells like mushrooms.—B'2H2PtCla. B'AgNO, —Picrate. [179°]. Golden n. p-TOLYL-GLYOXALYL MERCAPTAN Golden needles.

C,H,N < CH = CH [205°]. Got by action of boiling HClAq on the product of the action of amido-acetal on p-tolyl thiocarbimide (Marckwald, B. 25, 2363). Silvery leaflets, m. sol. hot water. Yields $(C_{10}H_{10}N_{2}S)_{2}PtCl_{4}$. MeI in alcohol forms $(C_{10}H_{3}MeN_{2}S)Hi$ [95°], which yields the base $C_{7}H_{7}N_{2}C(SMe):N$ [90°], which forms a

picrate [140°].

p-TOLYL-GLYOXYLIC ACID C.H.O. i.e. C.H.Me.CO.CO.H. [97°]. Formed by the action of AlCl, on a mixture of toluene and Cl.CO.CO₂C₃H₁₁ (Roser, B. 14, 1750). Formed also by oxidising p-tolyl methyl ketone with cold alkaline K.FeCy. (Buchka a. Irish, B. 20, 1762, 2213). Needles (from ligroin), sl. sol. hot water, v. sol. alcohol and ether. Yields p-toluic and terephthalic acids on oxidation. Benzene (containing thiophene) when shaken with tolyl-gly-oxylic acid and H₂SO₄ is turned red, and finally bluish-violet.

Salts.—KA'.—NaA' jaq (Claus a. Kroseberg, B. 20, 2048).—BaA',—BaA', 8aq.—CaA', aq.—AgA'. Needles, v. sol. hot water.

Ethyl ether EtA'. (260°-270°).

Amide C.H.NO. [160°]. Prisms.
Phenyl hydrazide [144°].
DI-o-TOLYL-GUANIDINE C.H., N. i.e.

C(NH)(NHC,H,), [179°]. Formed by the action of NH, lead acetate, and KOHAq on di-otolyl-thio-urea (Berger, B. 12, 1855). Crystals, sol. ether. Cyanogen passed into its alcoholic solution forms the dicyanide C₁₇H₁₇N₂ [174°] which is converted by HCl into the oxalyl derivative C₁,H₁₈N₂O₂ [207°], whence boiling with alcoholic HCl forms CO NC.H., CO. [214°]. The dicyanide is converted by boiling with aniline into C₂₂H₂₁N₂, which yields crystalline B'HCl aq.—B'₂H₄PtCl₆: yellow pp.
Di-p-tolyl-guanidine NH.C(NHC₂H₂)₂. [168°].

Formed by passing cyanogen chloride into fused p-toluidine (W. Wilson, C. J. 3, 154; A. G. Perkin, C. J. 37, 696). Formed also by desulphuration of di-tolyl-thio-urea in presence of NH_s (Hofmann, B. 7, 1739). Needles (from ligroin). Nitric acid (S.G. 1.5) gives a di-nitro- derivative [197°], which forms crystalline B'HNO. Alcohol and HNO, (S.G. 1.4) produce di-nitrop-tolyl-urea. Cyanogen passed into its ethereal solution forms $C_{1b}H_{17}N_3Cy_2$, crystallising from ether in prisms, converted by dilute HClAq into NH.C<N(C,H,).CO[188.5°] (Landgrebe, B. 10,

The cyanide is converted by boiling in 1587). alcoholic solution with aniline hydrochloride into

 $C_{22}H_{21}N_5\frac{e}{2}aq$ [110°-115°].—Salt B'₂H₂PtCl₆. Tri - o - tolyl - guanidine $C_{22}H_{23}N_3$ C(NC,H,)(NHC,H,)2. [131°]. Formed by the action of o-toluidine in alcoholic solution in presence of lead oxide on di-o-tolyl-thiourea (Berger, B. 12, 1857; cf. Girard, B. 6, 445). Formed also when di-o-tolyl-thiourea is boiled for a long time (Barr, B. 19, 1769). Minute prisms. Yields a dicyanide C24H23N5 [141°], con-

verted by cone. HClAq in alcoholic solution into $C_{24}H_{21}N_{1}O_{2}$ [179°].— $B_{2}^{\prime}H_{2}PtCl_{4}$: yellow prisms. Tri-p-tolyl-guanidine $C_{22}H_{23}N_{1}$. [123°]. S. (alcohol) 7.4 at 0°. Formed by heating di-ptolyl-thio-urea with copper (Merz a. Weith, Z. 1868, 610), or, in alcoholic solution, with ptoluidine and PbO (Hofmann, B. 2, 459). Got by heating p-toluidine with PCl₃ and p-tolyl cyanate (Weith, B. 9, 820). Formed also when di-p-tolyl-urea is boiled for a long time (Barr, B. 19, 1768), and likewise by the action of p-toluidine on the product of the action of chlorine on tolyl cyanate (Nef, A. 270, 322). Needles (from ligroïn). Yields a dicyanide Needles (from ligroin). Yields a dicyanide C₂₄H₂₃N₅ [184°], which forms B'HCl 3aq and B'₂H₂PtCl₆ and is converted by boiling with alcoholic HCl into di-tolyl-parabanic acid.

Salts.-B'HClaq. S. 6 at 0°.-B'2H2PtCl. S. 045 at 0°.—B'HNO₃. S. 07 at 0°.—B'₂H₂SO₄. Plates (from hot water). Neutral in reaction.

p-TOLYL HEPTADECYL KETONE [1:4]C_sH₄Me.CO.C₁,H_{2s}. $[67^{\circ}]$. $(278^{\circ}$ at 15 mm.). Formed from toluene, stearyl chloride, and AlCl, (Krafft, B. 21, 2268). Oxidised by HNO, (S.G. 1.12) to p-toluic acid.

p-TOLYL-HEXYL-TOLUTRIAZINE
HYDRIDE C_nH₂N₃ i.e.

C_oH₄M₀ N.C.H.C_oH₁₉ [165°]. Forme Formed by heating cenanthol with toluene-o-azo-toluidine at 175° (Goldschmidt a. Poltzer, B. 24, 1010). Needles, sol. hot benzene.—BHCl. [96 B'2H2PtCl₆. [171°]. Yellow crystalline pp. c-TOLYL-HYDANTOIN C₁₆H₁₀N₂O₂. [1 [96°].-

¯[176°]. Formed by heating o-tolyl-amido-acetic acid with urea at 180° (Ehrlich, B. 16, 742). Lightyellow plates, sol. alcohol and hot water. On boiling with baryta-water it gives o tolylhydantoic acid, which, when set free from its salts, at once splits up into water and o-tolylhydantoïn.

p-Tolyl-hydantoin C10H10N2O2 i.e. CO NH CO . [210°].

Formed, together with p-tolyl-hydantoic acid CO(NH₂).N(C,H₂).CH₂.CO₂H, by fusing p-tolylamido-acetic acid with urea (Schwebel, B. 11, 1128). Needles (from water), v. sol. alcohol. p-Tolyl-hydantoic acid is crystalline, sl. sol. hot alcohol and hot water.

 $CO < N(C, H_1).CO$ $N(C, H_2).CH_2$ Di-o-tolyl-hydantoïn Got from o-tolyl-amido-acetic toluide and COCl₂ (Bischoff, B. 25, 2275).

 $CO < N(C,H_1).CO \\ N(C,H_1).CH_2$ Di-p-tolyl-hydantoïn [175°]. Formed from p-tolyl-amido-acetic toluide and COCl₂ (Bischoff, B. 25, 2280). Plates (from alcohol), sl. sol. ligroin.

TOLYL-HYDRAZIDO-METHYL-THIAZOLE $\begin{array}{ll} \textbf{DIHYDRIDE} \stackrel{CHMe.S}{\leftarrow} C.NH.NHC_{7}H_{7}. \ \ \textbf{Formed} \end{array}$ by heating tolyl-allyl-thio-semicarbazide with conc. HClAq at 100° (Avenarius, B. 24, 270). The p- compound melts at 133°. Both the o- and the p- compounds form crystalline hydrochlorides.

o-TOLYL-HYDRAZIDO-PROPIONIC ACID C₆H₄Me.NH.NH.CHMe.CO₂H. [143°]. Formed by the action of sodium-amalgam on the o-tolylhydrazide of pyruvic acid (Japp a. Klingemann,

C. J. 53, 519). Small needles (from MeOH). o-TOLYL-HYDRAZINE C,H,NH.NH, [53°] (P.); [56°] (F.); [59°] (Gallinek a. Richter, B. 18, 3175). Prepared from o-toluidine in the same manner as phenyl-hydrazine is obtained from aniline (Fischer a. Bösler, A. 212, 338; Preund, B. 24, 4200). Glittering plates, slowly oxidised by air, forming a brown oil.—B'HCl aq: needles.—B'HNO₃: plates. With SOCl₂ and ether it forms C.H.NH.N:SO, a yellow oil of the state smelling like geraniums (Michaelis, A. 270, 119). Glucose yields o-tolyl-glucosazone C20H26N,O. [201°] (Raschen, A. 239, 229). Di-methyl di-ketone forms CH₃.C(N₂HC,H₇).C(N₂HC,H₇).CH₃ [198°] (Japp a. Klingemann, A. 247, 224

Formyl derivative C.H.Me.NH.NHCHO. [121°]. Formed from o-tolyl hydrazine and formamide. Needles (Gattermann, B. 25, 1078). Acetyl derivative C, H, NH. NHAc. [104°].

Formed from o-tolyl-hydrazine and Ac₂O (G.).

Propionyl derivative C,H,NH.NH.C,H,O. [84°]. Formed from otolyl-hydrazine and propionic acid (G.). Colourless tables.

C,H,.NH.NHBz. Benzoyl derivative [180°]. Got from the hydrazine and BzCl in Et,O (G.). Needles. Gives off all its nitrogen

in the free state on boiling with Fehling's solution (Strache a. Iritzer, M. 14, 38).

m - Tolyl-hydrazine C.H., NH.NH. (242°).
(Buchka a. Schachtebek, B. 22, 841; cf. V. Meyer a. Lecco, B. 16, 2976). Oil.—B'HCl. Needles, v. sol. water and alcohol.

p - Tolyl - hydrazine C,H,NH,NH₂. [61°]. (242°). Obtained from p-toluidine (Fischer, B. 8, 589; 9, 890). White plates (from ether), sl. sol. water.

Reactions.—1. With SOCL and ether it. forms C.H.NH.N:SO [112°], crystallising in yellow needles (Michaelis s. Ruhl, A. 270, 118).— CH.NH.NH.) PO [189°]. — 4. Acetone forms CM-,NH.NH.NH. forms CM-,NH.NH.NH. forms CM-,NH.NH.NH.] PO [189°]. — 5. Acetoacetic etherforms C₁₃H₁₈N₂O₂ [93°] (Knorr, B. 17, 550).— 6. Oxalic ether forms NHC₇H₂NH.CO.CO₂Et [133°] (Preund, B. 24, 4198).—7. Di - methyl diketone yields CH3. C(N2HC7H7). C(N2HC7H7). CH3 [230°], and CH3.C(N2HC7H7)Ac [161°] (Japp, C.J. 53, 544).

Salt.—B'C, H, PH₂O₂. [148°] (Michaelis, A. 270, 134).

Toluene-p-phosphinate B'C,H,PH,O2.

Formyl derivative

C_oH₄Me.NH.NH.CHO. [164°]. Formed by heating the alcoholic solution of p-tolyl-hydrazine with chloroform and potash. Formed also by heating p-tolyl-hydrazine with formamide at 130° (Ruhemann, C. J. 55, 248). Plates (from water), v. sol. alcohol.

C.H.Me.NH.NHAc. Acetyl derivative [121°]. Formed from p-tolyl-hydrazine and AcOH (Gattermann, B. 25, 1080). Plates.

Propionyl derivative [170°]. Needles. Dibenzoul derivative [188°] (Fischer).

Di-o-tolyl-hydrazine.

Formyl derivative (C,H,),N.NH.CHO. 9°]. Formed from C,H,NH.NH.CHO and [139°]. CuSO, (Gattermann, B. 25, 1078). Plates.

Acetyl derivative (C,H,)2N.NHAc.

Acetyl derivative (C₇H₇)₂N.NHAc. [191°]. Formed in like manner from acetyl-otolyl-hydrazine and CuSO4. Needles.

Propionyl derivative [167°]. Needles. Benzoyl derivative [209°]. Needles.

u-Di-p-tolyl-hydrazine (C,H₁)₂N.NH₂. [172°]. Formed by reduction of di-tolyl-nitrosamine with zinc-dust and HOAc (Lehne, B. 13, 1546). Plates, v. sol. alcohol and benzene.—B'HCl

Formyl derivative (C,H,)2N.NH.CHO. [146°]. Formed from formyl-p-tolyl-hydrazine and CuSO, (Gattermann, B. 25, 1079). Plates.

Acetyl derivative. [176°]. Needles. Propionyl derivative [171·5°]. Needles. Benzoyl derivative. [187°]. Needles. s-Di-o-tolyl-hydrazine C,H,NH.NHC,H,

o-Hydrazo-toluene. [165°] (P.); [146°] (S.). Formed by reduction of o-toluene-azo-toluene by sodium-amalgam (Petrieff, B. 6, 557; Schultz, Converted into di-amido-ditolyl B. 17, 467). [3:4:1]C,H₃Me(NH₂),C₆H₃Me(NH₂)[1:3:4] [123°] by heating with HCl. The diamido-derivative C,H₆(NH₂).NH.NH.C₂H₆(NH₂) [180°] is a product of the action of sodium-amalgam on nitro-ptoluidine (Buckney, B. 11, 1453).

C,H,NH.NHC,H,. s-Di-m-tolyl-hydrazine Formed by reducing m-toluene-m-azo-toluene by alcoholic ammonium sulphide (Goldschmidt, B. 11, 1626; Barsilowsky, A. 207, 116). Liquid, converted by H.SO, into a di-amido-ditolyl.

s-Di-p-tolyl-hydrazine C,H.NH.NHC,H. [126°]. Formed in like manner from p-toluenep-azo-toluene (Melms, B. 3, 553) and by the action of zinc-dust and NaOHAq on p-nitrotoluene (Janovsky, M. 9, 829). Tables or needles, v. e. sol. alcohol. In alcoholic solution it is readily oxidised by air to C,H,.N,2.C,H,. Dilute H2SO4 converts it into toluidine and tolueneazo-toluene. On treatment in alcoholic solution with hydrochloric acid it changes to tolylene-The di-amido-derivative tolyl-diamine. $[1:2:4]C_6H_3Me(NH_2).NH.NH.C_6H_3Me(NH_2)[4:1:2]$ obtained from C.H.Me(NH2).N2.C.H2Me(NH2) by reducing with sodium-amalgam, is crystalline and yields B"H₂SO₄, B"2HCl, B"2HBr, and B"H₂PtCl₆ (Graeff, A. 229, 352).

Reference. - OXY-TOLYL-HYDRAZINE.

o-TOLYL-HYDRAZINE SULPHONIC ACID C₆H₃Me(N₂H₃).SO₃H [1:2:4]. Formed by adding o-diazo-toluene p-sulphonic acid to a cold solution of SnCl₂ (Limpricht, B. 18, 2193). Colourless needles, v. sol. hot water, nearly insol. alcohol.—KA' 2aq: orange-red prisms.—BaA': yellow crystalline powder.

o-Tolyl-hydrazine sulphonic acid

 $C_{s}H_{s}Me(N_{2}H_{3}).SO_{s}H$ [1:x:2]. Formed by heating o-tolyl-hydrazine (1 pt.) with conc. H₂SO₄ (5 pts.) at 100° (Gallinek a. Richter, B. 18, 3175). Thin needles (containing gaq). By boiling with aqueous CuSO, it is quantitatively decomposed into a toluene sulphonic acid and nitrogen.-NaA'3½aq.—BaA'24aq.— ZnA'23aq.—PbA'26aq. —PbA'2(OH)4. Sparingly soluble needles. p-Tolyl-hydrazine-o-sulphonic acid

 $C_6H_3Me(N_2H_3).SO_3H$ [1:4:2]. [274°]. by reducing p-diazo-toluene o-sulphonic acid with Na₂SO₃ (Brackett a. Hayes, Am. 9, 401) or HCl and SnCl₂ (Pasche, B. 21, 3416). Tables, sl. sol. cold water. Decomposed by fusion. Not attacked by warm H2SO4.

-Tolyl-hydrazine m-sulphonic acid

 $C_{s}H_{3}Me(N_{2}H_{3}).SO_{s}H$ [1:3:4]. Obtained by reducing p-diazotoluene m-sulphonic acid by cooled SnCl₂ (Limpricht, B. 18, 2193). Slender prisms, sol. hot water. FeCl, evolves nitrogen in the cold. Conc. H₂SO₄ at 80° forms a red amorphous substance (Schneider, Am. 8, 271).

p-Tolyl-hydrazine disulphonic acid C₆H₂Me(N₂H₃)(SO₃H)₂. Formed by adding p-diazo-toluene disulphonic acid to a cold solution of SnCl₂ (Limpricht, B. 18, 2193). Nodules,

v. sol. water.—BaH₂A'', 2½aq: tables.
Di-tolyl-hydrazine disulphonic acid $C_bH_3Me(SO_2H).NH.NH.C_bH_3Me(SO_2H).$ A product of the reduction of the corresponding azocompound by SnCl₂ (Neale, A. 203, 72). Crystalline powder (containing 2 aq), sl. sol. water .-BaA" 5aq.—CaA" 3 aq: monoclinic efflorescent prisms.

Reference .- NITRO - TOLYL - HYDRAZINE SUL-PHONIC ACID

o-TOLYL-IMIDO-DIACETIC ACID

 $[1:2]C_6H_4Me.N(CH_2.CO_2H)_2$. [c. 160°]. Formed by heating o-tolyl-amido-acetic acid (1 mol.) with chloro-acetic acid (1 mol.), Na₂CO₃ (1½ mols.) and a little water at 140° (Bischoff, B. 25, 1994; 25, 2270). White crystals, insol. ligroin, sl. sol. ether, m. sol. alcohol.—NH,A'. V. sol. Aq. [160°].

Amide C,H,N(CH₂,CO.NH₂)₂. [164]. Plates. Di-o-toluide C,H,N(CH₂,CO.NHC,H,)₂. [150°].

Imide C₇H₇N < CH₂.CO NH. [146°].

Prisms (from alcohol), sl. sol. water.

o-Tolylamic acid C,H,N(CH2.CO2H).CH2.CO.NHC,H, [148°].

Crystals (from alcohol).

p-Tolylimido-diacetic acid [1:4]C₆H₄Me.N(CH₂.CO₂H)₂. Diglycotolylamic acid. [c. 140°]. Formed by the action of CH_Cl.CO.H on p-tolyl-glycocoll (Meyer, B. 14, 1323; Bischoff, B. 23, 2000). Needles (from water). Very unstable. — CuA', aq: green needles.—Ag₂A'(NO₂): needles.—p-Toluidine

salt (C,H,NH2)HA'. [119°]. Crystals, sol. | hot water.

Mono-amide. [222°]. A product of the action of Ac₂O on the compound [168°] got from chloro-acetic acid and p-toluidine (Bischoff, B. 25, 2280).

Di-amide C,H,N(CH₂.CO.NH₂)₂. [250°]. Di-p-toluide C,H,N(CH₂.CO.NHC,H,)₂. [251°]. Long needles (from alcohol).

Amide p-toluide C,H,N(CH,CO.NH,)CH,CO.NHC,H, [210°].

p-Tolylamic acid C,H,N(CH,CO,H).CH,CONHC,H, [222°]. Needles, insol. water, sol. hot alcohol.

p-Tolyl-Imido-Diacrylic ether C_eH₄MeN(CH:CH.CO₂Et)₂. [73°]. Formed from p-toluidine and formyl-acetic ether (Von Pechmann, B. 25, 1053). Yellow needles (from dilute alcohol).

o-TOLYL-8-IMIDO-BENZYL - MALONIC ETHER C21H23NO, i.e. C_8H_3 .C(NC, H_7).CH(CO₂Et)₂. [95°]. Formed by the action of w-chloro-benzylidene-o-toluidine upon sodium-malonic ether (Just, B. 19, 985).

Crystals, v. sol. ether. Split up by dilute HClAq at 120° into acetophenone and o-toluidine. The $m{p}$ -isomeride is liquid.

o-TOLYL-8-IMIDO-BUTYRIC ACID

C₆H₄Me.N:CMe.CH₂.CO₂H. [112°]. Formed by heating acetoacetic ether with o-toluidine at 150° (Knorr, B. 17, 542; Pawlewski, B. 22, 2203). Needles (from hot water), converted by conc. H₂SO₄ into (Py. 1)-oxy-(B. 4, Py. 3)-dimethyl-quinoline.

p-Tolyl-\$-imido-butyric acid. Formed in like manner from p-toluidine. Crystals. verted by H₂SO, into (Py. 1)-oxy-(B. 2, Py. 3)-

di-methyl-quinoline.

 $p ext{-} ext{TOLYL}$ IODO-ETHYL SULPHONE

C,H,SO,CH,CH,I. [100°]. Formed from tolyl oxy-ethyl sulphone and HIAq at 160° (Otto, J. pr. [2] 30, 357). Crystals, v. sol. hot

TOLYL IODO-METHYL SULPHONE

C,H,SO,CH,I. [126°]. Formed by heating sodium toluene sulphinate with CH,I, and alcohol (Otto, B. 21, 655). Small needles, v. e. sol. hot alcohol.

DI-p-TOLYL KETONE CO(C,H,Me)2. [92°]. (383°) at 725 mm. Formed by oxidation of ditolyl-methane (Weiler, B. 7, 1183), of u-di-tolylethane (O. Fischer, B. 7, 1195), and of u-di-tolyl-ethylene (Hepp, B. 7, 1414) by chromic acid mixture.

Preparation.—50 g. AlCl₃, 65 c.c. toluene, 50 c.c. CS2, and 11 c.c. of a solution of COCl2 in CS, saturated at 0° are put into a corked halflitre flask and gently warmed. After 20 minutes HCl is allowed to escape and a second 11 c.c. of CS₂ saturated with phospene is added. operation is repeated 2 or 3 times; the yield is 50 p.c. of the calculated (Elbs, J. pr. [2] 35, 467; cf. Ador a. Crafts, B. 10, 2174).

Properties.—Crystals (from alcohol). Converted by HNO, at 250° into benzoyl-terephthalic acid. Reduced by HI and P to di-tolyl-methane (Ador a. Rilliet, B. 12, 2303).

Oxim (C,H,)2C:NOH. [163°]. Prisms (Goldschmidt, B. 23, 2746). Decomposes on keeping with sudden evolution of nitrous fumes. Conc. I H₂SO, at 100° forms the p-toluide of p-toluic

Di-p-tolyl diketone C,H,.CO.CO.C,H,. [105°]. p- $Toli\bar{l}$. Formed by boiling toluoin (1 pt.) with HNO, (2 pts.) (Stierlin, B. 22, 381). Yellow plates (from warm alcohol). Gives a violet colour when boiled with alcoholic potash.

C,H,.C(NOH).C(NOH).C,H, (a)-0xim[217°]. Plates or needles, sl. sol. alcohol. Yields

a diacetyl derivative [134°].

(β)-Oxim [225°]. Formed, together with the (a)-oxim, by the action of hydroxylamine hydrochloride on the ketone. Needles, v. sol. alcohol. Yields a di-acetyl derivative [144°].

Di-p-tolyl tetraketone

 C_7H_7 . $CO.C(OH)_2$. $CO.CO.C_7H_7$. [88°]. Formed by the action of nitric acid (S.G. 1.3) on C₇H₇.CO.CH(OH).CO.CO.C₇H₇ (Söderbaum, B. 25, 3474). Prisms (containing $(C_{18}H_{16}O_5)_2CS_2$) (when crystallised from CS,).

Di-oxim C_7H_7 .C(NOH).CO.CO.C(NOH). C_7H_7 . [1819]. Plates (containing EtOH) (from alcohol).

Di-oximC.H., CO.C(NOH).C(NOH).CO.C.H., So called '(\(\beta\))-nitro-cymene.' [125°]. Mol. w. 324 (by Raoult's method). Formed by the action of nitric acid on cymene and on p-tolyl methyl ketone (Holleman, R. T. C. 6, 60; cf. Landolph, B. 6, 937; Fittica, A. 172, 314). Needles, v. sol. warm alcohol. Converted by NaOHAq into ptoluic acid. Zinc-dust and HOAc reduce it to C_1H_2 CO. CH_2 CH₂.CO. C_1H_1 [159°]. Alcoholic NH, forms p-toluic amide and C,0H,N,O, [162°]. Boiling Ac₂O forms C₁₈H₁₆N₂O₄Ac₂O [167°].

Reference.—OXY-DI-TOLYL KETONE.

TOLYL-MALONAMIC ACID v. MALONIC ACID. TOLYL-MELAMINE v. Cyanuramide in the article on Cyanic acids.

[1:2]C_eH₄Me.SH. o-TOLYL-MERCAPTAN [15°]. (188°) (H.); (193°) (V.). Formed by reducing (4,1,2)-bromo-tolyl mercaptan (Hübner, A. 169, 30; Vallin, B. 19, 2953). Prepared by boiling o-tolyl ethyl dithiocarbonate with alcoholic potash, the yield being 70 p.c. of the theoretical amount (Leuckart, J. pr. [2] 41, 188). Plates. - Pb(SC,H,)2. Brick-red pp., turning white in air.

m-Tolyl mercaptan [1:3]C₆H₄Me.SH. [c. 200°]. Formed by reducing toluene m-sulphonic chloride with tin and HCl (Hübner, A. 169, 51) and by boiling m-tolyl ethyl xanthate with alcoholic potash (L.). Colourless liquid with intense odour of mercaptan. Volatile with steam. In ammoniacal alcoholic solution it is oxidised by air to m-tolyl disulphide.

p-Tolyl mercaptan [1:4]C,H,Me.SH. [43°]. (188°) (J.); (191°) (Crafts, B. 19, 8180; Otto, B. 19, 8129); (194°) (Vallin, B. 19, 2953). Formed by reducing toluene p-sulphonic chloride with tin and HCl (Märcker, A. 136, 79; Javorsky, Z. 1865, 222). Formed also by heating p-tolyl ethylxanthate with alcoholic potash (L.). Formed also from di-p-tolyl disulphide, alcohol, and H.S (Otto a. Rössing, B. 19, 3130). Unctuous lamina from ether), volatile with steam. Conc. H.SO. forms a blue solution. Readily oxidised to dip-tolyl disulphide [46°]. Chloro-acetone forms C.H.S.CH.2CO.CH. (151° at 15 mm.); S.G. 11: 1.0986 which yields a phenyl-hydrazide [62°] (Delisle, A. 260, 268).—Hg(SC,H₇)₂.— C,H,S.HgCl. Plates.

Ethyl ether C,H,SEt. (221°). S.G. 11 1-0016. Colourless oil (Otto, B. 13, 1277).

References.-AMIDO- and BROMO-

MERCAPTAN.

o-TOLYL-MESITYL-THIO-UREA

C,H,NH.CS.NHC,H2Me3. [167°]. Formed from o-toluidine and mesityl thiocarbimide (Eisenberg, B. 15, 1014). Needles, insol. Aq.

DI-TOLYL-METHANE CH2(C,H2)2.

(286°).

Formation.-1. From formic aldehyde (or methylal), toluene, and H₂SO₄ (Weiler, B. 7, 1181).—2. By reducing di-tolyl ketone with HI and P (Ador a. Rilliet, B. 12, 2302).-3. A product of the action of AlCl, and toluene on CH₂Cl₂, on CH₃Cl, or on CCl₃.NO₂ (Friedel a. Crafts, A. Ch. [6] 11, 266; Bl. [2] 43, 50; Elbs a. Wittich, B. 18, 347).

Properties.—Prisms, v. sol. ether. Oxidised by chromic acid mixture to di-tolyl ketone, toluyl-benzoic acid, and CO(C₆H₄.CO₂H)₂. Yields

a di-nitro- derivative [164°]

Tri-tolyl-methane $CH(C_7H_7)_3$. [73°]. (377°). Obtained from tri-amido-tri-tolyl-carbinol (Rosenstiehl a. Gerber, A. Ch. [6] 2, 353). Crystalline.

References .- DI-BROMO- and DI-CHLORO-, DI-TOLYL-METHANE.

o-TOLYL-METHYL-p-AMIDO-PHENOL

Methyl derivative C,H,NMe.C,H,OMe. (336° cor.). From C,H,NH.C,H,OH, KOH, and MeI (Philip, J. pr. [2] 34, 59). Oil.

TOLYL-METHYL-AMINE METHYLv. TOLUIDINE, METHYL-BENZYL-AMINE, and AMIDO-

Di-p-tolyl-methyl-amine (C,H,)2NMe. (235°-240° at 20 mm.). Formed from ditolylamine, MeOH, and HCl at 250°-280° (Girard, Bl. [2] 24, 120). Does not combine with acids.

DI-TOLYL-DI-METHYL-DI-TRIAZYL

 $\underset{N.N(C_7H_7)}{N.N(C_7H_7)} \geqslant C.C \leqslant_N^{N(C_7H_7).N} \underset{CMe}{\overset{}{\sim}}$ (259°-260°). Formed by boiling p-tolyl-hydrazine cyanide (C,H,NH.NH₂)₂Cy₂ with Ac₂O (Bladin, B. 22, 3114). Crystals (from alcohol), m. sol. alcohol. p-TOLYL-METHYL-BENZYL KETONE

C.H.Me.COCH, C.H.Me. [102°]. Got by reducing p-toluoin with zinc and alcoholic HCl (Stierlin, B. 22, 383). Needles, insol. water, v. sol, benzene.

o-TOLYL-METHYLENE-AMINE

C,H,N:CH₂ (?). Formed by adding o-toluidine to a solution of formic aldehyde (Wellington a. Tollens, B. 18, 3307). Colourless syrup.

p-Tolyl-methylene-amine C,H,N:CH, (?). [c. 122°]. Formed, together with a polymeride sl. sol, benzene and alcohol, by adding p-toluidine to a solution of formic aldehyde (W. a. T.). Crystals, v. sol. benzene.

Di-o-tolyl-methylene-diamine (C,H,1(H)2CH2. (over 850°). Formed, together with the following solid isomeride, by heating o-to uidine with CH₂Cl₂ at 110°-115° (Grünhagen, A. 256, 305). Liquid, v. sol. ether and alcohol.—B' H2PtCl.: brown amorphous powder.

Isomeride [c. 135°]. Crystalline powder, sl. sol. cold alcohol. — Salts: B"2HCl. — B"2HBr.—B"H₂SO₃.—B"2H₂PO₄.—B"3H₄C₄O₄. -B"2H,CO. Radiating aggregates.

Di-p-tolyl-methylene-diamine

(C,H,NH),2CH, (above 350°). Formed, together with the solid isomeride, by heating p-toluidine with CH₂Cl₂ (G.). Oil, v. sol. ether.—B"HOl.—

B"<u>H_PtCl</u>₂.—B"HAuCl₄.

Isomeride C₁₆H₁₈N₂. [o. 156°]. (over 350°).

Amorphous, sl. sol. ether. — B"2HCl (?). —
B"H_PtCl₆.—B"2HAuCl₄.—B"2H₂C₂O₄. Orystal-

line crusts.

Di-p-tolyl-di-methylene-diamine

C,H,N CH₂NC,H, [90°]. A product of the action of CH₂Cl₂ on p-toluidine at 100° (Grünhagen, A. 256, 296). Granules, sol. hot alcohol. Converted by nitrous acid into a crystalline nitroso- derivative (C,H,N)₂C₂H₂(NOH) (?) -B"2HCl.—B"2HAuCl₂—B"2HBr.—B"H₂SO₄.

o-Tolyl-trimethylene-diamine $C_sH_e(NH_2).NHC_rH_r$ (281°). Formed by reducing o-tolyl-pyrazole dihydride in alcoholic solution by sodium (Balbiano, G. 18, 354). Oil.

p-Tolyl-trimethylene-diamine C₂H_a(NH₂).NHC₇H₇. [c. -15°]. (287°). A product of the action of Na on an alcoholic solution of p-tolyl-pyrazole (Balbiano). Liquid. Oxalate. [208°]. Spherical groups of minute needles.

p-TOLYL METHYLENE METHYL SUL-PHÔNE KETONE C,H,.SO,.CH,.CO.CH,. [51°]. Formed from chloro-acetone and sodium toluene p-sulphinate (Otto, J. pr. [2] 36, 401). Silky needles. Yields a bromo- derivative [130°].

m-TOLYL-METHYLENE-PHTHALIDE

 $C_8H_4 < C(CHC_7H_7) > 0$. [153°]. Formed by heating m-tolyl-acetic acid with phthalic anhydride and a little NaOAc (Heilmann, B. 23, 3157). Needles (from ligroin), sl. sol. ether.

p-Tolyl-methylene-phthalide C16H12O2 p-Xylal-phthalide. [151°]. Formed in like manner from p-tolyl-acetic acid (Ruhemann, B. 24, 3965). Yellow needles (from alcohol), sl. sol. ether. Nitrous acid passed into its solution in chloroform forms $C_0H_4 < \frac{C(C(NO_2),C_1H_1)}{CO} > 0$ [140°], which may be reduced to the compound C₄H₄<0.0,H₇. [116°].

Reference.—NITRO-TOLYL-METHYLENE-PHTHAL-TDE.

m-TOLYL-METHYLENE-PHTHALIMIDINE $C_0H_4 < CO_-$ NH. [165°]. Formed by heating m-tolyl-methylene-phthalide with alcoholic ammonia (Heilmann, B. 23, 3161). Yellowish needles.

p-Tolyl-methylene-phthalimidine C, H, NO. [204°]. Formed by heating p-tolyl-methylene-phthalide with alcoholic NH₂, the compound NH₂.CO.C₄H₄.CO.CH₂C₃H₄, being first formed (Ruhemann, B. 24, 3968). Converted by passing nitrous acid through its solution in chloroform into C₆H₄ CO NO₂).C₇H₇) NH [227°].

Isomeride $C_8H_4 < \stackrel{OH;C.C_7H_7}{CO.NH}$. [228°]. Formed by heating $C_6H_4 < \stackrel{CH:C.C_7H_7}{<}$ alcoholic ammonia (R.).

Reference.-NITBO-TOLYL-METHYLENE-PHTHAL-IMIDINE.

DI-p-TOLYL TRIMETHYLENE DISUL-PHONE C₄H_o(SO₂C₇H₇)₂. [125°]. Formed from trimethylene bromide and sodium toluene p-sulphinate (Otto, B. 24, 1834). Plates. Converted by alcoholic potash at 120° into the compound $O(CH_2.CH_2.CH_2.SO.C_7H_7)_2$ [80°]

TOLYL-METHYL-IMESATIN v. Toluide of

Methyl-isatin.

m-TOLYL METHYL KETONE C,H, CO.CH, Methyl-acetophenone. (219°). Formed by distilling calcium m-toluate with calcium formate (Buchka a. Irish, B. 20, 1766).

Tolyl methyl ketone. (225°). S.G. 20 .9891. V.D. 4.58. Formed, together with the p-compound, by the action of AcCl on toluene and AlCl₃ (Essner a. Gossin, Bl. [2] 42,95). This is

perhaps the o-compound.

p-Tolyl methyl ketone. (217°) (M.); (230° cor.) (W.). Formed by the action of AlCl₃ on a mixture of toluene and Ac2O or AcCl (Michaelis, B. 15, 185; Claus, B. 19, 234). Formed also by heating toluene with HOAc, ZnCl₂, and POCl₃ (Frey, J. pr. [2] 43, 114) and as a by-product in the oxidation of (a)-nitro-cymene (Widman, B. 19, 587). Oil. Yields a dibromide C_uH_uBr₂O [100°]. KMnO₄ forms terephthalic acid. Alkaline K₂FeCy₅ forms p-toluic and p-tolyl-glyoxylic acids (B. a. K.). HNO₃ forms C₁₈H₁₄N₂O₄ [125°], which is reduced by zinc-dust and HOAc to C₂H₄(CO.C₇H₇)₂ (Hollemann, R. T. C. 10, 211). Oxim [88°]. Crystals (from ligroïn).

Phenyl-hydrazide [97°]. Prisms.

Reference. - AMIDO-, IODO-, and OXY-, TOLYL METHYL KETONE.

o-TOLYL METHYL KETONE TETRA-HYDRIDE $CH_2 < \frac{CH_2 \cdot CMe}{CH_2 \cdot CH_2} > C.CO.CH_8$. (206°).

Formed from αω-di-acetyl-pentane by treatment with conc. H₂SO₄ (Kipping a. Perkin, jun., C. J. 57, 18). Oil, smelling like peppermint. Does not combine with NaHSO₃. In moist ether it is reduced by Na to the hexahydride.

Oxim C.H., NO. Liquid. Phenyl hydrazide C, H20N2. Oil. Tolyl methyl ketone hexahydride v. METHYL-HEXAMETHYLENYL METHYL KETONE.

m-TOLYL-METHYL-MALONIC ACID $[1:3]C_6H_4Me.CH_2.CH(CO_2H)_2.$ [133°]. Formed by saponification of the ether (Poppe, B. 23, 110). Rhombohedra, sl. sol. water. K2A": needles.

Methyl ether Me₂A". (c. 300°). Ethyl ether Et₂A". (320°). Formed from e-bromo-m-xylene and sodium malonic ether. Oil. Converted by alcoholic NH, at 150° into C₈H₄Me.CH₂.CH(CO₂Et).CO.NH₂[186°], which on warming with water forms the ammonium salt C₆H₄Me.CH₂.CH(CO₂Et).CO₂NH₄[77°]. Et₂A" is converted by methylamine into the methylamide C.H.Me.CH2.CH(CO2Et).CONHMe [120°], and by aniline into C.H.Me.CH.CH(CO.NHPh), [188°].

Di-m-tolyl-di-methyl-malonic ether $(C_0H_1Me.CH_2)_2C(CO_2Et)_2$. [122°]. A product of the action of ω -bromo-m-xylene on sodium

malonic ether (Poppe, B. 23, 109). o-TOLYL-METHYL-OXAZOLINE O11H13NO i.s. CHMe.O C.O.H. (258°). Formed by boiling C.H., CO.NH.CH. CHMeBr with alcoholic potash (Salomon, B. 26, 1828). Oil.—B'C.H.N.O., [129°].—B'.H.PtCl., Vol. IV.

p-Tolyl-methyl-oxazoline C, H, NO. (265°). Formed from the bromopropyl-amide of p-toluic acid (Salomon, B. 26, 1326). Oil.—Picrate B'C,H,N,O,. [183°].—B',H,PtCl,. [184°]. p-TOLYL-DI-METHYL-PHOSPHINE

C₇H₇PMe₂. (210°). Formed from C₇H₇POl₂ and ZnMe₂ (Czimatis, B. 15, 2014). Oil, oxidised by HgO to C,H,PMe2O. CS2 forms C,H,PMe2CS2 [110°] crystallising in red plates.

Methylo-iodide B'MeI. [255°]. Needles.

Yields B'MeI, and B'2Me2PtCl

Benzylo-chloride. Yields $B'_{2}(C_{7}H_{7})_{2}PtCl_{6}$ [226°], a yellow crystalline pp.

p-TOLYL-METHYLPHTHALIMIDE $[4:2:1]C_0H_3Me:C_2O_2:N.C_0H_4Me.$ [180]. Formed by distilling (4,2,1)-methyl-phthalic acid with p-toluidine (Niementowski, M. 12, 630). Crystalline mass, v. e. sol. chloroform, insol. water.

p-TOLYL-METHYL-PHTHALIMIDINE

 $C_8H_4 < CO \xrightarrow{CH(CH_2.C_8H_4Me)} NH.$ [149°]. Formed from p-tolyl-methylene-phthalimidine, HI and P (Ruhemann, B. 24, 3969). Needles.

DI-p-TOLYL-METHYL-PIPERAZINE

 $C_{19}H_{21}N_2$. [105°]. Formed from di-p-tolylpropylene-diamine, ethylene bromide. Na₂CO₃. Needles (from alcohol) (Bischoff, B. 25, 3278).

o-TOLYL- METHYL-PROPYLENE- 4-THIO-UREA $\stackrel{\rm CHMe.S}{_{\rm CH_2}} >$ C.NMe.C,H,. (c. 295°). Formed from o-tolyl-propylene-\psi-thio-urea and MeI (Prager, B. 22, 2999). Yields (\$)-methyl-taurine on oxidation.—B'C₆H₂N₃O₇. [138°].—B'H₂[166°].—B'₂H₂PtCl₆. Decomposes over 200°. [138°]. — B'HI.

p-TOLYL-DI-METHYL-PYRROLE

 $(C_7H_7)N < \stackrel{CMe:CH}{Cme:CH}$. [46°]. (255°). Got by heating the di-carboxylic acid (Knorr, B. 18, 308). Crystals, volatile with steam.

Reference.—AMIDO-TOLYL-DI-METHYL-PYRROLE. p-TOLYL-DI-METHYL-PYRROLE DICARB-

OXYLIC ACID (C,H,)N CMe:C.CO,H. Got by saponifying its ether, which is formed by mixing acetic acid solutions of di-acetyl-succinic ether and p-toluidine (Knorr, B. 18, 304). Needles, decomposes at about 250°, with evolution of CO. -K₂A".--AgHA".

Ethyl ether Et2A". [67°]. Tables.

Reference.—AMIDO-TOLYL-DI-METHYL-PYRROLE DICARBOXYLIC ACID.

 $p ext{-} ext{TOLYL-DI-METHYL-SULPHAMIDE}$

[91°]. Formed SO₂(NMe₂).NHC₇H₇. NMe2 SO2Cl and p-toluidine (Behrend, A. 222, Crystals, v. sol. alcohol and ether.

p-TOLYL METHYL SULPHONE C,H,SO,CH, [87°]. Formed by the action of conc. KOHAq on p-tolyl-sulphono-acetic acid Otto, B. 18, 161). Got also from C,H,SO,Na Needles (from very dilute and MeI (Otto). alcohol), v. sol. hot water.

O-TOLYL-METHYL-THIAZOLINE

 $CHMe.S > C.C_7H_7$. (284°-295°). Formed by heating o-toluic bromo-propylamide with P.S. at 150° (Salomon, B. 26, 1328). Oil. — B'C,H,N,O,. [186°

p-Tolyl-methyl-thiazoline C11H12NS. (295°). Oil.—B'C.H.N.O. [141°].—B'.H.PtCl.

o-TOLYL-METHYL-THIOHYDANTOIN

 $_{\mathrm{HMe}}<_{\mathrm{NH.\,CS}}^{\mathrm{CO}\ N(\mathrm{C,H,})}$ [198°]. Formed from o-tolyl-thiocarbimide and alanine (Marckwald,

B. 24, 3281). Crystals.

p - Tolyl-methyl - thiohydantoïn. Formed in like manner from p-tolyl-thiocarbimide and alanine (Aschan, B. 17, 427). Minute prisms. On heating with alcoholic potash it is converted by hydrolysis into a salt of the oily NHC,H,.CS.NH.CHMe.CO,H, which slowly changes, in the free state, back into the hydantoïn.

o-Tolyl-di-methyl-thiohydantoïn

 $\text{CMe}_2 < \frac{\text{CO.N}(C_7 \text{H}_7)}{\text{NH.CS}}$. [175°]. Formed from o-tolyland a-amido-isobutyric thiocarbimide (Marckwald, B. 24, 3284). Needles, v. sol. hot alcohol.

p-Tolyl-di-methyl-thiohydantoïn. Formed in like manner from p-tolyl-thiocarbimide. Crystals, v. sol. most solvents.

o-TOLYL-METHYL-THIO-UREA

NHMe.CS.NHC,H,. [153°]. Formed from methyl-thiocarbimide and o-toluidine in alcohol (Dixon, C. J. 55, 622). Pearly crystals, m. sol. hot water, sol. alcohol.

p-Tolyl-methyl-thio-urea. [126°]. Formed in like manner from p-toluidine. Vitreous prisms,

v. sol. hot water.

Reference.—TOLYL-THIO-UREA

o-TOLYL-(a)-NAPHTHYLAMINE C,,H,,N i.e. C₁₀H, NH.C₆H, Me. [95°]. Formed by heating (a)-naphthol (1 mol.) with o-toluiding (2 mols.) and CaCl₂ (1 mol.) in sealed tubes at 290°, the yield being 37 p.c. of the theoretical amount (Friedländer, B. 16, 2084). Flat needles, v. sol. alcohol and ether.

o-Tolyl-(8)-naphthylamine $C_{10}H_7NH.C_6H_4Me$. [96°]. Formed by heating a mixture of (β)-naphthol with o-toluidine and CaCl, at 280°. Silvery plates (from ligroin), v. sol. alcohol.— B'2C,H,N,O,. [110°]. Reddish-brown needles. Benzoyl derivative [118°]. Plates.

p-Tolyl-(a)-naphthylamine $C_{17}H_{18}N$. [79°]. (360° at 528 mm.). Formed by heating (a)-naphthol with p-toluidine and CaCl, at 280°, the yield being 50 p.c. Short prisms (from alcohol). Its solutions exhibit blue fluorescence.

p-Tolyl-(β)-naphthylamine. [104°]. Formed by heating p-toluidine hydrochloride with (8)-naphthol for 8 hours at 200° (Witt, B. 20, 578; cf. Friedländer, B. 16, 2078). Plates (from alcohol). Its solutions have violet-blue fluor-escence. Dissolved in isosmyl alcohol it is reduced by Na to a hydride [44°] (Bamberger a. Müller, B. 22, 1309). With nitroso-dimethylaniline hydrochloride and HOAc it yields a saffranine C₁₀H₆N₂Ol(O,H₇):C₂H₂NMe₂ (Witt, B. 21, 724). p-Tolyl-naphthyl-amine (10 g.), heated with nitroso-dimethylaniline hydrochloride (4 g.), ZnCl₂ (10 g.), and HOAc (100 c.c.), forms $C_{2e}H_{12}(NH.C_eH_4Me)$, [225°], which yields an acetyl derivative [225°].

Acetyl derivative [85°]. Needles Benzoyl derivative [189°]. Needles. Reference. - TETRA-BROMO-TOLYL-NAPHTHYL-

DI-p-TOLYL-NAPHTHYLENE-DIAMINE
C₁₀H₄(NHC₁H₂), [237°]. Formed by heating di-oxy-naphthalene with p-toluidine and p-tolu-

idine hydrochloride (Annaheim, B. 20, 1873). Needles (from xylene). Gives a red colour on heating with solid ZnCl.

o-TOLYL-(a)-NAPHTHYL-THIO-UREA

CS(NHC,eH,)(NHC,H,). [168°]. Formed by the action of o-tolyl thiocarbimide on (a)-naphthylamine or of (α)-naphthyl thiocarbimide on o-toluidine (Mainzer, B. 15, 1416). Short needles, sol. alcohol. Split up by conc. HClAq at 150° into o-toluidine, (a)-naphthylamine, o-tolyl-thiocarbimide, and (a)-naphthyl thiocarbimide.

o - Tolyl - (β) - naphthyl - thio - urea [194°]. Formed from o-tolyl thiocarbimide and (β) -naph-

thylamine (M.).

p - Tolyl - (α) - naphthyl - thio - urea Formed from p-tolyl thiocarbimide and (α)-naphthylamine (M.). Slender white needle-.

p - Tolyl - (β) - naphthyl - thio - urea Formed from p-tolyl thiocarbimide and (β) -naphthylamine (M.). Decomposed by HClAq at 150° into p-toluidine, (β) -naphthylamine, p-tolyl thiocarbimide, and (β) -naphthyl thiocarbimide.

p-TOLYL p-NITRO-BENZYL OXIDE

C₆H₄Me.O.CH₂·C₆H₄·NO₂. [91°]. Formed from p-nitro-benzyl chloride and alcoholic C,H,ONa (Frische, A. 224, 144). Yellowish plates, sol. ether.

TOLYL NITROSO-METHYL KETONE v. Oxim of Tolyl-Glyoxal

TOLYL-OXAMIC ACID v. vol. iii. p. 654. TOLYL-OXAMIDE v. OXALIC ACID.

o-TOLYL-OXAZOLINE CH2.O C.C,H7.

(255°). Formed by dissolving o-toluic bromo-ethyl-amide in hot water and adding alkali (Salomon, B. 26, 1322). Oil. Evaporating with aqueous HCl (1 mol.) forms the compound C,H, CO.O.CH, CH, NH, while excess of HCl

gives C,H,.CO.NH.CH,.CH,.CH. — B'C,H,N,O,. [145°].—B'2H,PtCl,. [189°].

p-Tolyl-oxazoline C,H,INO. [66°]. (265°).

Formed by decomposing C,H,.CO.NH.CH,.CH_Br with alkali. Needles. B'C, H, N,O,. [188°]. -

B'2H2PtCl6. [186°].

DI-0-TOLYL OXIDE (C.H.Me)2O. ether. (272°-278°). S.G. 24'8 1.047. A product of the distillation of Al(OC,H,Me), (Gladstone a. Tribe, C. J. 49, 27). Oil, smelling like geranium leaves.

Di-m-tolyl oxide (C,H,),O. (284°-288°). 1. 1.5576 at 16°. V.D. 193.9. A product of distillation of aluminium thymol Al(OC_{1e}H₁₈)₃ (Gladstone a. Tribe, C. J. 41, 13). Oil.

Di-p-tolyl oxide (C,H,)2O. [50°]. (270°-800°). Formed by distilling aluminium p-cresol (G. a.

T.). Prisms (from alcohol).

Isomeride [165°]. Fo Formed by heating p-cresol with ZnCl, (Busch, B. 17, 2638). Crystals, volatile with steam.

DI-TOLYL-OXINDOLE C.H. $\langle NH, NH, NH \rangle \rangle$ CO

Toluisatin. [200°]. Formed by adding toluene to a solution of isatin in H₂SO₄ (Baeyer a. Lazarus, B. 18, 2638). Needles, v. sol. alcohol and ether, sol. aqueous alkalis. Yields an acetyl derivative C₂₂H₁₆O(NAc) [148°] and an ethyl derivative C₃₂H₁₆O(NEt) [108°], both being crystalline.

TOLYL-OXY-ACETIC ACID v. Tolyl derivative of GLYCOLLIC ACID and OXY-TOLYL-ACETIC ACID.

p-TOLYL OXY-ETHYL SULPHONE C,H, SO, CH, CH, OH. [55°]. Got from sodium toluene p-sulphinate and glycol chlorhydrin (Otto, J. pr. [2] 30, 355). Needles (from alcohol) with bitter taste. Heating with KOHAq gives C₁₈H₂₂S₃O₄ [84°

Benzoyl derivative C,H,.SO2.C2H4.OBz.

[176°]. Needles, sl. sol. hot alcohol.

DI-o-TOLYL-PARABANIC ACID C17H14N2O3 **6.e.** $CO < N(C,H_1).CO \\ N(C,H_2).CO$ [208°]. Formed by boiling an alcoholic solution of oxalyl-di-tolyl-guanidine with cone. HClAq (Berger, B. 12, 1856). Small needles.

Di-p-tolyl-parabanic acid [144°]. Formed by boiling di- (or tri-) p-tolyl-guanidine dicyanide with alcohol and HCl (Landgrebe, B. 11, 978). Plates. Oxidised by potassium permanganate to

CO (N(C,H,CO,H).CO [159° ?]. P-TOLYL PENTADECYL KETONE

 $C_{15}\bar{H}_{31}.CO.C_6H_4Me.$ [60°]. (262° at 15 mm.). Formed from palmityl chloride, toluene, and AlCl, (Kinfft, B. 21, 2266). Needles (from

ohol). Yields p-toluic acid on oxidation.
o-TOLYL-PENTHIAZOLINE C₁₁H₁₈NS i.e. C,H,C,S-CH₂. Got by heating trimethylene N.CH, CH, chlorobromide with thio-o-toluic amide (Pinkus, B. 26, 1081). Oil, volatile with steam. Its hydrochloride gives a white compound with HgCl2.

 $p ext{-} ext{Tolyl-penthiazoline}$ [53°]

p-tolyl-phentriažinė. Dihydride C₆H₄ N=N (151°). Formed by the action of HNO2 on o-amido-benzyl-p-toluidine hydrochloride (Busch, B. 25, 450). Yellow plates (from alcohol).—B'HCl.—B',H,PtCl.. [190°].— $(C_3H_3N_3O_7$. [132°]. Dark-yellow needles. p-TOLYL-PHENYL-ACETAMIDINE $C_{1p}H_{1p}$. B'C,H,N,O,

i.e. C.H., CH2.C(N.C.H.Me).NH2. [119°]. Formed by heating phenyl-acetonitrile with p-toluidine hydrochloride (Bernthsen, A. 184, 342). Crystals (from alcohol).— $B'_2H_2PtCl_4$: yellow prisms. DI - p - TOLYL - DIPHENYLACETYLENE -

DIAMINE C28H21N2 i.e. C,H,N:OPh.CPh:NO,H, [161°]. Formed, together with Bz.CPh:NC,H, [116°], by heating benzil (1 mol.) with p-toluidine (2 mols.) at 130° (Bandrowski, M. 9, 690). Triclinic plates.

DI - TOLYL - PHENYLENE -DIAMINE v. PHENYLENE-DI-TOLYL-DIAMINE.

TOLYL PHOSPHATES.

Tri-o-tolyl-phosphate (C,H,O),PO. Formed by heating o-cresol with POCl, the yield being 95 p.c. (Heim, B. 16, 1767; Rapp, A. 224, 173). Oil, sol. alcohol and ether.

p-Tolyl phosphate PO(OC,H,)(OH). [116°]. Formed by the action of water on PO(OC,H,)Cl, which is a product of the action of POCL on p-cresol (Rapp, A. 224, 168). Plates, sol. alcohol, ether, and water.

Tri-p-tolyl phosphate PO(OC,H,),

Formed by heating p-cresol with POCl, the yield being 95 p.c. Colourless tables, insol. Aq. p - TOLYL - PHOSPHINE C.H.NH. [4°]. (178°). Formed by the action of AlCl, on a mixture of toluene and PCl, the resulting C.H.,PCl₂ [20°] (245°), being converted by water into C.H.,P(OH)₂, which splits up on heating into tall heavy in the splits up on heating into tolyl-phosphine and toluene phosphonic acid C₇H₄PO₃H₂ (Michaelis, B. 12, 1009; 13, 653; A. 212, 230). Oil with strong odour. Rapidly oxidised by air to C,H,P(OH)2. HI forms C,H,PH,I, crystallising in colourless needles.

p - TOLYL - PHTHALIDE $C_{15}H_{12}O_{2}$ $C_eH_i < \frac{CO}{CH(C,H_2)} > 0$. [129°]. Formed by reducing C,H,.CO.C,H,.CO2H in alcoholic solution with Zn and HCl (Gresly, A. 234, 235). Needles, sol. hot alcohol, insol. NH, Aq.

Di - p - tolyl - phthalide $C_0H_4 < \frac{CO}{C(C_1H_1)_0} > 0$. [116°]. Formed by heating phthalyl chloride (100 g.) with toluene (450 g.) and AlCl, (Berchem, Bl. [2] 42, 168; cf. Friedel a. Crafts, Bl. [2] 85, 505). Prisms, sol. alcohol and ether.

TOLYL - PHTHALIMIDE v. Tolylimide of

PHTHALIC ACID

m-TOLYL-PROPIOLIC ACID C,H,.C:C.CO2H. [109.5°]. Formed by boiling di-bromo-tolyl-propionic acid C,H,CHBr.CHBr.CO,H with alcoholic potash (Müller, B. 20, 1215). AgNO. gives an explosive white pp.
0-TOLYL-PROPIONIC ACID C.H., O.H., O.

Monoclinic prisms (Young, B. 25, 2102).

m-Tolyl-propionic acid C,H,CH,CH,CH,CO,H. [43°]. Formed by reducing m-tolyl-acrylic acid with sodium-amalgam (Müller, B. 20, 1214; von Miller, B. 23, 1899). Needles (from ligroin), sol. water, alcohol, and ether. Volatile with steam. m-Tolyl-propionic acid [125°]. Formed by oxidation of m-isobutyl-toluene by HNO, (Effront, B. 17, 2330). Needles, sl. sol. hot water.—AgA': crystals, sol. hot water.

p-Tolyl-propionic acid. [116°] (Von Miller, B. 23, 1898); [120°] (Kröber, B. 23, 1033). Formed by reducing p-tolyl-acrylic acid. Condensed by hot H₂SO₄ to oxy-methyl-indonaph-

Di-α-tolyl-propionic acid CH₂.C(C,H₂)₂.CO₂H. [152°]. Prepared by adding toluene to a solution of pyruvic acid in H₂SO₄ at -10° (Böttinger, B. 14, 1596; 15, 1474). Monoclinic crystals, sol. hot alcohol. Give u-di-tolyl-ethane on distillation with lime. Oxidised by KMnO, to di-phenyl-ethane tri-carboxylic acid.—AgA'.

Ethyl ether EtA'. [145°]. Prisms. References. - Bromo-, Nitro-, and Oxx-, TOLYL-PROPIONIC ACIDS.

p-TOLYL-PROPIONIC ALDEHYDE

C,H,CH,CH,CHO. S.G. 13 .9941. (228°). Formed by the action of water on the compound of CrO₂Cl₂ with the cymene got from camphor and P₂O₄ (Richter a. Schüchner, B. 17, 1981). Got also by distilling calcium p-tolyl-propionate with calcium formate (Miller, B. 23, 1082). Oil, smelling like peppermint. Volatile with steam. Yields p-toluic and terephthalic acids on oxida-

DI-0-TOLYL-PROPYLENE-DIAMINE

CHMe(NHC,H.,).CH.,NHC,H., (280° at 120 mm.). Got by heating o-toluidine with propylene bromide (Bischoff, B. 25, 8276).

Acetyl derivative C₂₁H₂₂N₂O₂. [120°].

Di-p-tolyl-propylene-diamine C.H. (NHC,H.). (277° at 48 mm.) (Bischoff, B. 25, 3277). Got from p-toluidine and C.H.Br.

3 x 2

Acetyl derivative C₂₁H₂₈N₂O₂. [114°]. Benzoyl derivative C₃₁H₃₀N₂O₂. [152°]. o-TOLYL-PROPYLENE-4-THIO-UREA

 $C_7H_7NH.C \leqslant_{N.CH_2}^{S CHMe}$. $[126^{\circ}].$ Formed from s-o-tolyl-allyl-thio-urea and HCl (Prager, B. 22, 2999). Tables, sl. sol. hot water.—B'2H2PtCl8. [178°]. Prisms.—B'C, H, N, O, [176°]. Cubes.

DI-o-TOLYL-PROPYLENE UREA

 $C_3H_6 < \frac{NC_7H_7}{NC_7H_7} > CO$. [93°]. Formed from di-otolyl-propylene-diamine and COCl2 (Bischoff, B. 25, 3276). Prisms (from dilute alcohol).

Di-p-tolyl-propylene-urea. [130°]. d-TOLYL ISOPROPYL KETONE

C.H.Me.CO.Pr. (236°). Formed from toluene, isobutyryl chloride and AlCl₃ (Claus, J. pr. [2] 46, 480). Oil, with bitter taste and aromatic odour. Yields p-toluic and terephthalic acids on oxidation. Yields an oxim [92°]. May be reduced to p-tolyl-isopropyl-carbinol; an oil boiling above 300°

DI-0-TOLYL-PYRAZINE HEXAHYDRIDE Di-o-tolyl-

piperazine. [171°]. Formed from ethylene bromide, o - toluidine, and Na₂CO₃ at 160° (Bischoff, B. 22, 1781; 23, 1982; cf. Mauthner a. Suida, M. 7, 233). Crystals (from ether).

Isomeride. [154°]. Formed by heating dio-tolyl-ethylene-diamine with chloro-acetic acid and NaOAc (Bischoff, B. 23, 2031). Thin needles (from alcohol).

Di-p-tolyl-pyrazine hexahydride

C,H,N CH2 N.C,H, [188°]. Formed from p-toluidine and C,H,Br, in presence of NaOAc or Na₂CO₃. Prisms, sl. sol. alcohol. With NaNO2 and HOAc it yields a di-nitrosoderivative [167°].

 $_{\rm CH} < _{\rm CH:N}^{\rm CH.N.C,H,Me}$ o-TOLYL-PYRAZOLE

(246.5° cor.). S.G. & 1.0868; 13 1.0753. Formed by boiling epichlorhydrin (1 mol.) with o-tolylhydrazine (2 mols.) and benzene, boiling off the benzene, and heating the residue at 150° (Balbiano, G. 18, 368). Not reduced by boiling with alcohol and Na.—B'2H2PtCl8. [201°].—B'EtI.

[100°]. Needles, v. e. sol. water. Dihydride $C_3H_5N_2\cdot C_7H_7$. (271°). Formed from acrolein tolyl-hydrazide and dilute (2 p.c.) H_2SO_4 . Oil. Sol. fuming HClAq, but reppd. by water. $K_2Cr_2O_7$ and

H2SO, give an indigo-blue colour.

p-Tolyl-pyrazole. (259° cor.). [33°]. Formed, in like manner, from epichlorhydrin and p-tolyl-hydrazine. Yellowish plates, sl. sol. hot water.-B'2H2PtCl 2aq: needles. [184°].-B'EtI. [105°]. Prisms, v. sol. water.

Dihydride. [60.5°]. (282° i.V.). Formed from acrolein p-tolyl-hydrazide and dilute H2SO. Got also by reducing p-tolyl-pyrazole in alcoholic solution by Na. Iridescent plates, sl. sol. hot water. Its acidified solution is coloured violetred by K,Cr,O,

p-TOLYL-PYRIDINE HEXAHYDRIDE

C.H.Me.N.C.H. (262°). Formed by heating piperidine with p-bromo-toluene at 270° (Lellmann a. Just, B. 24, 2099). V. sol. alcohol and benzene.

p-TOLYL-PYRROLE C,H,N i.s.

CH:CH>NC_eH,Me. Got by dry distillation of p-toluidine mucate (Lichtenstein, B. 14, 933). Converted by AcCl into crystalline C,H,NC,Ac,. -B',HgCl2.

Di-p-tolyl-pyrrole CH:C(C,H,) NH.

[197°]. Formed by treating di-p-tolyl-furfurane or (C,H,CO),C,H, with NH,OAc (Hollemann, R. T. C. 6, 73). Small plates with bluish lustre. Does not form a K derivative. Boiling HClAq gives a red resin. With isatin, after several days, it gives an intense red colour.

o-TOLYL-QUINAZOLINE DIHYDRIDE

 $C_{15}H_{11}N_2$ i.e. $C_6H_4 < \stackrel{N=CH}{<}_{CH_2.N.C_6H_4Me}$. Formed by reducing the formyl derivative of o-nitro-benzyl-o-toluidine with tin and HClAq (Paal a. Busch, B. 22, 2701). Oil.—B'HSnCl.—B HCl: amorphous.—B',H,PtCl., [210°]. Needles (from alcohol_containing HCl).

 $[120^{\circ}].$ dihydride, p.Tolyl-quinazoline Formed in like manner. Plates, v. sol. alcohol. -B'HSnCl₃. [165°].—B'HCl 2aq. [85°].— 'HCl. [251°].—B'₂H₂PtCl₅. [216°¹. Yellow B'HCl. needles.

Methylo-iodide B'MeI. [186°]. Needles. p-Tolyl-quinazoline tetrahydride

C₆H₄ CH₂, NC₆H₄Me [127°]. Formed by reducing the dihydride in alcoholic solution by sodium. Needles, m. sol. alcohol. Yields a nitrosamine [100°]. Yields a crystalline hydro-

Reference.—OXY-TOLYL-QUINAZOLINE

(Py. 3)-o-TOLYL-QUINOLINE C16H13N i.e. C₆H₄CH:CH N=C.C₆H₄Me [77°]. Formed by distilling oxy-tolyl-quinoline (pseudoflavenol) with zinc-dust (Weidel a. Bamberger, M. 9, 108). Silky needles (from benzene).—B'2H2PtCl6 (dried at 105°). Orange tables.

m-Tolyl-isoquinoline CoH. CH: CH: N.

[52°]. Formed by heating (1)-chloro- (3)-mtolyl-isoquinoline with HI and P for three hours at 170° (Heilmann, B. 23, 3168). Needles (from MeOH).

p-Tolyl-isoquinoline C.H. CH:N.

[78°]. Formed in like manner (Ruhemann, B. 24, 3975). Needles.—B'HI.—B'H $_2$ CrO $_4$.—B' $_2$ H $_2$ PtCl $_6$.—B'C $_6$ H $_3$ N $_3$ O $_7$. Yellow needles.

Reference. -- OXY-TOLYL-QUINOLINE.

TETRA-m-TOLYL-SILICANE Si(C,H,)4. [151°]. (above 550°). S.G. 20 1·1188. Formed from m-bromo-toluene, SiCl,, and sodium (Polis, B. 19, 1021). Prisms (from ether), v. sol. benz-

Tetra-p-tolyl-silicane $Si(C_7H_7)_4$ S.G. 20 1.0793. Formed from (above 450°). SiCl,, p-bromo-toluene, ether, Na, and a little EtOAc (Polio, B. 18, 1542; 19, 1019). Monoclinic crystals (from chloroform). Decomposed by conc. HNO, forming silica and di-p-nitrotoluene $[72^{\circ}]$.

TETA A -o-TOLYL SILICATE Si(OC,H.),. (c. 437°). Formed by heating o-cresol with SiCl, the yield being 90 p.c. of the theoretical quantity (Hertkorn, B. 18, 1686). Oil, v. sol.

alcohol. Decomposed by hot water into silica and o-cresol.

Tetra-m-tolyl silicate Si(OC,H,)4. (445°)

at 720 mm. Syrup (H.).

Tetra-p-tolyl silicate Si(OC,H,)4. [70°]. (444°). Tables or prisms, v. sol. benzene (H.).

p-TOLYL SILICO-CHLORIDE C,H,SiCl₃. (219°). Formed from $Hg(C_1H_1)_2$ and SiCl, at 310° (Ladenburg, A. 173, 165). Liquid, converted by dilute aqueous ammonia into C,H, SiO.OH [c. 150°], which is converted at 200° into solid $(C_7H_7.SiO)_2O.$

TOLYL SULPHATES.

o-Tolyl sulphuric acid

[1:2] C₆H₄Me.O.SO₂.OH. Occurs in small quantity in the urine of horses (Preusse, H. 2, 355). Prepared from C₆H₄Me.OK and K₂S₂O₇ (Baumann, 3. 11, 1911). The K salt crystallises in plates, m. sol. water.

m-Tolyl sulphuric acid C6H4Me.O.SO2.OH. Occurs in very small quantity in horses' urine

p-Tolyl sulphuric acid C₆H₄Me.O.SO₂.OH. Occurs in the urine of horses and cows (Baumann, P. 9, 1389, 1716; Städeler, A. 77, 18). Formed by boiling potassium p-cresol with K₂S₂O₇. The K salt is sl. sol. cold alcohol.

DI-o-TOLYL-SULPHAZIDE C14H16N2SO2 i.e. Tolyl-[2:1] $C_0H_4Me.NH.NH.SO_2.C_0H_4Me$ [1:2]. Obtained by the toluene-sulphazide. [142°]. action of SO₂ upon o-diazo-toluene in alcoholic solution. White glistening needles (Limpricht, B. 20, 1241). When boiled with baryta-water it decomposes into o-toluene-sulphinic toluene, and nitrogen.

 $\begin{array}{cccc} \textbf{Di-p-tolyl-sulphazide} & \textbf{C}_{11}\textbf{H}_{16}\textbf{N}_2\textbf{SO}_2 & \textit{i.e.} \\ \textbf{[4:1] C}_{6}\textbf{H}_{4}\textbf{Me.NH.NH.SO}_{..}\textbf{C}_{6}\textbf{H}_{4}\textbf{Me} \textbf{[1:4]}. & \textit{Tolyl-toluene-sulphazide}. & \textbf{[140°]}. & \textbf{Formed} & \textbf{in} & \textbf{like} \\ \end{array}$ manner from p-diazo-toluene (L.). needles. By boiling with baryta-water it is decomposed into p-toluene-sulphinic acid, toluene,

and nitrogen.

Reference.—Nitro-di-tolyl-sulphazide.

DI-p-TOLYL SULPHIDE S(C₆H₄Me)₂. [57°]. Formed by distillation of lead (above 300°). p-tolyl-mercaptide (Otto, B. 12, 1175). Small needles, insol. water, sol. alcohol. Yields di-ptolyl sulphone on oxidation by KMnO.

Di-m-tolyl disulphide $C_{14}H_{14}S_2$ (C.H.Me) S2. Formed from m-tolyl mercaptan by treatment with dilute HNO. (Hubner a. Post, A. 169, 51). Liquid at -22°. Decomposed on boiling or by atmospheric oxidation of its ammoniacal alcoholic solution (Leuckart, J. pr. [2] 41, 189).

Di-p-tolyl disulphide $(C_0H_4Me)_2S_2$. Formed from p-tolyl mercaptan by atmospheric oxidation (Märcker, A. 136, 88) or by the action of ClSO₃H (Beckurts a. Otto, B. 11, 2066), or by treatment with ICy (Thurnauer, B. 23, 769). Formed also by the action of H2S on a warm dilute solution of toluene p-sulphinic acid in alcohol (Otto, J. pr. [2] 37, 211). Crystals, v. e. sol. ether. Begins to boil at 307° i.V. but decomposes, on distillation. Alcoholic K,S forms p-tolyl mercaptan (Otto a. Rössing, B. 19, **3129**)

Di p-tolyl tetrasulphide (C_eH₄Me)₂SO₄. [75°]. Formed by the action of H2S on a conc. solution of toluene p-sulphinic acid (Otto, J. pr. [2] 37, 211). Small plates, insol. water, v. sol. ether.

References.-DI-AMIDO-, DI-BROMO-, and OXY-

DI-TOLYL-SULPHIDE.

o-TOLYL SULPHOCYANIDE C.H.Me.S.CN. (c. 245°). Formed by adding cuprous sulphocyanide in KSCy to a solution of o-diazo-toluene (Thurnauer, B. 23, 770). Yellowish-red oil with unpleasant odour.

p-Tolyl sulphocyanide. (c. 248°). Formed in like manner, and also by passing CyCl through

alcohol containing lead p-tolyl mercaptide. Liquid, solidifying below 0°. DI-p-TOLYL SULPHONE (C.H.Me)2SO2. $\mathtt{DI} ext{-}p ext{-}\mathtt{TOLYL}$ $(C_6H_4Me)_2SO_2$ [158°]. (405°) at 714 mm. Prepared by the action of AlCl₃ on a mixture of toluene and toluene p-sulphonic chloride (Beckurts a. Otto, B. 11, 2068; 12, 1177; cf. Michael, B. 10, 584). Formed by the action of fuming H₂SO₄ or SO₅ on toluene (Deville, A. 44, 306; Otto a. Grüber, A. 154, 193) and by the action of KMnO, and HOAc on di-p-tolyl sulphide. Prisms (from benzene), sl. sol. cold alcohol and ether. Potashfusion yields p-cresol and diphenyl (Otto, B. 19,

DITOLYL DISULPHONIC ACID

[1:4:5] C₈H₃Me(SO₃H).C₈H₃Me(SO₃H) [5:1:4]. Formed from di-amido-ditolyl disulphonic acid by diazotisation followed by boiling with alcohol (Halle, A. 270, 363). Crystals, v. e. sol. water, m. sol. alcohol, insol. ether.—K₂A". Hexagonal plates.—KHA": prisms.—BaA" 5aq.

Chloride $C_{14}H_{12}S_2O_4Cl_2$. [229°]. Amide $C_{14}H_{12}S_2O_4(NH_2)_2$. Needles. p-TOLYL-SULPHONO-ACETIC ACID

C₆H₄Me.SO₂.CH₂.CO₂H. [118°]. Formed from sodium benzene sulphinate and chloro-acetic acid (Gabriel, B. 14, 834). Crystals (from benzene), sl. sol. hot water. Cl passed into its aqueous solution forms C,H,.SO,.CHCl, [114°]. Br forms C,H, SO, CHBr, and C,H, SO, CH, Br (Otto, J. pr. [2] 40, 542).—AgA': trimetric tables. Ethyl ether EtA'. Converted by bromine

(2 mols.) at 90° into C,H,.SO,.CBr,.CO,Et which on saponification by cold NaOHAq yields

C,H,,SO,CHBr, [117°].

p-TOLYL-SULPHONO-ACETONE
CH,CO.CH,SO,C,H,Me. [51°]. Formed by heating sodium toluene p-sulphinate with chloroacetone in alcohol (Otto, J. pr. [2] 36, 425). Needles, v. sol. alcohol. Bromine forms CH,Br.CO.CH,SO.C,H, [130°], m. sol. alcohol.

Di-p-tolyl-di-sulphono-acetone CO(CH₂·SO₂·C₆·H₄Me)₂. Formed by heating CH₂Br.CO.CH₂·SO₂·C₇·H₄, with sodium toluene p-Formed by heating sulphinate in alcoholic solution. Plates, v. sol.

hot HOAc and chloroform. DI-TOLYL DI - SULPHONO - DI - ETHYL -

AMINE NH(C₂H₄,SO₂C₂H₄)₂. Formed by heating di-tolyl ethylene disulphone with NH₄Aq (Otto, 7. pr. [2] 30, 359).—B'HCl. [201°].—

B'HAuCl.: dark-yellow needles

DI - TOLYL - DI - SULPHONO - DI - ETHYL OXIDE O(C₂H₄, SO₂, C₇H₇)₂. [84°]. Formed by the action of conc. KOHAq on C₂H₄(SO₂C, H₇)₂ and as a by-product in the preparation of C,H,SO2.C2H4OH [55°] by the action of dilute KOHAq on the same body (Otto a. Tröger, B. 26, 944; cf. J. pr. [2] 80, 171, 821).

TOLYL DISULPHOXIDE v. Tolyl ether of

TOLUENE THIOSULPHONIC ACID.

Di-p-tolyl sulphoxide SO(C₆H₄Me)₂. [92°]. Formed from toluene, SOCl₂, and AlCl₃ (Parker, B. 23, 1844). Flat needles (from ligroin), v. e. sol. benzene. Reduced by sodium and alcohol to di-tolyl sulphide [56°]. KMnO, in HOAc forms di-tolyl sulphone [158°].

DI-o-TOLYL-TETRAZINE C16H16N4 i.e.

C,H,N < N:CH NC,H,. [141°]. Formed from o-tolyl-hydrazine, chloroform, and alcoholic potash (Ruhemann, C. J. 57, 52). Crystals. orystalline C16H13(SO3H)N4 Yields

CleH₁₈(NO₂)N₄ [207°].—B'HCl: needles.

Methylo-iodide B'MeI. [198°]. Nodules.

Di-p-tolyl-tetrazine CleH₁₈N₄. Formed in like manner. B'HCl: very unstable white needles. Methylo-iodide B'MeI. [242°]. Needles.

o-TOLYL-THIAZOLE DIHYDRIDE $\text{C}_{\bullet}\text{H}_{\bullet}\text{Me.C} \underset{S.CH_{2}}{\overset{N.CH_{2}}{\bigcirc}}.$ (282°). Formed from thioo-toluic amide and ethylene bromide (Gabriel a.

Heymann, B.24, 786). Got also by heating o-toluic bromo-ethyl-amide with P2S, (Salomon, B. 26, 1329). Converted by bromine-water into taurine and o - toluic acid. - B'aH,PtCl. [199°].-

B'C.H.N.O.. [132°]. p-Tolyl-thiazole dihydride. [81°]. Prepared in like manner. Tables.

p-TOLYL-THIOBIURET C,H11N,S, [158°]. Formed by fusing p-toluidine (2 pts.) with persulphocyanic acid (1 pt.) (Tursini, B. 17, 584). Minute needles, sol. NaOHAq. Yields C₂H₁₀N₃S₂Et [134°], crystallising from alcohol in needles, and C.H. AcN.S. [166°], crystallising in needles. o-TOLYL-8-THIOCARBAMIC ACID

[1:2]C,H,Me.NH.CS.OH.

Ethyl ether C,H,N:C(SH).OEt. Tolyl-thiowrethane. Formed by heating o-tolyl-thiocarbimide with alcohol at 130° (Liebermann a. Natanson, B. 13, 1576; A. 207, 161). Oil, sol. alkalis. - C10H12AgNSO: white pp. insol. NH2Aq. Converted by MeI into C,H,N(OEt).SMe, and by Etlinto C.H.N(OEt). SEt, both being oils boiling above 250°

m-Tolyl-\$-thiocarbamic acid

[1:8]C,H,Me.NH.CS.OH.

Ethyl ether C,H,N:C(SH).OEt. 168°1. Formed in like manner (L. a. N.). Yields $C_{10}H_{12}AgNSO$ and oily $C_{10}H_{12}EtNSO$.

p.Tolyl-\$-thiocarbamic acid. Ethyl ether C.H.NC(SH).OEt. [87°]. Formed in the same way (L. a. N.). Triclinic crystals. Ammoniacal AgNO, added to its alcoholic solution ppts. C,H,NC(OEt).SAg, which is converted by MeI into C,H,NC(OEt).SMe, and by EtI into C,H,NC(OEt).SEt, both being oils boiling above 250°. C,H,NC(OEt).SEt is decomposed by dilute H2SO, at 180° into p-toluidine and thiocarbonic ether

o-Tolyl-a-thiocarbamic acid C,H,NH.CO.SH. Methyl ether MeA'. (70°]. Fo.med by heating C.H.NH.C(NC.H.). SMe with dilute H.SO. at 140° (Will a. Bielschow.ki, B. 15, 1317). Plates.

Ethyl ether EtA'. [60°]. Tables. p-Tolyl-a-thiocarbamic acid NHC,H,.CO.SH.

Methyl ether MeA'. [107°]. Formed by ating C.H.NH.C(NC,H.).SMe with dilute heating C,H,NH.C(NC,H,).SMe H,SO, at 170°. Needles, v. sol. alcohol and ether.

Ethul ather Eth! [799]. Formed from

p-toluidine and Cl.CO.SEt. Needles. EtSH and tolyl cyanate on distillation. Needles. Yields

Ethylene derivative

 $co<_S^{N(C,H,)}>c_2H$. Long slender [88°]. needles.

o-Tolyl-di-thio-carbamic acid

C,H,NH.CS.SH.—NiA', Brown needles, insol. water.—BaA', Plates (Losanitsch, B. 24, 3027). Forms di-o-tolyl-thio-urea on heating.

Methyl ether MeA'. [132°]. Fo from the Ba salt and MeI. White needles.

Ethyl ether EtA'. [72°]. Formed by heating C₇H₇NH.C(NC₇H₇).SEt with CS₂ at 160° (Will a. Bielschowski, B. 15, 1817). Prisms. Ethylene derivative C₁₀H₁₁NS₂ i.s.

 $CS < N(C_7H_7) > C_2H_4$. [129°]. Formed by heating

 $C_7H_7N.C < S_{NC.H} > C_2H_4$ with CS_2 at 200°. Yields

C₁₀H₁₁NS₂MeI, crystallising in prisms [151°]. m-Tolyl-di-thio-carbamic acid

C,H,NH.CS.SH.

Salts.—BaA'₂. Formed from m-toluidine, CS₂, and Ba(OH)₂ in alcohol (Losanitson, B. 24, 3027). Needles, sol. water, insol. cold alcohol.— NiA'2. Yellowish-brown plates.

Methyl ether MeA'. [89°]. Needles. p-Tolyl-di-thio-carbamic acid C,H,NH.CS,H. Salts.—NH,A'. Formed from p-toluidine, CS₂, alcohol, and NH₃Aq (Losanitsch, B. 24, 3026). Yellowish prisms, sol. water, insol. alcohol.—BaA'₂. Needles.—NiA'₂. Formed from *p*-toluidine, CS₂, ammoniacal NiSO₄, and alcohol. Brown needles, sol. alcohol, insol. water.

Methyl ether Med. [84°]. Prisms. Ethyl ether EtA. [74°]. Formed by heating NHC,H,C(NC,H,).SEt with CS₂ at 160° (W. a. B.). Needles, split up by heat into

p-tolyl thiocarbimide and EtSH.

 $CS < N(C,H,) > C_2H_4$ Ethylene ether [126°]. Crystals. Yields a methylo-iodide [107°] which is converted by aniline into C₁₆H₁₆N₂S [128°], and by o-toluidine into $C_{17}H_{18}N_2S^{\circ}[82^{\circ}]$.

Reference. - NITRO - TOLYL - THIOCARBAMIC

o-TOLYL-THIOCARBAZIC ACID o-Tolyl C,H,NH.NH.CS.SNH,NHC,H,. hydrazide Formed from o-tolyl-hydrazine, ether, and CS, (Preund, B. 24, 4200). Prismatic tables, insol. ether, decomposed by water and alcohol. The p-isomeride melts at 109°.

DI-o-TOLYL-THIOCARBAZIDE C15H18N4S i.e. CS(NH.NHC,H,Me). [130°]. Formed from o-tolyl-thiocarbazic acid and o-tolyl-hydrazine CS(NH.NHC,H,Me)2. at 90° (Preund, B. 24, 4201). Needles, sol. alcohol, insol. Aq

DI-p-tolyl-thiocarbaside. [121°]. Formed in like manner at 110° (Preund, B. 24, 4194).

Plates, v. sol. hot alcohol

Di-o-tolyl-thiosemicarbaside C18H17N2S i.e. Formed by NHC,H,.CS.NH.NHC,H,. [149°]. heating o-tolyl-thiocarbimide (1 mol.) with o-tolyl-hydrazine (1 mol.) in alcehol (Dixon, C.J. 61, 1017). Octahedra, m. sol. hot alcohol, insol. Aq.

Di-p-tolyl-thiosemicarbaside. [154°]. Formed in like manner (D.). Prisms, v. e. sol. hot alco-

op-Di-tolyl-thiosemicarbazide

[4:1]C₆H₄Me.NH.CS.NH.NHC₆H₄Me[1:2]. [142°]. Formed from p-tolyl-thiocarbimide and o-tolylhydrazine (D.). Prisms, sl. sol. hot alcohol.

Isomeride [2:1]C,H,Me.NH.CS.NH.NHC,H,Me[1:4]. [163°]. Formed from o-tolyl-thiocarbimide and p-tolylhydrazine (Dixon, C. J. 61, 1015). Prisms, m. sol. hot alcohol.

DI-o-TOLYL-THIOCARBAZONE C13H16N4Si.e. C,H,N:N.CS.NH.NHC,H,. [168°]. Formed from di-o-tolyl-thiocarbazide and alcoholic potash (Preund, B. 24, 4201). Blue-black needles, v. sol. chloroform, sl. sol. alcohol. Conc. H2SO4 forms a bluish-green solution.

Di-p-tolyl-thiocarbazone. [105°]. Formed in like manner. Dark-blue amorphous flakes.

o-TOLYL-THIOCARBIMIDE C,H,NS [1:2]C₆L,Me.N:CS. (239°). Formed from di-o-tolyl-thio-urea by distilling with P₂O₅ (Staats, B. 13, 136), by heating with aqueous H,PO. (Hofmann, B. 15, 986; Mainzer, B. 16, 2017), or by boiling with fuming HClAq (Girard, B. 6,

Reactions.—1. Chlorine forms C,H, NCl.CSCl (218°), which is converted by alcohol into a mixture of di-o-tolyl-urea and o-tolyl-carbamic ether (Lachmann, B. 12, 1349).—2. Chloro-acetic acid and alcohol at 150° form $C_7H_7.N:C < S.CH_2 O.CO$ crystallising in needles [120°], decomposed by boiling baryta-water into o-toluidine, CO2, and thioglycollic acid (Völtzkoff, B. 13, 1580).— 3. PhOMe and AlCl, on warming form 3. PhOMe and AlCl, on warming form C,H,NH.CS.C,H,OMe [95] (Tust a. Gattermann, B. 25, 3528). PhOEt and AlCl, give C, H,NSO

m-Tolyl-thiocarbimide $C_7H_7.N:CS$. at 732 mm. Formed by boiling di-m-tolyl-urea with conc. HClAq (Weith a. Landolt, B. 8, 719). Heavy oil. Converted by copper-powder at 220°

into m-toluic nitrile.

p-Tolyl-thiocarbimide [1:4] $C_6H_4Me.N:CS$. [26°]. (237°). Formed by heating di-p-tolylthio-urea with P.O. or aqueous H.PO. of S.G. 1.7 (Hofmann, B. 1, 173; 15, 986; Staats, B. 13, 135). Formed also, together with phenylthiocarbimide, by heating PhN:C:NC,H, with CS, at 190° (Huhn, B. 19, 2409). Needles (from

ether), v. sol. alcohol.

Reactions.-1. Ammonia forms tolyl-thiourea, and other bases act in like manner. 2. Bromine forms C,H,NBr.CSBr, which readily gives off Br, leaving (C,H,NCS),Br, crystallising in plates decomposing at 210° (Helmers, B. 20, 790).—2. Chlorine forms (C,H,NCS), Cl., whence dilute alcohol gives (C,H,NCS), O [189°] crystallising in needles.—3. CH_CLCO_H and alcohol at 150° form C,H,N:C (S.CH 162°) (Völtzkoff, B. 13, 1579) .- 4. Warmed with toluene and AlCl, it gives C,H,NH.CS.C,H, [166].— 5. Heated with AlCl, it yields (C,H,N.CS),28 [176°] (Friedmann a. Gattermann, B. 25, 3525). 6. C.H.OMe and AlCl. react, with formation of C.H.NH.CS.C.H..OMe [157°].—7. C.H.OEt and AlCl. give C.H.,NOS [151°], which may be oxidised to C.H.,NOS [170°].

DI-p-TOLYL DI-THIO-CARBONATE CO(SC_eH₄Me)₂. [91°]. Formed from p-tolyl mercaptan and COCl₂. It is also a by-product

in the action of p-diazo-toluene chloride on potassium xanthate (Leuckart, J. pr. [2] 41, 190). Needles (from alcohol).

TRI-TOLYL TRI-THIOCYANURATE (C_sH₄Me.SCy)_s. [114°]. Formed from sodium p-tolyl mercaptan and cyanuric chloride (Klason, J. pr. [2] 33, 120). Crystals (from HOAc).

o-TOLYL-THIOHYDANTOÏN C.H. NSO i.s $CS < N(C_6H,Me) > CO$. [136°]. Formed from o-tolyl-thiocarbimide and amido-acetic acid (Marckwald, B. 24, 8281). Plates, v. sol. alcohol, sl. sol. ligroïn.

p-Tolyl-thiohydantoin [210°] (M.). Formed as above (M.; cf. Aschan, B. 17, 426). Flat prisms, sol. alkalis.

Isomeride [183°]. Formed by fusing CH2Cl.CO.NHC, H, with thio-urea (P. Meyer, B. 10, 1966). Small crystals.

p-Tolyl-\psi-thiohydantoic acid C.H.,N:C(NH.).S.CH.,CO.H. [176° - 182°]. Formed by boiling chloro-acetic acid with ammonium sulphocyanide, p-toluidine, and alcohol (Jäger, J. pr. [2] 16, 21). Prisms, not attacked by Ac₂O and Br.

DI-p-TOLYL-THIOPHENE C, H, S $CH: C(C_eH_{\downarrow}Me) > S.$ $CH: C(C_eH_{\downarrow}Me) > S.$ Formed [171°]. bv heating di-p-tolyl-furfurane with P₂S₃ (Hollemann, R. T. C. 6, 74). Small plates (from alcohol). Gives an intense dark-green colour with isatin and H₂SO₄.

o-TOLYL-THIO-UREA NH2.CS.NHC6H4Me. [155°]. Formed from o-tolyl thiocarbimide and NH₂Aq (Staats, B. 13, 136). V. sol. hot water and alcohol, sl. sol. ether.

Acetyl derivative NHAc.CS.NHC,H,Me. [184°]. Formed from acetyl thiocarbimide and otoluidine in alcohol (Dixon, C. J. 55, 304). Pale lemon-yellow prisms, insol. water, sol. alcohol.

Benzoyl derivative NHBz.CS.NHC.H.Me. [119°]. Formed in like manner (Dixon, C. J. 55, 623). Pale-yellow

prisms, blackened by AgNO.
Di-o-tolyl-thio-urea CS(NH.C.H.Me). [158°]. Formed from o-toluidine and CS2 (Girard, B. 4, 985; Berger, B. 12, 1854; Ador a. Rilliet, B. 12, 2301). Long needles, v. sol. hot alcohol. Converted by heating with MeI into the hydroiodide of di-tolyl-methyl-thiourea C₁₆H₁₈N₂S, which may be represented as 'methyl di-o-tolylimido- thiocarbamate ' C,H,N:C(SMe).NHC,H, This body melts at 60°, while the corresponding ethyl derivative melts at 51°. The ethylene derivative [91°] may be represented by the formula $C_{7}H_{7}N:C<\frac{N(C_{7}H_{7})}{S}C_{2}H_{4}$ (Will a. Bielschowski, The isomeric compound 1316). [1:2] $C_6H_4Me.N:C < N(C_6H_4Me[1:4] > C_2H_4$ [82°] is got by boiling the methylo-iodide of the ethylene derivative of p-tolyl-di-thio-carbamic acid with o-toluidine.

m-Tolyl-thio-urea NHC,H,.CS.NH, [103°]. Formed from m-tolyl thiocarbimide and ammonia (Weith a. Landolt, B. 8, 719). Prisms m. sol. hot water, v. sol. alcohol and ether.

Di-m-tolyl-thio-urea CS(NHC,H₁)₂. [122°

Formed from m-toluidine, CS2, and alcohol (W.

a. L.). Needles, v. sol. alcohol, nearly insol. hot

p-Tolyl-thio-urea NHC,H,.CS.NH,. (C. a. W.); [182°] (S.). Formed by heating ptoluidine hydrochloride with ammonium sulphocyanide (De Clermont a. Wehrlin, C. R. 83, 347). Formed also from p-tolyl-thiocarbimide ammonia (Staats, B. 13, 136). Plates (from alcohol) with bitter taste, sl. sol. cold water, m. sol. alcohol.

Acetyl derivative NHC,H,.CS.NHAc. [176°]. Formed from acetyl sulphocyanide and p-toluidine (Miguel, Bl. [2] 28, 103). Needles,

v. e. sol. hot alcohol, v. sol. ether.

Di-p-tolyl-thio-urea $CS(NHC_6H_4Me)_2$. [176°]. Formed by boiling p-toluidine with CS₂ and alcohol (Sell, A. 126, 160). It is also a product of the action of allyl-thiocarbimide on an alcoholic solution of p-toluidine (Maly, Z. [2] 5, 258). Trimetric prisms (Levin, J. 1882, 384), insol. water, nearly insol. cold alcohol. alcoholic solution it is converted by HgO into di-p-tolyl-urea. In benzene solution COCl₂ forms $C_{16}H_{14}N_{5}SO$ [116°] (Will, B. 14, 1487). CSCl₂ forms $C_{16}H_{14}N_{2}S_{2}$ [109°] (Freund a. Wolf, B. 25, 1465). MeI forms C,H,N:C(NHC,H₇).SMe [1286], crystallising in needles, split up by heat into C(NC,H,), [60°] and HSMe, converted by heating with acids or alkalis into di-o-tolylurea and MeSH, and forming the salts B'HCl [173°] and B'H₂SO₄ [156°]. EtI forms the corresponding C₁,H₂₀N₂S [87°], while ethylene bromide gives $C_rH_rN:C < N(C_rH_r) > [112^\circ]$, which yields B'H₂SO, [194°] and a hydrochloride [219°] (Will a. Bielschowski, B. 14, 1492; 15, 1309).

References.—Nitro- and Oxy- tolyl-thio-

p-TOLYL - TOLUTRIAZINE DIHYDRIDE

C₁₅H₁₅N₃ i.e. CH : CH.C.N.CH₂ CMe: CH.C.N.NC₈H₄Me [178°]. Formed by heating o-amido-azo-toluene with formic paraldehyde and alcohol at 140° (Goldschmidt a. Poltzer, B. 24, 1008). Prisms, v. sol. hot alcohol.—B'HCl. [220°].—B'₂H₂PtCl_e. [216°]. Yellow crystalline pp. o-TOLYL-URAZOLE C₂H₂N₃O₂ i.e.

 $\mathbf{c}_{r}\mathbf{H}_{r}\mathbf{N} <_{\mathbf{CO.NH}}^{\mathbf{NH.CO}}$ [170°]. Got by heating otolyl-hydrazine hydrochloride with urea (Pinner, B. 21, 1219). White leaflets, v. sol. hot water.

p-Tolyl-urazole. [274°]. Yellowish needles, v. sl. sol. hot water.

o-TOLYL-UREA C,H10N2O i.e.

NH₂.CO.NH.C₆H₄Me [1:2]. [185°]. Formed from o-toluidine hydrochloride and potassium cyanate (Cosack, B. 13, 1089). Plates, sol. alcohol, ether, and hot water.

Bensoyl derivative CO(NHBz).NHC,H,. [210°]. Formed from o-tolyl cyanate and benzamide at 125° (Gattermann a. Cantzler, B. 25, 1088). Needles.

m-Tolyl-urea NH2.CO.NHC6H4Me [1:3]. [142°]. Formed from m-toluidine hydrochloride and potassium cyanate (Cosack, B. 12, 1450; 13, 1089). Needles or tables (from alcohol) or plates (from water).

NH2.CO.NHC6H4Me [1:4]. p-Tolyl-urea [172°] (C.); [180°] (St.; P.). Formed in like manner (C.; cf. Sell, C.J. 16, 190). Formed also by the action of p-toluidine on mercuric fulminate (Steiner, B. 8, 519), and also from toluamidoxim, benzene sulphonic acid, NaOĤ, and CHCl, (Pinnow, B. 24, 4167). Needles, sol. hot water, v. sl. sol. ligroin.

[246°] (B.); Di-o-tolyl-urea CO(NHC,H,)2.

[254°] (W.); [256°] (Barr, B. 19, 1769). Formation.—1. By the action of alcohol or water at 100° on the dichloride of o-toluic nitrile (Lachmann, B. 12, 1349).—2. From otoluidine hydrochloride and cyanamide (Berger, B. 12, 1859).—3. From tolyl cyanate and water (Nevile a. Winther, B. 12, 2325).—4. From o-toluidine and COCl₂ (Girard, B. 6, 444).—5. By heating o-toluidine with urea (G.).—6. A product of the distillation of o-tolyl-amido-acetic acid (Widman, J. pr. [2] 38, 303).—7. By the action of Ac₂O and benzene on the o-toluide of o-tolyl-imido-diacetic acid (Bischoff, B. 23, 1995).

Properties.—Needles (from HOAc), insol. ether, sl. sol. alcohol. Not volatile with steam.

Di-m-tolyl-urea CO(NH.C₆H.Me)₂. [203°] (G. a. C.); [217°] (C.). Formed from m-tolyl cyanate and m-toluidine (Gattermann a. Cantzler, 25, 1089), by the action of ClCO₂Et on m-toluidine, by heating m-tolyl-carbanic ether with water, and by heating m-tolyl-urea with m-toluidine at 160° (Cosack, B. 12, 1450; 13,

1090). Needles (from alcohol), insol. water. Di-p-tolyl-urea CO(NH.C₆H₄Me)₂. [255°]. Formation.—1. By boiling an alcoholic solution of di-p-tolyl-thio-urea with HgO as long as HgS is formed (Sell, C. J. 16, 190; A. 126, 161).—2. By passing COCl₂ into p-toluidine dissolved in chloroform (Michler, B. 9, 710; Kühn, a. Henschel, B. 21, 505).—3. By heating tolylurea with p-toluidine at 160° (Weith, B. 9, 821). 4. From p-tolyl-cyanate and p-toluidine (Gattermann a. Cantzler, B. 25, 1089).

Properties.—Needles, insol. water, sl. sol. cold alcohol.

Tri.p-tolyl-urea $C_{22}H_{22}N_2O$. [189°]. Got from $(C_1H_1)_2N$.COCl and p-toluidine (Hammerich, B. 25, 1822). Needles, sol. warm benzene, insol. ether. Gives a benzoyl derivative [137°].

Tetra-p-tolyl-urea $C_{20}H_{28}N_2O$. [80°]. Got m $(C_7H_7)_2N$.COCl and di-p-tolyl-amine.

Needles, v. e. sol. alcohol.

References .- AMIDO., NITRO-, and OXY-TOLYL-UREA.

TOLYL-XYLIDINE C, H17N i.e.

C₆H₄Me.NH.C₆H₃Me₂. [70°]. (c. 300° at 487 mm.). Silky needles (from alcohol) (Girard a. Vogt, Bl. [2] 18, 69).

o-TOLYL m-XYLYL KETONE C18H18O i.e. [1:2] $C_sH_sMe.CO.C_sH_sMe_s$ [1:2:3]. (330°) at 728 mm. Formed from o-toluic chloride and (330°) m-xylene (Smith, B. 24, 4050). Oil. Converted by hydroxylamine at 120° into a mixture of anilides.

TONKA BEANS, the fruit of Dipterix odorata, contains coumarin (q. v.) (Boullay a. Boutron-Charlard, J. Ph. 11, 426; [8] 7, 160).
TRAGACANTH GUM. Exudes from Astra-

galus verus, a tree growing in Armenia and the north of Persia. It swells up in water, about one-half of it dissolving. It contains arabin, bassorin, starch, and water.

TRANSPIRATION. Graham, in 1846 and 1849, applied the term 'transpiration' to the passage of gases through capillary tubes into a vacuum (v. T. 1846. 573; 1849. 349). The rates of transpiration of different gases bear constant relations to each other; but these rates have not been connected in any definite ways with the compositions of the gases. M. M. P. M.

TREE GUM v. XYLAN. TREHALOSE v. SUGAR.

TREHALUM $C_{24}H_{42}O_{21}$. S. ·06 at 17°; 1·8 100°. [a]_D = +179°. Occurs in *Trehala* manna (Scheibler a. Mittelmeier, B. 26, 1331). Tasteless, minute prisms. Very hygroscopic. Does not reduce Fehling's solution. Does not react with phenyl-hydrazine. Boiling dilute H₂SO₄ forms glucose. At 180° it slowly forms trehalin, which is v. sol. water and ppd. by alcohol. Trehalin dissolves in phenyl-hydrazine. Iodine colours trehalum and trehalin violet. Diastase, yeast, and invertin have no action. Ac₂O and NaOAc yield an acetyl derivative [above 240°].

TRI-. In the alphabetical arrangement of this dictionary the prefix 'tri,' when indicating the presence of three radicles, is treated as if it did not form part of a name, except where the en-

tire name is numerical, as in 'tridecane.'
TRIANOSPERMIN. A crystalline substance occurring in Trianosperma ficifolia, a climbing plant of Brazil (Peckholt, Ar. Ph. [2] 113, 104; Parodi, Ph. [3] 10, 667). It has a pungent taste, is alkaline in reaction, sol. water, alcohol, and ether, and is ppd. by Pb(OAc)2 and PtCl4.

TRIAZOLE N\left\(\bigcolemn{c} \text{CH:N} \\ \text{CH:NH} \\ \end{c} \end{c}. Pyrrodiazole. [120.5°]. (260°). V.D. 2.39 (calc. 2.49). Formed by heating its carboxylic acid alone or with boiling water (Andreocci, B. 25, 229; Bladin, B. 25, 745). Needles (from ether), v. sol. water and alcohol, may be sublimed. The di-oxy-derivatives of alkyl-triazoles $N \leqslant_{C(OH),NR}^{C(OH);N}$, named by Pinner 'urazoles,' are obtained by heating hydrazines with urea (Pinner, B. 20, 2358).

TRIAZOLE CARBOXYLIC ACID

·CH ——N $N \leqslant \overline{C(CO_2H)}.\overline{NH}.$ [137°]. Obtained by oxidising methyl-triazole (Andreocci, B, 25, 229) or amido-phenyl-triazole carboxylic acid and alkaline KMnO₄ (Bladin, B. 25, 744). Amorphous powder.—CuA'₂ aq. Green crystalline powder.

TRICARBALLYLIC ACID v. vol. i. p. 679.

TRICOSANE v. TRI-ICOSANE.

TRIDECANE $C_{18}H_{28}$. [-6°]. (234°). S.G. ·771; \$ ·761. Formed by reduction of methyl dodecyl ketone or of tridecoic acid by HI and P (Krafft, B. 15, 1699). Formed also by distilling barium myristate with NaOMe (Mai, B. 22,

TRIDECOIC ACID C₁₂H₂₅.CO₂H. [41°]. (236° at 100 mm.). Formed by oxidation of methyl tridecyl ketone (Krafft, B. 12, 1669). Crystals.

AgA'.

Amids C₁₂H₂₂.CO.NH₂. [98.5°]. Formed by digesting the nitrile with conc. H₂SO₄ and pouring into water (Lutz, B. 19, 1439). Plates (from alcohol).

Nitrile C₁₂H₂₃CN. (275°). tridecylamine; Br, and NaOH. (275°). Formed from Oil, v. sol.

alcohol and ether.

TRIDECYL ALCOHOL (C.H.,3)2CH.OH. [42°]. Formed by reducing di-hexyl ketone (Kipping, C. J. 57, 536). Plates (from dilute alcohol), insol. water. May be distilled. HBr forms

(C₆H₁₃)₂CHBr [39°]. TRIDECYLAMINE C₁₃H₂₇NH₂. [87°]. (265°). Formed by boiling myristyl-tridecyl-urea with KOHAq (Lutz, B. 19, 1437). Unctuous mass, v. sol. alcohol and ether. Absorbs water and CO₂ from the air.—B'HCl: needles, decomposing at 100°.—B'2H2PtCl8.—B'2H2SO4: needles, insol. cold water.

TRIDECYLENE C₁₅H₂₆. (233° cor.). S.G. 2 ·8445. Occurs in petroleum from Burmah

(Warren a. Storer, Z. 1868, 232).

TRIDECYL-UREA. Myristyl derivative C₂₈H₃₆N₂O₂ i.e. C₁₃H₂₇NH.CO.NHC₁₁H₂₇O. [103°]. Formed by the action of Br and KOHAq on the amide of myristic acid (Reimer a. Will, B. 18, 2016). Crystals (from alcohol), almost insol. cold water, sol. ether.

TRIGENIC ACID v. ETHYLIDENE-BIURET. TRIGONELLIN C,H,NO2 i.e.

CH_C -co

. Occurs in the seeds of CH:NMe Trigonella Fænum-græcum (Jahns, B. 18, 2518; 20, 2840). It is also produced by saponification of the product of the reaction of MeI on potassium pyridine (β) -carboxylate (Hantzsch, B. 19, 31). Colourless prisms (containing aq), converted by HClAq at 265° into pyridine (\$\beta\$)-carboxylio acid (q. v., Reaction 4).—B'HCl.—B',H,PtCl.,—B'HAuCl., [198°].—B',48HAuCl., [186°]. Slender needles

n-TRI-ICOSANE C₂₃H₄₈. [48]. (234° at 15 mm.). S.G. 46° 7785; 31° 7570. Formed from laurone (C11H23)2CO by treatment with PCl5 and reduction of the resulting (C₁₁H₂₃)₂CCl₂ with HI and P at 240° (Krafft, B. 15, 1712). Obtained also by fractional distillation of paraffin oil from brown coal (Krafft, B. 21, 2263). Glittering plates (from ether-alcohol), sl. sol. alcohol.

TRI-ICOSANE DICARBOXYLIC ACID

C₂₃H₄₆(CO₂H)₂. [102·5°]. Formed by heating di-oxy-penta-icosylene with soda-lime (Stärcke, A. 223, 300). Flocculent pp. (from ligroïn-ether).

—PbA".

TRI-ICOSYL ALCOHOL (C11H23)2CH.OH. [76°]. Formed by reducing laurone with Na and water (Kipping, C. J. 57, 983). Plates (from ether), insol. water. Yields an acetyl derivative (C₁₁H₂₃)₂CH.OH [35°]

TRI-ICOSYL ALCOHOL C23H47.OH. [82°]. A wax-like body extracted by hot alcohol from flax fibres (Cross a. Bevan, C. N. 59, 135). Yields an acetyl derivative [65°].

TRIMELLITIC ACID v. vol. iii. p. 201. TRIMESIC ACID v. vol. iii. p. 230.

TRIMESITIC ACID v. Pyridine TRICARB-OXYLIC ACID.

TRIMETHYLENE v. vol. iii. p. 304

TRITICIN $C_{12}H_{22}O_{11}$. $[a]_{D}=-43.6^{\circ}$ (Reidemeister, J. Th. 1881, 69). Extracted by dilute alcohol from the root of couch-grass (Triticum repens) (H. Müller, Ar. Ph. [3] 2, 500; 3, 1). Tasteless hygroscopic powder, v. sol. water, insol. alcohol and ether. Leworotatory. On boiling with water, especially in presence of acids, it changes into lævulose. Its solutions are not ppd. by metallic salts, nor coloured by iodine.

TROPECLINES v. Azo- compounds.
TROPEINES. A name given by Ladenburg to alkoyl derivatives of tropine (q. v.).

TROPIC ACID v. OXY-PHENYL-PROPIONIC

TROPIDINE v. METHYL-ETHYLENE PYRIDINE

(a)-Methyl-tropidine C_bH₁₅N i.e. CH₂:CH.CH:CH.CH:CH.CH₂NMe₂ (Ladenburg) or CH CH.CH. CH.CH. NMe. (Merling). Formed by distilling tropidine methylo-hydroxide with water (Roth, B. 17, 157; Merling, B. 24, 8118). Oil. HClAq forms 'hydrochloro-(α)methyl-tropidine' which gradually changes into the isomeric tropidine methylo-chloride. --B'₂H₂PtCl₈. [174°].—B'HAuCl₄: golden-yellow pp.

Methylo-iodide B'MeI. [162°]. Needles. (β)-Methyl-tropidine C₂H₁₃N. (205°). S.G. 15 922. Formed by heating (a)-methyl-tropidine at 150°-200°. Oil. Converted by HCl into

tropilene

TROPIGENIN C,H,NO. [161°]. oxidation of tropine by alkaline KMnO, (Merling, B. 15, 287). Got also by boiling 'homohydroapotropine' with baryta-water (Pesci, G. 12, 329). Needles, sol. water and alcohol. Reconverted by Oxidised by chromic acid MeI into tropine. mixture to ecgonic acid and a small quantity of tropic acid (Liebermann, B. 24, 615).—B'HI.— B'₂CO₂: crystalline pp. — B'₂H₂PtCl₈ aq. B'HAuCl₄: golden-yellow plates, sol. alcohol. TROPILENE C₇H₁₀O i.e. pp. — B'2H2PtCl6 aq.

CH₂ CH: CH CH. CHO (?) (Merling) (182°) (L.); (187°) (M.). S.G. 2 1.0091. V.D. 102.3 (calc. 110). Formed by distilling tropidine methylo-iodide with KOH (Lacenburg, B. 14. 2403; 15, 1028; A. 217, 138). Formed also from (β) -methyl-tropidine and cold HClAq (Merling, B. 24, 3123). Oil, smelling like oil of bitter almonds and acetone, sol. dilute HClAq, v. sol. alcohol and ether. Reduces warm ammoniacal AgNO_s, forming a mirror. Reduces Fehling's solution and KMnO, in the cold. Not attacked by AcCl. Combines with NaHSO, (M.). Nitric soid forms an adipic acid CaH₁₀O₄. Slowly combines with methylamine, forming (8)-methyltropine.

TROPILIDENE C,H, i.e.

CH < CH: CH / CH: CH2 / C: CH2 (114°). S.G. 2 ·9129. Formed by distilling methyl-tropine methyloiodide with KOH (Ladenburg, A. 217, 133; B. 14, 2403; 26, 1067). Formed also, together with methylamine, by distilling tropine with soda-lime. Oil, smelling somewhat like toluene, but combining readily with bromine, forming C,H,Br. Gives no pp. with ammoniacal Cu,Cl. Oxidised by CrO, to benzoic aldehyde and acid (Merling, B. 24, 3122)

TROPINE C.H., NO i.e. CH.CH2.CH.CH2.CH2.OH (Ladenburg) or CH.CH₂.NMe CH_2 CH CH CH_3 CH_4 CH_4 CH_5 CH_4 CH_5 CH_5

Methyl-oxyethyl-pyridine tetrahydride. [6∡°] Formed, together with tropic acid, by hydrolysis of atropine or hyoscyamine by baryta or HClAq (Kraut, A. 128, 281; 183, 87; Ladenburg, A. 206, 292; 217, 115; B. 13, 608; 20, 1653; 23, 1780; 26, 1067). Hygroscopic tables (from ether), v. e. sol. water and alcohol. Does not absorb CO, from the air. Its aqueous solution ppts. metallic oxides from their salts. Not attacked by nitrous acid. Not volatile with steam. According to Eykman (B. 26, 1400) the refractive index agrees best with Merling's formula.

Reactions.—1. Decomposed by distilling with soda-lime into tropilidene, methylamine, and H₂O.—2. Loses H₂O, being converted into tropidine by heating with fuming HCl and HOAc at 180° or by heating with H₂SO₄ (1 pt.) and water (1½ pts.) at 230°.—3. Conc. HIAq and red P at 150° form tropidine and tropidine periodide, but at 140° they form 'tropine iodide' C₂H₁,NI₂ which crystallises from water in prisms [115°], whence silver chloride followed by PtCl, give (C_sH₁₇NICl)₂PtCl₄ crystallising in red octahedra. By treatment with Ag₂O followed by HCl and PtCl₄ very soluble (C₈H₁,NCl₂)PtCl₄ is got.—4. Alkaline KMnO, forms tropigenin and, finally, oxalic acid and NH_a (Merling, A. 216, 341).-5. CrO, forms tropinic acid C₆H₁₁N(CO₂H)₂.—6. Aqueous HOCl yields prisms [111°], whence alcohol produces C₆H₅NCl₄.CH.OH [108°], which forms a hydrochloride [162°] crystallising in plates (Einhorn a. Fischer, B. 25, 1391).

Salts. - B'HCl. -B',H,PtCl. Orange-red monoclinic crystals. [200°] (Schmidt, A. 208, 214).—B'HAuCl₄. [212°].—B'HCl6HgCl₂. [246°] (Ladenburg, B. 24, 1631).—B'C₄H₂N₃O₇.

Nitroxyl derivative $C_8H_{14}(ONO_2)N$. Nitro-tropein. Formed by warming tropine (2 g.) with HNO₂ (12 g. of S.G. 1.25) at 100° (Ladenburg, B. 15, 1025). Alkaline liquid, sol. water, alcohol, and ether. Yields KNO₃ on boiling with KOH.—B'₂H₂PtCl₆. Needles.—B'HI. Prisms. Benzoyl derivative C₈H₁₄(OBz)N.

Benzoyl-tropeine. Formed by heating tropine hydrochloride with benzoic acid and dilute HCl (Ladenburg, B. 13, 1083; A. 217, 96). Orystallises as $C_{15}H_{19}NO_22aq$ [58°], $C_{15}H_{19}NO_2\frac{1}{2}aq$ [37°], or anhydrous [42°]. Sl. sol. water, v. sol. alcohol and ether.—B'HNO₈.—B'₂H₂PtCl₆ 2aq.—B'C₆H₂N₂O₇.

o-Oxy-bensoyl derivative Salicyl-tropeine. $C_8H_{14}(O.CO.C_6H_4.OH)N.$ Formed in like manner, using salicylic acid (Gaebe a. Caro, B. 13, 106; L.). Silky plates. Feeble poison, '025 g. killing a frog in a few hours. Has no apparent effect on the pupil.-B'HCl.—B'H2PtCl.—B'HAuCl4: yellow plates.

m-Oxy-bensoyl derivative C13H19NO3. [226°]. Small plates, v. sl. sol. water. Acts slightly on the pupil of the eye.—B'HCl.—B₁₂H₂SO₄4sq.—B'₂H₂PtCl₅: orange plates.

p. Oxy-bensoyl derivative C₁₁H₁₀NO₅

[227°]. Trimetric plates (containing 2sq).—
B'HNO₃.—B'₂H₂PtCl₂ 2sq. Orange plates.

Phenyl-acetyl derivative C₁₆H₂₁NO₂
i.e. C₆H₁₁N(0.CO.CH.Ph). Phenylacettroperne.

Aromatic oil.—B'HBr.—B'₂H₂PtCl₅.—B'HAuCl₄.

-B'2H2SO4. Soluble tables.

Mandelyl derivative C16H21NO4 i.e. C.H., N(O.CO.CHPh.OH). Homatropine. Formed by heating tropine with mandelic acid, HCl (1 pt.), and water (40 pts.), the yield being 50 p.c. of the theoretical amount (Ladenburg, C. R. 90, 921; A. 217, 82). Deliquescent prisms (from ether), al. sol. water. A solution of its hydrochloride is ppd. by potassio-mercuric iodide, but not by tannin. It enlarges the pupil of the eye almost as energetically as atropine, but the enlargement passes off much more rapidly (Völkers, A. 217, 86; Bertheau, Berl. Klin. Wochenschrift, 1880, No. 41; Tweedy a. Ringer, Lancet, 1880, No. 21).-B'HBr. Trimetric crystals; a:b:c = ·414:1: ·472.—B'C₆H₃N₃O₇. Yellow plates .-

B'HAuCl,.-B',H,SO,. Needles.

Cinnamoy l derivative C₁₇H₂₁NO₂. [70°]. Small plates (from dilute alcohol). Gives pps. with tannin, picric acid, potassio-mercuric iodide, and I in KI. Strong poison, 03 g. killing a frog in three minutes. Has little action on the pupil.

-B'HOl.-B'₂H_.PtCl₆.-B'HAuCl₄. Needles.

Atropyl derivative C₁₇H_{.21}NO₂. Atroptroperne. Oil.-B'HAuCl₄. Small needles.

Atrolactyl derivative C17H23NO20 Pseudo-atropine. [120°]. Formed from atro-lactic acid, tropine, and dilute HCl (1:400) by repeated evaporation (Ladenburg, A. 217, 87). Needles (from water). Resembles atropine in physiological action, two drops of a 1 p c. solution expanding the pupil and paralysing accommodation for a week.—B'HAuCl₄. [114°].

Phthalyl derivative C₂, H₂₂N₂O₄. [70°].

Formed, in very small quantity, by evaporating phthalic acid with tropine and dilute HCl.

Mass of silky needles.—B'H2PtCl

C₈H₁₈NOMeI. Methy . o - iodide crystals, nearly insol. alcohol. Yields B'MeOH and B'2Me2PtCl6 (Merling, B. 14, 1829; Ladenburg, A. 217, 131)

Ethylo-iodide B'EtI. Crystalline. Yields

B'2Et2PtCl6, a yellow crystalline powder.

(a)-Methyl-tropine C₈H₁₄MeNO. (243°). Formed by distilling tropine methylo-iodide. Liquid, v. sol. water and alcohol. Strongly alkaline. Its hydrochloride, distilled with solid KOH, yields dimethylamine.—B'HAuCl. unstable.

Methylo-iodide (C,H,MeNO)MeI. Deliquescent needles, decomposed by distilling with solid KOH into tropilidene and trimethylamine.

—(C₆H₁MeNO)₂Me₂PtCl₂. Orange crystals. (β)-Methyl-tropine C₆H₁₇NO. (198°-205°). Formed by shaking tropilene with an aqueous solution of dimethylamine (Ladenburg, B. 14, 2404). Liquid, split up by gaseous HCl into

tropilene and NMe.H.—B'HAuCl. Prisms.
(7)-Methyl-tropine C.H., NO. Formed in small quantity by distilling the methylo-hydroxide of (a)-methyl-tropine (Merling, B. 15, 288). V. sl. sol. Aq.—B'2H,PtCl_s. V. sl. sol. Aq. Metatropine C_sH₁₈NO. (238°). Formed by

shaking tropine iodide with water and Ag2O.

Not solid at -30°.-B'HCl: tables.

Hydrotropidine C₈H₁₅N. (168°). S.G. 2.937; 15 926. Formed by reducing tropine iodide C.H., NL with zinc-dust and HClAq (Ladenburg, B. 16, 1408). Liquid, m. sol. water.—B'HCl:

deliquescent crystals.—B',H,PtCl,. Ta Norhydrotropidine C,H,2N. [60°]. (161°). Formed by distilling hydrotropidine hydrochloride in a current of HCl (Ladenburg, B. [281°]. 20, 1649). Crystalline.—B'HCl.

20, 1649). Crystalline.—B'HCl. [281°].—
B'_2H_PtCl_*.—B'HHgCl_*.—B'C_*H_N_0_*. Needles.

Nitrosamine C_*H_1N_0. [117°]. Cubes.

Paratropine C_*H_1N_0. (202°). Formed from hydrotropine and K_FeCy, (Ladenburg, B. 24, 1626).—B'_2H_PtCl_*. [197°].—B'HAuCl_*. [182°].—B'HOl6HgCl_*. [225°]. V. sol. water.

\$\psi\$-Tropine C_*H_1N_0. [108°]. (242°). Formed, together with tropic acid, by warming hyoscine with heavets water (Ladenburg, B. 18, 1551: 17.

with baryta-water (Ladenburg, B. 13, 1551; 17,

151; Liebermann, 24, 2587). Formed also by dissolving its benzoyl derivative in hot conc. HClAq (Hesse, A. 271, 210). Hygroscopic prisms (from chloroform), v. e. sol. water. Yields tropinic and ecgonic acids on oxidation.—B',H,SO,. Hygroscopic crystals. — B'HAuCl. [198°] (Ladenburg); [202°] (H.); [225°] (Liebermann). — B'₂H₂PtCl₆ 4aq. [206°, anhydrous]. Trimetric prisms; a:b:c=.702:1:.879.

Methylo-iodide B'MeI. [270°]. bohedra. Yields B'MeCl and B'Me,PtCl.

[216°] crystallising from water.

Bensoyl derivative C15H19NO2. [48°]. Occurs in the leaves of Java coca (Liebermann, B. 24, 2336; Hesse, A. 271, 208). Colourless ether). — B'HCl. plates (from [270°]. -B',H,PtCl, B',H,PtCl,. Pale-yellow needles.—B'MeI.—B'MeCl.—B'MeAuCl,.—B'Me,PtCl, 2aq.

Mandelyl derivative $C_8H_{14}NO(CO.CH(OH).C_6H_5)$. ψ-Homatropine. Formed by heating ψ -tropine with mandelic anhydride at 200° (Liebermann a. Limpach, B. 25, 931).—B'HCl.—B',H,PtCl.—B',HAuCl.,

Tropyl derivative C,H,NO(C,H,O).

Isomeride of hyoscine. [88°]. $[a]_D = -4.9^{\circ}$ Nodules, insol. water, sol. alcohol. - B'HCl. [183°].—B'₂H₂PtCl₆.—B'HAuCl₁. [135°].

Atropyl derivative. [129°]. Formed from tropide and ψ -tropine at 200°. Crystals, Formed sol. water. Split up by HClAq into atropic acid and ψ -tropine.

Oxy-tropine carboxylic acid

CH.CH₂.CH.CH(OH).CH(OH).CO₂H Dioxyan-CH.CH₂.NMe

hydroecgonine. Formed by oxidising hydroecgonine with KMnO4 and aqueous Na2CO3 at 10 (Einhorn a. Rassoff, B. 25, 1395). Decomposes at 280°. V. e. sol. water and MeOH, insol. EtOH.—B'HCl. [251°].—B'C H,N,O,—KA'.

Methyl ether MeA'. [139°]. Tables, v.

e. sol. CHCl. Yields B'2H.PtOl. [210°].

Benzoyl derivative of the methyl

C,H,MeN.CH(OH).CH(OBz).CO,H. ether

[108°]. Needles.—B'HNO_s. [216°].—B'HCl. [203°].—B'HAuCl_s. [173°].—B'₂H₂PtCl_s. [208°].

Di-bensoyl derivative of the methyl ether C_sH₂MeN.CH(OBz).CH(OBz).CO₂H. Needles.—B'HNO_s. [190°].—B'HCl. [280°].—B'₂H₂PtCl_s. [205°].—B'HAuCl_s. [208°].

Reference.—OXYTROPINE.
TROPINE DIHYDRIDE C.H., NO i.e.

ÇH₂.CH₂.ÇH.CH₂.CH₂OH Hydrotropine. CH2.CH2.NMe Methyloxyethylpyridine hexahydride. Formed from oxy-ethyl-piperidine and KMeSO. (Ladenburg, B. 24, 1622). V. sol. water and

alcohol. — B'HCl5HgCl2. [214°]. — B'HAuCl4. Crystals

TROPINIC ACID C.H., NO. [258°] (L.); [220°] (M.). A product of oxidation of tropine and of econine by chromic acid mixture (Merling, A. 216, 348; Liebermann, B. 23, 2519). Needles, v. 30l. water, sl. sol. alcohol, insol. benzene. Gives off CO₂ when heated. Forms salts with acids and bases.—BaA'₂. Very hysalts with acids and bases.—BaA'₂. Very hygroscopic.—CaA'₂.—AgA'. Readily reduced.— (HA').PtCl..—HA'HAuCl.. HA'HCl aq. Golden prisms.-

TROPYLENE v. TROPILENE. TRUXENE (C,H,)x. Formed by heating truxone with red P and HIAq (S.G. 1.7) at 180°

(Liebermann a. Bergami, B. 22, 786; 23, 817). Formed also by heating hydrindone with conc. HClAq, and by heating phenyl-propionic acid with P₂O₅ (Hausmann; Kipping, C. J. 65, 269). Needles or plates, melting above 860°, insol. CHCl₃. Oxidised by boiling HOAc and CrO₃ to crystalline 'tri-benzoylene-benzene,' which is not melted at 360°. HNO, forms (4,2,1)-nitrophthalic acid.

(α)-TRUXILLIC ACID C18H16O4 i.e.

ÇHPh.ÇH.CO2H (?) (γ) -Isatropic acid. CO2H.CH-CHPh [274°]. Formed, together with (\$)-truxillic acid, by the action of boiling HClAq on isatropylcocaine, which is a by-product in the preparation of cocaine (Liebermann, B. 21, 2342; 22, 124, 783, 2242). Small needles (from alcohol), sol. hot HOAc. Yields cinnamic acid on distillation. Nitric acid (S.G. 1.52) forms two di-nitro-derivatives, [229°] and [290°]. Yields two isomeric sulphonic acids.—Na₂A" 10aq (Drory, B. 22, 2256).—BaA" 8 aq. Crystalline, v. sol. water. — CaA" aq. —Ag₂A". Flocculent pp., sol. NH₂Aq.

Methyl ether Me, A". [174°]. (c. 300°). Ethyl ether Et, A". [146°]. H.C.p H.C.p. 2,720,990 (Liebermann, B. 25, 92). C₁₈H₁₂(NO₂)₂Et₂O₄ [138°] (Homans, B. 24, 2590).

Isoamyl ether (C₅H₁₁)₂A". [83°] Prisms.

Ac₂O at 150° forms (α)-truxillic anhydride, while at 170° the product is (γ) -truxillic anhydride.

Di-amide $C_{18}H_{14}O_2(NH_2)_2$. [265°]. Needles, v. sl. sol. hot water, sl. sol. alcohol (Drory, B. 22, 2261).

 $Di-piperidide C_{16}H_{14}(CO.NC_5H_{10})_2$ [259°]. Got from the chloride and piperidine (Herstein, B. 22, 2264). Crystalline powder, sol. alcohol.

Mono-piperidide C₁₈H₁₄(CO₂H).CO.NC₅H₁₀. [250°]. Yields MeA' [151°] crystallising from ether in needles.

Anhydride $(C_{3a}H_{30}O_{7})_{2}$. [191°]. Mol. w. 1712 (1703 obs. by Raoult's method). Formed by heating the acid with Ac₂O and NaOAc at 100°. Needles (from benzene-ligroïn). Gives no fluorescein. When heated for an hour at 200° it changes to (γ) -truxillic anhydride. (β) -Truxillic acid $C_{18}H_{16}O_4$ i.e.

CHPh.CH.CO₂H CHPh.CH.C'O₂H (?)

(8)-Isatropic acid. [206°]. Formed at the same time as the (a)-acid (Liebermann, B. 21, 2342; 22, 783, 2243; 25, 90; 26, 837). More sol. water than the (a)-isomeride. Yields benzil on oxidation. Nitric acid (S.G. 1.52) forms a di-nitro- derivative [216°] (Homans, B. 24, 2590). Forms cinnamic acid on distillation.—Na2A" 2aq.

—BaA" 2aq. Sl. sol. water.—CaA"3 aq.—Ag,A".

Methylether Me,A". [76°]. Mol. w. 309
(by Raoult's method); calc. 324. H.C.p. 2.422,900. Monoclinic prisms; a:b:c=

 $\cdot 826:1:2\cdot 019$; $\beta = 89^{\circ} 22^{\circ}$.

Ethylether Et,A". $[47^{\circ}].$

Chloride C₁₆H₁₄O₂Cl₂. [96°]. Tabular prisms, v. sol. ether (Drory, B. 22, 2260).

Monopiperidide C₁₆H₁₄(CO₂H).CONC₅H₁₆. Tabular

[224°]. Formed from (B)-truxillic anhydride and piperidine (Herstein, B. 22, 2264). Needles, sl. sol. cold alcohol.

Dipiperidine ClaH14(CO.NC,H10)2. [180°]. Formed from the chloride and piperidine. Prisms, v. sol. alcohol.

Phenylimide C16H14 CO NPh. [180°].

Mol. w. 351 (obs.). Got by heating the anhydride with aniline. Colourless needles (from alcohol). Cold alcoholic potash forms the compound CO₂H.C₁₆H₁₄.CO.NHPh [197°].

Phenyl hydraside C16H14CO.NH

[218°]. Formed from the acid, phenyl-hydrazine, and HOAc on warming. Crystals (from HOAc).

 $C_{16}H_{14} < CO \xrightarrow{C(C_6H_2(OH)_2)_2} O.$ Fluorescein Formed by heating the acid or its anhydride with resorcin at 240°. Amorphous brownish-red powder, sol. alkalis forming fluorescent solutions. v. sol. alcohol, insol. benzene.

Anhydride C₁₈H₁₄O₃. [116°]. Me 278 calc., 289 obs. by Raoult's method. Mol. w. not change to an isomeride when heated.

(γ)-Truxillic acid C₁₈H₁₈O₄. ε-Isatropic acid. [228°]. Formed by heating the anhydride of the (a)-isomeride with HCl at 160°. Needles (from dilute alcohol), v. sl. sol. hot water, v. sol. ether. Yields cinnamic acid on distillation. Gives the same anhydride as the (a)-isomeride (Ladenburg, B. 22, 124). Heated with HClAq all 260° it is changed to (a)-truxillic acid. HNO₃ (S.G. 1.52) forms a di-nitro- derivative [293°].

Salts.—BaA" 11aq.—CaA" 3½aq.-CaA" 6½aq.—Ag₂A": crystalline pp.

Methyl ether Me₂A". [126°].

Needles. Mol. w. 332 (calc. 324).

Mono-ethyl ether C16H14(CO2H).CO2Et. [172°]. Formed, together with the di-ethyl ether, by saturating an alcoholic solution of the acid with HCl (Liebermann, B. 22, 2240). Needles, yielding AgA'. At 320° it is converted into a mixture of (a)-truxillic acid and (a)-truxillic

Di-ethylether Et2A". [98°]. Needles, v. sol. alcohol (Drory, B. 22, 2260).

Mono-piperidide

C₁₀H₁₄(CO.NC₅H₁₀).CO₂H. [261°]. Plates (from water or dilute alcohol). Yields MeA' [201°] and a piperidine salt (C₅H₁₁N)HA' 3aq [218°], which crystallises (with Saq) from alcohol (Herstein, B. 22, 2262).

Di-piperidide C16H14(CO.NC5H16). [248°]. Needles, insol. water, sol. alcohol.

C₁₆H₁₄(CO.NHPh).CO₂H. Mono-anilide [220°]. Formed by heating the acid with aniline (Liebermann, B. 26, 838). Needles (from dilute alcohol).

Aniline C₁₆H₁₄(CO.NHPh)₂. [255°].

Anhydride C18H14O3. Mol. w. 282 (calc. 278). Gives no fluoresceïn.

(δ)-Truxillic acid $C_{18}H_{18}O_4$. [174°]. Formed by fusing (β)-truxillic acid with potash. Needles (from water), v. sol. alcohol (Liebermann, B. 22, 2250; Hesse, A. 271, 205). Yields a di-nitroderivative [226°]. Yields cinnamic acid on distillation.—CaA": rosettes of needles.—BaA" 4aq: prisms, sl. sol. Aq.—CuA" 2aq.—Ag₂A".

Methyl ether Me₂A". [77°]. Ne

Needles. Reference.—OXY-TRUXILLIC ACID.

TRUXONE $(C_0H_sO)_x, x = 2 \text{ or } 3. \lceil 289^\circ \rceil$. Formed from (a)-truxillic acid and fuming H.SO. (S.G. 1.96) at 15° (Liebermann a. Bergami, B. 22, 784; 23, 320). Plates (by sublimation), insol. water, acids, and bases. Not attacked by HNO₃. Yields (C.H.Cl.), [178°]. Aniline and HOAc on boiling form the anilide (C, H,: NPh), crystallising in needles, [270°], decomposed by fusion.

Phenyl-hydrazide (C₂H₆:N₂HPh)_x. 270°]. Needles, v. sl. sol. alcohol.

Oxim (C₂H₆:NOH)_x. Very unstable, easily changing to an anhydride. Boiling Ac.O yields (B.H.: NOAc), [261°], m. sol. HOAc.

TULUCUNIN C20H14O4. Occurs in the bark of Tarapa Tulucuna (Caventou, J. Ph. [3] 35, 189). Light-yellow amorphous resin, with very bitter taste, sl. sol. water, v. sol. alcohol, insol. ether. Turned blue by cold H₂SO₄.

TUNGSTATES, and derivatives of; v. Tung-STEN OXYACIDS, SALTS, AND DERIVATIVES OF, p. 802; also Tungsten thio-acids, and salts of, p. 810.

TUNGSTEN. W. (Wolfram.) At. w. 183-6. Mol. w. is unknown. S.G. 18-77 (Waddell, Am. 8, 280); 18-77 (Moissan, C. R. 116, 1225); 19-13 at 4° (Roscoe, C. J. [2] 10, 286); for other determinations v. Bernoulli (P. 111, 576), Zettnow (P. 111, 16). S.H. (6° to 15°) ·035 (De la Rive a. Marcet, A. Ch. [2] 75, 113). S.V.S. c. 9.7.

Historical.—In 1785 a new acid was prepared by J. and F. d'Elhujar from the mineral wolframite, and shown to be identical with an acid obtained three years before from tungstein by Scheele (Opusc. 2, 119). The metal of the new acid was isolated by J. and F. d'E. For many years the new metal was known as wolfram or tungsten (from Swedish = heavy stone); the latter name has gradually driven out the former in England and France, but the metal is generally called wolfram in Germany; the symbol W is universally employed.

Occurrence.-Never uncombined. Wolframite (tungstate of Fe and Mn) is found in considerable quantities in Cornwall, Saxony, Bohemia, &c.; tungstenite, or scheelite (tungstate of Ca), scheeletine (tungstate of Pb), and some other tungstates occur in various localities; wolframine or wolfram-ochre, WOs, is also found. Some tin ores contain compounds of W, and W is therefore not infrequently found in specimens

of tin.

Formation.—1. By reducing WO, by heating to redness in H (Berzelius, P. 4, 147; Wöhler, A. 77, 262; Zettnow, P. 111, 16; Roscoe, C. N. 25, 61, 78).-2. By heating WO, with Na and NaCl (Z., l.c.).—3. By reducing WO, by mixing with 10 p.c. charcoal and 2 p.c. resin, and heating in a closed crucible for some hours to a white heat (Filsinger, S. C. I. 1878. 229).—4. By passing vapour of oxychloride of W and H through a red-hot tube (von Uslar, A. 94, 255).

5. By heating WCl, in H, in a zinc bath (Roscoe, l.c.).—6. By strongly heating the salt (NH₄)₂WO₄ in a carbon crucible (Bucholz, P. 111, 576).—7. By strongly heating the nitride (q. v., p. 800) in H (Wöhler, A. 73, 190).

Preparation .- 1. Pure WO, is heated to bright redness, in a Pt tube, in a stream of pure dry H (v. Formation, No. 1).—2. A mixture of pure WO, and dry lampblack is heated in an electric

furnace (Moissan, C. R. 116, 1225).

WO, is prepared from wolframite by heating the very finely powdered mineral with conc. HClAq, pouring off the solution from time to time and adding more HClAq, and after some time adding a little HNO, Aq until most of the brown solid is changed to yellow WO₃.xH₂O, washing the residue thoroughly by decantation, adding a considerable quantity of NH, Aq and warming, filtering from quartz, unchanged wolframite, &c., evaporating the solution until small lustrous crystals of an acid ammonium tungstate separate, boiling these crystals for a long time with HNO₃Aq, washing, and heating the residue (Scheibler, J. pr. 83, 239; cf. Bernoulli, P. 111, 590).

Properties.—A steel-grey, hard, brittle, crystalline powder (Riche, A. Ch. [3] 50, 5). By reducing the nitride in H, Wöhler (A. 73, 190) obtained W as a black powder. By reducing WO, by C in an electric furnace, Moissan (C. R. 116, 1228) obtained a lustrous, very hard, greyish white solid. As produced by passing the vapour of WO2Cl, mixed with H through a red-hot tube, W forms a shining, dark steel-grey, mirror-like deposit, which can be detached from the tube in hard, brittle crusts (von Uslar, A. 94, 255). W can be melted in the O-H flame, part of it burning to WO₃ (Riche, l.c.); or by using a current from 600 Bunsen cells, in an atmosphere of N (Desprez, C. R. 29, 549). Considerable masses may be melted by employing the current from a dynamo (v. Huntington, C. N. 46, 163). W was regarded by Faraday as diamagnetic (T. 1846. 49). The emission spectrum is described by Thalen (A. Ch. [4] 18, 202). Wis unchanged in air, but when the pulverulent metal is heated it burns to WO3; compact W burns in air only at very high temperatures; as obtained by reducing WCl, by H, W is said to be pyrophoric. Water is decomposed by W at a red heat. The metal is scarcely acted on by HClAq or H₂SO₄Aq; it is oxidised by HNO, Aq to WOs. xH2O; the pulverulent metal dissolves slowly in caustic alkali solutions. Finely divided W reduces and ppts. many metals from solutions of their salts. combines directly with Br, Cl, I, or S. W is not acted on by heating in NH, nor in CO (v. Rideal, C. J. 55, 45; Smith a. Oberholtzer, Zeit. f.

anorg. Chemie, 5, 63).

The at. w. of W has been determined (1) by reducing WO₃ in H, and again oxidising W to WO₃ (Berzelius, P. 4, 151 [1825]; Schneider, J. pr. 50, 158, 161 [1850]; Marchand, A. 77, 263 [1850]; von Borch, J. pr. 54, 254 [1851]; Riche, [1850]; Von Bordi, J. M. 34, 294 [1801]; Metal. A. Ch. [3] 55, 143 [1859]; Bernoulli, P. 111, 597 [1860]; Persoz, A. Ch. [4] 1, 93 [1864]; Roscoe, C. N. 25, 61, 73 [1872]; Waddell, Am. 8, 280 [1887]); (2) by determining H2O in BaW4O13.9H2O (Scheibler, J. pr. 83, 324 [1861]); (3) by reducing WO₃ in H and determining H₂O produced (Bernoulli, P. 111, 597 [1860]); (4) by analysing Ag. WO., and FeWO. (Zettnow, P. 130, 16, 240 [1867]); (5) by analysing WCl, (Roscoe, C. N. 25, 61, 73 [1872]);
(6) by determining S.H. of W (De la Rive a. Marcet, A. Ch. [2] 75, 113 [1840]); (7) by determining V.D. of WCl., WCl. and WOCl. (v. these compounds). The older determinations gave values for at. w. varying from 186 to 189; the most recent determination (by Waddell) gave 184.04; Rescoe's determinations, made by reducing WO, to W and oxidising W again to WO, by finding the ratio of WCl, to AgCl and to Ag. and by reducing WCl, in H, gave values varying from 183.25 to 183.77. The number 183.6 is probably correct to half a unit. The atom of W is pentavalent in the gaseous molecule WCls and hexavalent in the gaseous molecule WCl.

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W is the fourth member (the third is as yet anknown) of the even series, or chromium, family of Group VI. in the periodic arrangement of the elements. W is followed in its family by U, and it succeeds Mo and Cr. W very closely resembles Mo; like that element W is both metallic and non-metallic; the balance of these properties being, however, not quite so evenly maintained in W as in Mo. It is doubtful whether any definite salt has been isolated derived from an oxyacid by replacing H by W. The oxide WO, acts as a fairly acidic oxide; like MoO, it is remarkable for the large number of compounds it forms, simultaneously, with oxides that are more basic and oxides that are less basic than itself. The chemical relations of Ware discussed in the article Chromium group of elements, vol. ii. p.

Reactions.-1. Heated in air burns to WO3; finely divided W burns easily; compact W only at a very high temperature (v. Roscoe, C. N. 25, 61, 73).—2. Heated to redness in steam gives H and oxide (? WO_s).—3. Reacts with hot nitric acid, or aqua regia, to form WO3.xH2O; the same product is obtained by heating W with conc. sulphuric or hydrochloric acid, but oxides intermediate between WO2 and WO3 are formed at first (Riche, A. Ch. [3] 50, 15).—4. Pulverulent W dissolves in boiling cone. potash solution, giving H and a tungstate of K (Riche, l.c.). 5. Heated in carbonyl chloride to 150°-200° WOCl, is produced, mixed with C (Smith a. Oberholtzer, Zeit. f. anorg. Chemie, 5, 63).-6. By heating to redness with sulphur chloride a red, crystalline thiochloride, probably W2S,Cls, is formed as a sublimate, unstable in air (S. a. O., l.c.).-7. Finely divided W reduces solutions of several metallic salts; AgNO, and AuCl, solutions are reduced to Ag and Au, HgCl₂ to HgCl, Cu, Pd, Pt and Rh are partially ppd. from solutions; Bi, Cd, and Pb are not ppd. (Smith, Zeit. f. anorg. Chemie, 1, 360).

Combinations.—1. Heated in oxygen, WO, is formed.—2. Combines with chlorine, to form WCl_s, at c. 300°; with bromine, to form WBr_s, at a higher temperature; and with *iodine*, to form WI₂, when strongly heated.—8. Heated with sulphur, forms WS₂—4. The compound W₂P₄ is said to be formed by strongly heating W in phosphorus vapour (Wöhler a. Wright, A.

79, 244).

Detection and Estimation.—Compounds of W form colourless beads with borax or microcosmic salt in the outer blowpipe flame. The borax bead is yellow in the inner flame if a considerable quantity of a compound of W is present. The microcosmic salt bead is blue in the inner flame in the absence of compounds that themselves form coloured beads; in presence of iron compounds the bead is blood-red, but addition of tin causes it to become blue. Soluble alkali tungstates are formed by fusion with alkali carbonate or nitrate; addition of excess of H₂SO₄Aq, HClAq, H₂PO₄Aq, H₂U₂O₄Aq, or HC2H2O2Aq to an aqueous solution of a tungstate, followed by immersion of a piece of zinc in the liquid, produces a blue colour; addition of H2S to an acidulated solution of a tungstate gives no pp., but a blue colour. According to Mallet (C. J. 28, 1228), the successive addition of small pieces of zinc to the solution obtained

by adding excess of cone. HClAq to an alkaline tungstate produces various colours, the most marked of which is a brilliant magenta-red. By adding KCNSAq and then small pieces of zinc to a solution of an alkaline tungstate in excess of conc. HClAq, a deep green colour is produced; and an amethyst colour is noticed when KCNSAq is added to the solution of an alkaline tungstate, the solution is then diluted, HClAq is then added, and lastly zinc is placed in the liquid. Very minute quantities of W can be detected by adding SO₂Aq and Zn to solution of an alkaline tungstate, whereby a light-blue colour is obtained (M., l.c., p. 1233). W is estimated as WOs, obtained by evaporation and strongly heating; for separation from other elements a Manual of Analysis must be con-

Tungsten, alloys of. By reducing mixtures of WO₈ with oxides of Sb, Bi, Co, Cu, Pb, Ni, or Zn, Bernoulli (P. 111, 573) obtained alloys of W with these metals, provided there was not more than 10 p.c. of the foreign metal present. An alloy with aluminium, approximately WAl,, was obtained by Wöhler a. Michel (A. 115, 102) by heating a mixture of WO₃, Al, cryolite, and NaCl and KCl. W alloys with iron; v. Poleck a. Grützer (B.26,35) for an alloy approximately W₂Fe. An alloy of 9 or 10 p.c. W with steel is extremely hard (v. Bernoulli, l.c.; Philipp, Hof-mann's Ber. über chem. Industrien, 745; Le Guen, C. R. 56, 593; 59, 786; 63, 967; 64, 619; 68, 592; Caron, A. Ch. [3] 68, 143; Osmond, C. R. 104, 985; Gruner, C. R. 96, 197).

Tungsten, amidonitrides of, v. Tungsten nitrides, and allied compounds, p. 799.

Tungsten, bromides of. The elements combine when heated together to form WBr, and by partial reduction in H this gives WBr₂.

WBr. TUNGSTEN PENTABROMIDE formula is probably molecular, from the analogy of WCls. Prepared by heating W in excess of Br vapour, taking care that every trace of water and O are excluded. Dark, violet-brown needles, resembling I; melts at 276° and boils at 333°, giving off a dark-brown vapour; decomposed on distillation, with separation of Br; moist air or water forms HBrAq and blue oxide of W: heated to 350° in a stream of H, WBr, is formed (Roscoe, C. N. 25, 73; cf. Borck, J. pr. 54, 254; Blomstrand, J. pr. 82, 408).

Tungsten dibromide WBr₂. Obtained, as a bluish-black, velvety solid, when WBr, is heated at c. 350° (in a bath of ZnCl₂) in a stream of dry H; WBr_s, WOBr_s, and Br distil over, and WBr₂ remains. Heated above c. 400° in H gives W and HBr; with HNO, Aq gives WO, HBrAq,

and NH₄NO₂Aq (Roscoe, l.c.).

Tungsten, chlorides of. When W is heated in Cl the compound WCl, is formed, and by reduction in H or CO, this gives WCl, WCl, and WCl. Great confusion existed about the compositions of the chlorides of W until Roscoe's researches in 1872 established the formulæ.

TUNGSTEN HEXACHLORIDE WCl. Mol. 895.82. Prepared by heating W in Cl (Roscoe, C. N. 25, 61). The materials must be perfectly dry, and every trace of air must be excluded, else WOCl, is formed; the WCl, sublimes on the cooler part of the tube, and is then distilled several times in a stream of dry Cl, and then in

a current of dry H (a small quantity of WCl, is formed, and distils off with the H). Teclu (A. 187, 255) obtained WCl, by heating WO, and PCl, in a sealed tube at 170°. WCl, forms darkviolet crystals; crystallises from CS, in brown, six-sided tablets, from POCl, in steel-blue, metallike, regular crystals (T., l.o.). Melts, out of contact with air, at 275°, and boils at 346.7° at 760 mm. pressure (R., l.c.). V.D. 190 at 350° (Debray, C. R. 60, 820; Roscoe, l.c.); V.D. 168-8 at 440° (D., l.c.; R., l.c.; Rieht, B. 3, 666). Dissolves readily in CS₂, or POCl₃ (T., l.c.). Distilled in CO₂ gives WCl₅ and Cl; this fact taken in connection with V.D. at 440° shows that at somewhat above b.p. WCl_s probably dissociates to WCl_s and Cl. WCl_s is unchanged in air; but if a trace of WOCl, is present, HClAq and WO are formed. Hot water produces WO₃ and HClAq. Heated in air, or in O, forms WOCl4 (Roscoe, l.c.; Blomstrand, J. pr. 82, 417). WOCl, is also formed by heating WO, with WCl, WCl, interacts with NH, at the ordinary temperature, to form NH₄Cl and W₂N₂ (Rideal, C. J. 55, 44).

TUNGSTEN PENTACHLORIDE WCl_s. Mol. w. 360·45. Prepared by repeatedly heating WCl_s somewhat above its b.p. (346·7°) in a stream of dry H, and when a solid residue has been formed removing the WCl_s from less volatile lower chlorides by heating in CO₂ (Roscoe, C. N. 25, 61). Black, lustrous, needle-shaped crystals; very deliquescent; melts at 248°, and boils at 275·6° (R., l.c.). V.D. 175·6 to 179·9 at 350°; 185·7 to 186·4 at 440° (R., l.c.). Slightly soluble in CS₂ forming a blue liquid; forms an olivegreen solution in water, but is mostly decomposed to HClAq and blue oxide of W. Heated in O forms WOCl₄ and Cl (R., l.c.; cf. Blomstrand, J. pr. 82, 425; 89, 230).

TUNGSTEN TETRACHLORIDE WOl. This compound is present in the solid residue obtained by heating WOl. in H in the preparation of WOl. (v. supra); it is prepared by distilling this residue, in a bath of H.SO. in a stream of dry CO. returning the distillate to the distilling vessel and heating again, and repeating these processes several times (Boscoe, l.c.). A soft, crystalline, greyish-brown powder; very hygroscopic; has not been fused or volatilised; heated strongly, it gives WOl. and WCl. Heated in H to c. 440° pyrophoric W is produced. Decomposed by water to WO. and a greenish-brown solution (R., l.c.).

Tungsten dichloride WCl. Prepared by

TUNGSTEN DICHLORIDE WCl.. Prepared by heating WCl., in a bath of zinc, in a stream of dry CO.; WCl. distils off and WCl. remains. A loose, grey, amorphous powder; water forms WO. and HClAq, with evolution of H; interacts with H which has been passed through HNO. Aq to form WO., HCl. and NH. NO. (R., l.c.).

Tungsten, cyanides of. No cyanides of W

Tungsten, cyanides of. No cyanides of W have been isolated; according to Wyrouboff (A. Ch. [5] 8, 444; of. Atterberg, Bl. [2] 24, 855), compounds containing W, K, and FeCy, are obtained by adding HOLAq to a mixture of K tungstates and K ferrocyanide.

Tungsten, fluorides of. No fluoride of W has been isolated. According to Berzelius (P. 4, 147), when a solution of WO.H.O in HFAq is evaporated and the residue is treated with water, a solid remains which is free from F after being

heated in NH₂. According to Riche (A. Ch. [8] 50, 41), a solution of tungstic hydroxide in HFAq gives crystals of WO₂.H₂O on evaporation.

Tungstoxyfluorides. (Fluotungstates.) By dissolving tungstates of the form M₂WO₄ in HFAq, Marignac obtained a series of compounds of the forms 2M¹F.WO₂F₂.xH₂O and also a few compounds of the form M¹F.WO₂F₃.H₄O (A. Ch. [3] 69, 67). These compounds may be regarded as salts of the hypothetical acids H₂WO₂F₄ and HWO₂F₃; they are similar to some classes of the compounds described as fluomolybdates (this vol. p. 425).

The tungstoxyfluorides etch glass even when dry; they react slowly with acids, giving WO₃.H₂O. The salts of the forms $M^1_2WO_2F_4$ and $M^{11}WO_2F_4$ were prepared by dissolving $M^1_2WO_4$ and $M^{11}WO_4$ in HFAq and evaporating; in some cases also by dissolving WO₃.H₂O in HFAq, adding MOH, and evaporating. The salts of the form $M^1WO_2F_3$ were formed by dissolving acid tungstates in HFAq and evaporating. The following compounds were isolated: (1) $M^1_2WO_2F_4.xH_2O$; $M=NH_4$, x=0; $M=NH_4$, x=0; $M=NH_4$, x=1; $M=NH_4$, x=1; $M=NH_4$, x=1; $M=NH_4$, x=1; $M=NH_4$, x=1. The salts $(NH_4)_4WO_2F_6.(NH_4)_2WO_4$ and $CuWO_2F_4.NH_3$ were also obtained.

were also obtained.

By dissolving K₂WO₂F₄.H₂O in 4 p.c. H₂O₂Aq, and crystallising from dilute H₂O₂Aq containing a little HF, Piccini (Zeit. f. anorg. Chemie, 2, 21) obtained K₂WO₃F₄.H₂O. P. calls this compound fluoroxypertungstate; it might also be named pertungstoxyfluoride.

Tungsten, haloid compounds of. When W is heated in a stream of Cl the compound WCl₄ is formed, and this by reduction in H gives WCl₅, WCl₄, and WCl₂. The compound WBr₅ is formed by heating W in Br vapour, and WBr₅ is obtained by partially reducing WCl₅ in H. Small quantities of WI₂ are obtained by heating W in vapour of I. No fluoride of W has been isolated. Oxychlorides and oxybromides of the forms WOX₄ and WO₂X₅ are obtained by heating W oxides in Cl or Br, and in other ways. The following compounds have been vaporised, and the simplest formulæ are molecular: WCl₅ WCl₅, WOCl₄. The formulæ WBr₅, WOBr₆ WCl₅, WOCl₄, and WO₂Br₅ are probably molecular; but if the analogy of the chlorides and bromides of Mo of the form Mo₂X₂z is to be followed (v. vol. iii. pp. 427, 428) it is probable that the molecular formulæ of the dichloride and dibromide of W are not less than W₂X₅.

Tungsten, hydroxides of, v. Tungsten oxides and hydrated oxides (p. 800), also Tungsten oxyacids (p. 802).

Tungsten, iodide of, WI. This, the only iodide of W that has been isolated, is obtained, in very small quantities, by passing I vapour, mixed with CO₂, over red-hot W. It forms a metal-like, greenish crust; heated in air it gives off I and leaves WO₃; it is not decomposed by water (Roscoe, C. N. 25, 73).

Tungsten, nitrides of; and allied compounds. Compounds of W with N, and probably also with N and H, are formed by heating WCl. or WOCl. in NH, the interaction of NH, and WO₂ probably produces a compound, or compounds, of W, N, H, and O; and a compound of W, N, and O is perhaps formed by heating WO₂ with NH,Cl.

Tungsten nitrides. By passing dry NH2 over WCl_s, and washing away the NH₄Cl produced, by water, Rideal (C. J. 55, 44) obtained a black lustrous powder, agreeing fairly with the composition W₂N₂. This substance is insoluble in HNO₂Aq, dilute H₂SO₄Aq, or NaOHAq; hot conc. H₂SO₄ produces NH₃ and WO₃; fusion with NaOH forms Na tungstate; heating in air,

or with aqua regia, oxidises it to WO₃ (R., l.c.).

By continued heating WCl₅ in NH₅, to a temperature difficult to regulate, as slightly too high a temperature produces W, Uhrlaub obtained a black substance to which he gave the formula W₂N₂ (Die Verbindungen einiger Metalle mit Stickstoff, Göttingen, 1859).

Tungsten amidonitrides. By the action of NH_s on WCl_s, Wöhler (A. 73, 198) obtained a black substance containing from 86.76 to 90.8 p.c. W, and 824 p.c. N; W. supposed this substance to be a mixture of two amidonitrides, 2WN₂,W(NH₂)₂ and 2WN.W(NH₂)₂. W. obtained similar black substances, which he did not analyse, by heating chlorides of W with NH,Cl

(A. 105, 258).

Compounds of tungsten with N, H, and O. Wöhler (A. 73, 198) obtained a black substance by heating WO₃ to dull redness in dry NH₃; the percentage of W in this substance varied from 87.65 to 88.47, and it contained 7.15 p.c. N, and c. 2 p.c. H. To this substance W. gave the formula $W_8N_{10}H_1O_4$ (=4WN₂.W₂(NH₂)₂.2WO₂). Rideal (C. J. 55, 44) passed dry NH₃ over WO₃ heated to a dull redness until the yellow WO₃ was changed to a black, amorphous powder; he allowed to cool, and removed excess of NH, by passing a current of dry air through the tube. The black substance thus obtained contained 85.26 p.c. W, and 7.4 p.c. N; assuming c. 3 p.c. H, the numbers agreed fairly with the formula W,N,H,O,.

Compound of tungsten with N and O. Rideal (l.c.) obtained a black powder, agreeing in composition approximately with the formula WN₂.WO₃, by heating WO₃ with NH₄Cl until the weight became constant. According to Rideal (l.c.) finely divided W does not change when heated in a stream of dry NH_s, after having been heated to redness in dry H; nor does NH, react with the blue oxide of W when

heated therewith.

Tungsten, oxides and hydrated oxides of. When powdered W is heated in O it combines to form WOs; by heating this oxide in H a blue oxide is formed, to which various formulæ intermediate between WO, and WO, have been assigned; by further heating in H, or with C, the dioxide WO2 is produced. There are indications of an oxide with more O than WO. WO2 dissolves in acids, but corresponding salts have not been isolated; the blue oxide separates from these solutions; WO2 is also soluble in alkali WO, is insoluble in acids; it dissolves in alkali and alkaline carbonate solutions, forming tungstates (v. under Tungsten oxyacids, p. 802). WO, combines with many acidic oxides to form compounds which react as acids (v. p. 801). WO, is the anhydride of more than

one tungstic acid (v. p. 801). None of the oxides has been vaporised; the mol. w. of none is

TUNGSTEN DIOXIDE WO. (Brown oxide of

tungsten.)

Formation.—1. By heating a mixture of WO₃ and C to redness (Buchholz, S. 3, 1).—2. By heating WO₃ to incipient redness in H (Wöhler, A. 78, 198; 77, 262).—8. By decomposing WOl₄, or WOl₂, by H₂O (Roscoe, C. N. 25, 61, 78).—4. By the reaction of Zn and HClAq on WO, (Wöhler, P. 2, 345), or on solution of a metatungstate (Riche, l.c.).

Preparation.-WO, is placed in a porcelain tube closed at one end and provided with a long opening in the middle; this tube is placed inside another porcelain tube which is kept at red heat, while H is passed through as long as water continues to be formed (Wöhler, A. 77, 262 note); the product is allowed to cool in H, and is kept in H for 24 hours (Riche, A. Ch. [3] 50, 29).

Properties.—A brown powder with a slight violet sheen; prepared by reducing WO, by Zn and HClAq, the oxide is obtained in metal-like lustrous crystals, pseudomorphs of WO₃. S.G. 12·11 (Karsten, S. 65, 394).

Reactions.—1. Moist WO2 oxidises rapidly in air to WO,; as prepared by reducing WO, in H it is pyrophoric, but if allowed to cool slowly in H the product is not pyrophoric (Berzelius, 1. 4. 147; 8, 267; Wöhler, l.c.; Riche, l.c.).—
2. Heated strongly in hydrogen gives W.—
3. Chlorine forms WO₂Cl₂ when heated with WO₂ (Roscoe, l.c., p. 63).—4. Heated to dull redness in a stream of ammonia, a compound, or compounds, of W with N, H, and O is formed (v. Compounds of W with N, H, and O, under Tungsten nitrides, supra). — 5. By heating with sal ammoniac, Rideal (C. J. 55, 44) obtained a black powder approximately WN₂.WO₃ (v. Compound of W with N and O, supra).—6. Moist WO₂ dissolves easily in warm hydrochloric or sulphuric acid, forming reddish-brown solutions from which blue oxide of W separates. WO₂ prepared in the dry way is scarcely acted on by acids, except aqua regia, which oxidises it to WO₃.—7. WO₂ dissolves in conc. potash solution, giving off H, and forming K tungstate. 8. Reduces mercuric chloride to HgCl, and ppts. Cu₂O from solutions of copper salts (Riche, A. Ch. [3] 50, 5).—9. Heated in nitric oxide to below 500° gives the blue oxide; oxidised to WOs by heating in nitrogen dioxide to c. 300° (Sabatier a. Senderens, C. R. 114, 1429; 115, 236).

BLUE OXIDE OF TUNGSTEN. Blue compounds of W and O are obtained by heating WO, with reducing agents; analyses lead to formulæ such as W_2O_5 , W_3O_9 , or W_4O_{11} , intermediate between WO_2 and WO_3 . Malaguti (A. Ch. [2] 60, 273) gave the formula W_2O_3 to the blue product of heating WO_3 in H to c. 250°; von Uslar (Beiträge zur Kenntniss des Wund Mo [Göttingen, 1855]) gave the formula W₂O₂. Gmelin said that a blue oxide of the composition W₄O₁₁ is formed by strongly heating Wo, in CO.

Blue compounds are also obtained by strongly heating NH, tungstates out of contact with air (Malaguti, I.c.; von Uslar, I.c.). Aqueous solutions of WO, or tungstates are coloured blue by the action of very weak reducers, e.g. by exposing moist WO, on paper to sunlight (Liesegang, C. C. 1865. 943), or by the action of $SnCl_2Aq$ on tungstates (Bunsen, A. 188, 289; v. also Scheibler, J. pr. 83, 313). Conc. H_2SO_4 or HClAq gives a blue colour when heated with W (Riche, A. Ch. [3] 50, 15). Tungstates boiled with acetic acid and then electrolysed give blue coloured substances (Smith, B. 13, 753).

PEROXIDES OF TUNGSTEN. Fairley (C. J. 81, 141) obtained indications of an oxide with more O than WO₃, by dissolving WO₃.2H_O in H₂O₂Aq, and evaporating over H₂SO₄ in vacuo; the green, transparent, yellow scales thus obtained dissolved in water, forming a solution which could be titrated with KMnO₄Aq, with separation of WO₃.xH₂O. Cammerer (Chem. Zeitung, 15, 957) found that WO₃ dissolved in boiling H₂O₂Aq, with evolution of O; on spontaneous evaporation he obtained a yellow p-wder, to which he gave the formula WO₃.H₂O₃.H₂O (cf. reaction of H₂O₂Aq with CrO₃Aq, vol. ii. p. 166). According to Péchard (C. R. 112, 1060), a salt Na₂O.W₂O₂.2H₂O is formed by boiling Na paratungstate solution with H₂O₂Aq and evaporating in vacuo.

TUNGSTEN TRIOXIDE WOs. (Tungstic anhydride. Sometimes erroneously called tungstic

acid.)

Occurrence.—As tungstic ochre or wolframine in Cumberland, Connecticut, North Carolina, &c. Formation.—1. By strongly heating in air tungstate of NH₄ or of Hg¹.—2. By strongly heating WO₃.xH₂O.—3. By burning W in air or O

Preparation.—An aqueous solution of commercial Na, WO, is poured into boiling HClAq (1:1); the ppd. WO, is washed with HClAq until NaCl is completely removed; it is then washed with water to remove all HOl, and dissolved in NH₃Aq; the solution is evaporated to dryness, and the NH, salt thus obtained is heated in an open crucible until every trace of NH, is driven off (Roscoe, C.N. 25, 73). Bernoulli (P. 111, 590) recommends to boil the NH, salt for a long time with successive quantities of HNO, Aq (to remove traces of Na salts), and to wash the residual WO_s quite free from acids (cf. Zettnow, P. 130, 16, 240). WO, is obtained in crystals by very strongly heating the amorphous oxide (Bernoulli, P. 111, 595; Schafarik, W. A. B. 47, 246); by dissolving WO, in molten borax (Nordenskjold, P. 114, 612); by strongly heating the oxide in a current of HCl (Debray, C. R. 55, 287); or by heating to whiteness a mixture of Na₂WO₄ and Na₂CO₂ imbedded in NaCl, when crystals of WO, sublime (D., l.c.).

Na, WO, may be prepared from wolframite by heating a mixture of 150 pts. of the finely powdered mineral with 100 pts. calcined Na, CO, and 15 pts. NaNO, for four or five hours, in an iron vessel, lixiviating with water and crystallising (v. Franze, J. pr. [2] 4, 238; cf. Huntington, B. 17, 203; Scheibler, J. pr. 83, 239).

Waddell (Am. 8, 880) prepares pure WO, from

Waddell (Am. 8, 880) prepares pure WO₂ from tungstenite, by treating the powdered mineral with aqua regia, washing the residue, fusing it with KHSO₄, dissolving in water, and fractionally ppg. by HgNO₂Aq; the first pps. are then decomposed by aqua regia, the WO₂ thus obtained is dried and fused with Na₂CO₂, the fused mass is dissolved in water, tartaric acid is added, and H₂S is passed into the solution; the filtrate Voil. IV.

from ppd. MnS, &c., is boiled till H₂S is expelled, and is then fractionally ppd. by HgNO₃Aq; the later pps. are collected, washed, and decomposed by heating in an open vessel. For a method of preparing WO₃ from wolframite v. Preparation of TUNGSTEN, p. 797.

Properties. - A heavy, canary-yellow powder; becoming orange-coloured when heated (Roscoe, C. N. 25, 73). Crystalline WO, is described as wine-yellow, very lustrous, rhombic tablets (Nordenskjold, P. 114, 612). A trace of Na tungstate gives a green colour to WO, (R., l.c.). S.G.: amorphous, 7:13 to 7:16 (Zettnow, P. 130, 16, 240; Karsten, S. 65, 394); crystalline, 7.23 at 17° (Z., L.c.); 6·38 (N., Lc.). S.H. (8° to 98°) 07983 (Regnault, A. Ch. [3] 1, 129); (22° to 52°) ·0894 (Kopp, T. 1865. 71). Melts readily in the blowpipe (Riche, A. Ch. [3] 50, 29); can be sublimed by mixing with Na CO3 and heating to whiteness (Debray, C. R. 55, 287); also by strongly heating in HCl (Schafarik, W. A. B. 47, Not decomposed at m.p. of Pt (c. 1750°) (Read, C. J. 65, 313 [1894]). On exposure to light WO₃ turns greenish, owing to partial reduction (Roscoe, l.c.; cf. Liesegang, C. C. 1865. 943). WO, is insoluble in water or acids, even in boiling conc. H2SO4; it dissolves in hot KOHAq, NH, Aq, or K2CO, Aq, also in molten K2CO, KHSO,, or K, WO,; also in molten alkali chlorides, in air, with evolution of Cl (Schultze, J. pr. [2] 21, 437, 441). WO₃ acts as an acidic oxide, forming tungstates (v. Tungsten oxyacids,

Reactions.-1. Heated in hydrogen, WO, is reduced to the blue oxide, then to WO2, and finally to W (v. Blue oxide of Tungsten, p. 800; Tungsten dioxide, p. 800; and Tungsten p. 797).—2. Reduction to WO₂ or W is effected by heating with carbon.—3. Heated with potassium or sodium, WO, gives W.-4. Reduction to the blue oxide, and then to WO₂, is brought about by zinc and hydrochloric acid, by stannous chloride, and also by heating with water and different organic compounds .- 5. Heating with carbon in chlorine produces WO,Cl, and WOCl4.--6. WOCl4 is formed by passing vapour of tungsten hexachloride over heated WO (Roscoe, C. N. 25, 63) .-- 7. A mixture of WO, and calcium chloride heated to redness in carbon dioxide gives WO₂Cl₂ (Schultze, J. pr. [2] 21, 441).—8. Mixed with carbon and heated in bromine, WO₃ gives WOBr₄; and WO₂Br₂ is formed by passing vapour of tungsten pentabromide over hot WO₃ (Roscoe, L.c. p. 73).— 9. WO, is said to dissolve in molten alkali chlorides, in air, giving off Cl; heated in absence of air (in a stream of CO₂) with chloride of calcium, cobalt, iron, magnesium, or nickel, it is said to give WO2Cl2 and MWO, (Schultze, l.c.).—10. Heating with phosphorus pentachoride produces WCl, and POCl, (Teclu, A. 187, 255); but WO.Cl, and WOCl, are also formed, according to the relative masses of WO, and PCl, and the temperature (Schiff, A. 197, 185).— 11. WO.Cl., WOCl., and CO. are formed by heating WO, in vapour of carbon tetrachloride Watts a. Bell, C. J. 33, 442).—12. Compounds of W with N, H, and O (v. p. 800) are formed by heating WO, in ammonia; and a compound of W with N and O is probably produced by heating WO, with sal ammoniac (v. p. 800).—13. WS2 is formed by passing sulphur vapour, or hydrogen sulphide, over WO, heated to whiteness (Berzelius); also by heating to redness a mixture of WO₂ with 6 pts. of mercuric sulphide (B.; Brock, J. pr. 54, 254).

HYDRATES OF TUNGSTEN TRIOXIDE. Various compounds of the form WO_s.xH₂O have been isolated; as these compounds act as acids, they are described under Tungsten oxyacids.

Tungsten oxyacids, salts, and derivatives of. Several hydrates of WO, that react as acids are known. WO₃ also combines with many acidic oxides—such as P₂O₃, B₂O₃, SiO₂, &c.—and the compounds so formed combine with basic oxides to form salt-like bodies.

TUNGSTIC ACIDS. The hydrates of WO. fall into two classes: tungstic acids, which are insoluble, or nearly insoluble, in water; and metatungstic acids, which dissolve in water. Tungstic acids. The monohydrate WO₃.H₂O₃

or orthotungstic acid WO2(OH)2, is obtained by dissolving WO, in hot alkali or alkaline carbonate solutions, and boiling these solutions with excess of a mineral acid. It is also said to be obtained by decomposing an ammoniacal solution of WO, by Cl (Dumas, A. Ch. [3] 55, 144); and also by boiling powdered wolframite with aqua regia. This acid is a yellow solid; it reddens litmus; is insoluble, or almost insoluble, in water. According to Braun (J. pr. 91, 39), it is changed to $2WO_3$. H_2O at 100° ; according to Zettnow (P. 130, 16, 240), this change is effected at 50°. The acid is slowly reduced by Zn and HClAq, the final product being WO₂ (O. v. d. Pfordten, B. **16,** <u>5</u>08).

The dihydrate WO₃.2H₂O is said to be formed by adding a mineral acid to a dilute solution of an alkali tungstate M2WO4, and drying in the wo_cl_ by water (Forcher, W. A. B. 44 [2], 173; cf. Riche, A. Ch. [3] 50, 36; Anthon, J. pr. 9, 6). Forms WO_H_O by drying over H_SO_ (Braun, J. pr. 91, 39). This hydrate is described as an amorphous, white solid; reddens litmus; somewhat soluble in water, but insoluble if a little acid is present. By boiling with dilute acids is said to give WO, H₂O.

The hemihydrate 2WO.H.O, or ditungstic acid H₂W₂O₇, is said by Braun (l.c.) to be formed by heating the monohydrate to 190°; the change is complete at 50° according to Zettnow (P. 130,

16, 240).

METATUNGSTIC ACIDS. Acids of the composition 4WO.xH2O are obtained by decomposing hot conc. BaW,O,Aq by the equivalent quantity of H2SO4, filtering from BaSO4, and evaporating (Scheibler, J. pr. 83, 310); also by decomposing PbW 0₁₈Aq by H₂S, filtering, and evaporating (Forcher, W. A. B. 44 [2], 173). The composition of the acid formed by evaporating in vacuo over H₂SO₄ is probably H₂W₄O₁,7H₂O (Scheibler, l.c.; cf. Persoz, C. R. 84, 185; Lotz, BaCl,Aq; PbW,O,, by ppg. (NH,),W,O,,Aq by Pb(C,H,O,),Aq; (NH,),W,O,, is formed by boiling (NH,),WO, with WO,Aq and evaporating.

According to Zettnow (P. 150, 16, 240), the pp. obtained by decomposing Na, WO, Aq by a mineral acid has the composition

 $4WO_3.H_2O(=H_2W_4O_{13})$ after drying at 200°.

Metatungstic acid crystallises in sulphuryellow octahedra; the acid is readily soluble in water, forming a very sour, intensely bitter liquid. A conc. solution gives a pp. of WO, 2H,O on standing; when a dilute solution is boiled it coagulates and gives a pp. of white WO₃.2H₂O, and then of yellow WO₃.H₂O. The heats of The heats of neutralisation of H₂W₄O₁₃Aq by Na₂OAq, K₂OAq, BaOAq, and SrOAq are given by Péchard (C. R. 108, 1167); the values are nearly the same as those for HNO, Aq and H2SO, Aq. When four

equivalents of base are added to H₂W₄O₁₃Aq, orthotungstates M₂WO₄ are formed (P., Lc.).

By long-continued boiling 2 pts. Na₂WO₄ with 1 pt. As₂O₅, evaporating, treating with alkali solution, and repeatedly crystallising, Lefort (A. Ch. [5] 25, 205) obtained fine, very lustrous crystals, which were very soluble in water and in alcohol; to this acid L. gave the composition H₂W₄O₁₈.6H₂O, and the name luteo-

metatungstic acid.

Colloidal (meta?) tungstic acid. By adding to 5 p.c. Na, WO, Aq rather more HClAq than sufficed to neutralise the Na, and dialysing, with addition from time to time of a little HClAq, Graham (C. J. 17, 318) obtained a solution in water of tungstic acid. The solution was not gelatinised by acids, salts, or alcohol at the ordinary temperature; it remained clear at 200°. According to Sabaneeff (J. R. 21, 515; Abstract in C. J. 58, 1215), determinations of the freezing-point of an aqueous solution of Graham's soluble acid point to the formula $3WO_3.H.O(=H_2W_3O_{10})$. As the values for mol. w. obtained by S. varied from 679 to 995, there seems little reason for adopting the formula H₂W₄O₁₀ (713) rather than H₂W₄O₁₅ (944).

TUNGSTATES. A great many tungstates

have been prepared. The proportion of basic to acidic radicle varies much in these compounds. The better studied tungstates have been divided into three classes - orthotungstates MO.WO, 8MO.7WO, paratungstates perhaps or 5MO.12WO, and metatungstates MO.4WO. but as many tungstates are known which do not fit into any of these classes it seems better to adopt a nomenclature and arrangement similar to those employed for the molybdates (cf. vol. iii. p. 423), based on the number of WO, radicles in the salts. Arranging the tungstates on this plan we get the following table. R = divalent metal, or two atoms of a monovalent metal.

(Ortho) Monotungstates RO.WO, or RWO4;

derived from the acid H2WO4.

Ditungstates RO.2WO, or RW,O,; derived from the acid H₂W₂O₇.

(8:7 tungstates 8RO.7WO, or R₂W₂O₃₄.
(Para) 5:12 tungstates 5RO.12WO₃ or R₂W₁₂O₄₁.
2:5 tungstates 2RO.5WO₃ or R₂W₂O₁₇.

3:8 tungstates BRO.8WO3 or R3W8O27 Tritungstates RO.3WO, or RW,O10; ?derived from Graham's colloidal acid.

(Meta) Tetratungstates RO.4WO, or RW4O18; derived from the acid H2W4O13.

Pentatungstates RO.5WO, or RW, O10. Hexatungstates RO.6WO, or RW.O. Octotungstates RO.8WO, or RW, O22.
The tungstates which have been most

thoroughly examined are the monotungstates or orthotungstates; the paratungstates, which are either 3:7 or 5:12 salts; and the tetra- or metatungstates.

For thiotungstates v. Tungsten, Thio-ACIDS

AND SALTS OF, p. 810.

MONOTUNGSTATES, **ORTHOTUNGSTATES** or BO.WO, or RWO, Salts of H₂WO, which is the monohydrate of WO₂. The alkali orthotungstates are obtained by dissolving WO, or WO.H.O in alkali or alkali-carbonate solutions; or by fusing WO, with alkalis, alkaline carbonates, or alkali acid sulphates, and dissolving in water. The other orthotungstates are generally obtained from solutions of the alkali salts by ppn.; also by strongly heating WO, with metallic oxides or carbonates; also by fusing the alkali salts with metallic thlorides, and washing with water (Manross, A. 81, 243; 82, 356; Geuther a. Forsberg, A. 120, 268; Schultze, A. 126, 56). The orthotungstates are insoluble in water, except the alkali salts and the Mg salt. Solutions of the alkali salts give pps. of WO₃.H₂O (yellow) or WO₃.2H₂O (white), with dilute H₂SO₄Aq, HClAq, HNO₅Aq, or H₃PO₄Aq; the pp. by HClAq dissolves in a considerable excess of conc. HClAq (Mallet, C. J. 28, 1228); cold solutions of metatungstates are not ppd. by acids. According to Marignac (A. Ch. [3] 69, 5) the pp. obtained by adding an acid to a solution of an alkali tungstate sometimes contains alkali, and sometimes the ppg. acid, according to the relative quantity of acid used, the dilution, and probably also the temperature. Orthotungstates which are insoluble in water, and the metals of which form carbonates that are insoluble in alkali carbonate solutions, can be brought into solution by fusion with alkali carbonates and treatment with water.

Solutions of orthotungstates give a white pp. with HgNO,Aq; a bluish-white pp. Cu(NO_s)₂Aq; a brown flocculent pp. on adding K, FeCy, Aq, after adding HClAq; no pp. with tincture of galls until an acid is added, when a copious chocolate-coloured pp. is formed; and a yellow pp. with SnCl2Aq, which pp. turns blue when it is warmed with a little HClAq or H.SO,Aq. For the reactions with Zn and acid, and with KCNSAq, Zn and acid, v. TUNGSTEN, DETECTION OF, p. 798.

Barium orthotungstate BaWO. Prepared by fusing 7 parts BaCl, with 4 parts NaCl and 2 parts Na2WO4, and washing with water; white octahedra (Geuther a. Forsberg, A. 120, 270); fusible with difficulty (Zettnow, P. 130, 256); S.G. 5·0035 at 13·5° (Clarke, Am. S. [3] 14, 281). Hydrates: (1) 2BaWO₄·H₂O, by ppg. a hot solution of 3Na₂O.7WO₅ by BaOAq (Soheibler, B. B. 1860, 208); (2) BaWO₄·H₂O and BaWO. 2H.O., by adding four equivalents BaO in solution to H.W.O., Aq (Péchard, C. R. 108, 1167).
Cadmium orthotungstate CdWO. Obtained

as a yellow crystalline powder by fusing together 11 parts CdCl₂, 16 parts NaCl, and 4 parts Na₂WO₄ (G. a. F., l.c., p. 268; Z., l.c., p. 240). The dihydrate CdWO₄.2H₂O is obtained by ppg. a solution of a salt of Cd by Na₂WO₄Aq (Anthon, J. pr. 8, 399; 9, 337; Smith a. Bradbury, B. 24, 2935).

Calcium orthotungstate CaWO. Occurs native as scheelite, S.G. 6.02 (Bernoulli, J. 13, 783). Obtained by adding K.WO, Aq to CaCl, Aq. Quadratic octahedra are formed by fusing Na₂WO₄ with excess of CaCl₂ (Manross, A. 81, 243; 82, 348); also by heating the ppd. salt with CaO in a stream of HCl (Debray, C. R. 55,

Chromium orthotungstate Cr_s(WO₄)_{s.20}H_sO. Obtained as a green pp. by adding Na₂WO₄Aq to CrCl₈Aq; loses 13H₂O at 100° (Lotz, A. 91, 66).

Cobalt orthotungstate CoWO.2H.O. violet powder; obtained by ppg. Co salt solutions

by K₂WO₄Aq (Anthon, J. pr. 9, 344).

Copper orthorungstate CuWO.2H,O. A green powder, melting at a red heat and cooling to chocolate-coloured crystals; formed by ppg.

Cu salts by Na, WO, Aq (A., l.c.).

Ferrous orthotungstate FeWO. Obtained as opaque crystals by heating WO, and Fe₂O, in a stream of HCl (Debray, C. R. 55, 288); also by fusing 2 parts FeCl₂, 2 parts NaCl, and 1 part Na.WO4 (Geuther a. Forsberg, A. 120, 273; Zettnow, P. 130, 30). The trihydrate FeWO.3H2O was obtained by Anthon (J. pr. 9, 343) as a brown powder by ppg. a ferrous salt by K2WO4Aq. Double compounds of the form $m\text{FeWO}_{\bullet}.n\text{MnWO}_{\bullet}$ with the ratios m:n=7:1, 4:1, 3:1, 3:2, 1:2, and 1:7 were obtained by fusing FeCl₂, MnCl₂, NaCl, and Na₂WO₄ in varying proportions (G. a. F., l.c. p. 270; cf. Z., l.c. p. 250). Wolframite has approximately the composition FeO.MnO.WO.

Lead orthotungstate PbWO. Occurs native as scheeletine; obtained, as a white pp., by adding K₂WO₄Aq to solution of a salt of Pb (A., l.c., p. 342); quadratic octahedra are formed by fusing 10 parts Na2WO, with 47 parts PbCl,

S.G. 8.235 (Manross, A. 82, 357).

Magnesium orthotungstate MgWO. Formed, in white octahedra, by fusing 2 parts MgOl, with 2 parts NaCl and 1 part Na,WO, (G. a. F., *l.c.* p. 272).

Manganese orthotungstate MnWO. net-brown, lustrous, rhombic crystals; S.G. 6.7; obtained by fusing 2 parts MnCl₂, 2 parts NaCl, and 1 part Na₂WO₄ (G. a. F., *l.c.*; Z., *l.c.*).

Mercurous orthotungstate Hg₂WO₄. A

yellow pp., formed by adding solution of a mercurous salt to solution of an alkali tungstate

(A., l.c.).
Nickel orthotungstate NiWO4. Formed by fusing 2 parts NiCl₂ with 2 parts NaCl and 1 part Na₂WO₄; brown, lustrous crystals (Schultze, A. 126, 56); S.G. 6.8845 at 20.5°, 6.8522 at 22° (Clarke, Am. S. [3] 14, 281). The hexahydrate, NiWO₄.6H₂O, is obtained by ppn. (A., l.c.).

Potassium orthotungstate K2WO4. triclinic needles; obtained by boiling WO. with an equivalent weight of KOHAq or K,CO,Aq, and evaporating (Marignac, A. Ch. [3] 69, 18: Anthon, J. pr. 8, 899; 9, 337). Decrepitates when heated; melts at red heat (M., i.c.), taking up CO₂ (Ullik, W. A. B. 56 [2], 148; cf. Knorre, J. pr. 27, 89). Various hydrates are obtained by crystallising a solution of WO, in K.CO.Aq under different conditions (M., l.c.; A., Lc.; Riche, A. Ch. [3] 50, 45).

Silver orthotungstate Ag₂WO₄. yellow, amorphous pp. by adding Na, WO, Aq to solution of a salt of Ag; melts below redness, and becomes crystalline on cooling; easily soluble in HNO,Aq; soluble in NH,Aq (Zettnow, P. 130, 30; cf. Muthmann, B. 20, 984; Wöhler a.

Rautenberg, A. 114, 120). By evaporating a solution in NH₂Aq over CaO and NH₂Cl, Widmann obtained the compound Ag2WO4.4NH3

(Bl. [2] 20, 64).

Sodium orthotungstate Na2WO4.2H2O. Obtained by crystallising a solution of WO, in the equivalent quantity of NaOHAq or Na,CO,Aq. Transparent, colourless, rhombic tablets (Marignac, A. Ch. [3] 69, 22). Effloresces in vacuo, or in dry air. Loses all H₂O at 100°. S.G. 8-2588 at 17.5°, 3-2314 at 19°; S.G. of Na₂WO₄ = 4.1743 at 20.5° , 4.1833 at 18.5° (Clarke, Am. S. [3] 14, 281). Solubility = 41 at 0° , 55 at 15°, 123.4 at 100° (Riche, J. pr. 69, 10). For S.G. of Na₂WO₄Aq of different concentrations, v. Franz (J. pr. [2] 4, 238). For E.C. of Na₂WO₄Aq v. Walden (Z. P. C. 1, 529).

Strontium orthotungstate SrWO. Obtained by ppn (Anthon, l.c.); also by fusing 2 parts SrCl₂, 2 parts NaCl, and 1 part Na₂WO₄ (Zett-

now, l.c.; Schultze, A. 126, 56).

Zine orthotungstate ZnWO. Colourless tablets; formed by fusing 2 parts ZnCl₂ with 2 parts NaCl and 1 part Na₂WO₄ (Geuther a. Forsberg, A. 120, 270; cf. Zettnow, P. 130,

Paratungstates. A number of acid tungstates have been isolated, the composition of which is expressed by one of the formulæ 3RO.7WO, or 5RO.12WO₃. The former formula was given by Lotz (A. 91, 49) and Scheibler (J. pr. 83, 273); the latter formula by Laurent (A. Ch. [3] 21, 54) and Marignac (A. Ch. [3] 69, 5). The two formulæ represent salts of almost identical composition, and no way has been found of deciding which formula is the better (cf. Knorre, J. pr. [2] 27, 83). The acid tungstates of this series are more easily formed and crystallise better than the other acid salts; they are generally called paratungstates, following the suggestion made by Marignac.

The alkali paratungstates are obtained by saturating hot alkali solutions with WO₃ and crystallising; most of the other paratungstates are prepared by ppn. from solutions of the alkali salts. The alkali salts are soluble in water, the other salts are insoluble. Solutions of alkali paratungstates give, with dilute HClAq, white pps. which become yellow on boiling $(WO_{s}.x\hat{H}_{s}O).$ Metatungstates, RO.4WO, are formed by gradually adding HClAq to boiling solutions of paratungstates; meta-salts are also formed by boiling para-salts with WO3.H2O. Solutions of paratungstates are not ppd. by SO2Aq, HIAq, citric, tartaric, or oxalic acid; but the presence of one of these acids does not hinder ppn. by HClAq. Solutions of alkali paratungstates give pps. with salts of almost all the heavy metals. H₂S reacts with alkali paratungstates in solution to form solutions of thiotungstates which give brown pps. with acids, soluble in (NH₄)₂SAq. Paratungstates eadily react with H₂PO₄, H₃AsO₄, &c., to form complex acids (v. p. 807), solutions of which are not ppd. by acids, except these are added in considerable excess; solutions of paratungstates are, there-Hore, not ppd. by HClAq in presence of H₂PO₄, H₂AsO₄, H₂B₂O₄, &c. For the reactions of paratungstates with PtO₄H₄ v. Rosenheim (B. 24, 2397; the reaction is rather indefinite).

If the simpler (8:7) formula is adopted, the

paratungstates are represented as 3R120.7WO. 3Ruo.7Wo, and Ruoo.7Wo, i.e. as salts of the hypothetical acid How, O24, wherein 6H are replaced by 6R1, 3R11, or 2R111; if the less simple formula (5:12) is adopted, the paratungstates of monovalent and divalent metals are regarded as alts of the hypothetical acid $H_{10}W_{12}O_{11}$ = 12 WO_3 .5 H_2O), wherein 10H are replaced by 10R1 or 5R11; paratungstates of R111 must be represented in this scheme by the complicated formula R^{III}₁₀(W₁₂O₄₁)₃. In the paratungstates which have been best examined R^I = NH₄, Li, Hg, K, Ag, and Na; RII = Ba, Cd, Ca, Co, Cu, Pb, Mg, Mn, Ni, Sr, and Zn; and RIII = Al and Cr. The simpler formulæ are employed in the descriptions of the individual salts.

Aluminium paratungstate Al₂W₇O₂₄.9H₂O. A flocculent pp., drying to gum-like lumps; obtained by adding (NH₄)₆W,O,Aq to a solution of a salt of Al (Lotz, A. 83, 65; Anthon, J. pr.

8, 399; 9, 337).

Ammonium paratungstate

(NH₄), W,O₂₄.6H₂O. Obtained by dissolving WO, H₂O in NH, Aq and crystallising at the ordinary, or a little above the ordinary, temperature (Lotz, A. 91, 55). Crystallises in white, rhombic needles (Marignac, A. Ch. [3] 69, 25; Kerndt, J. pr. 41, 190); also in rhombic tablets (M., l.c.); and is, therefore, dimorphous. Solubility = 2.8 to 4.5 at $15^{\circ}-22^{\circ}$; when the solution is boiled the very soluble meta-salt (NH₄)₂W₄O₁₂ is formed. Loses 3.9 p.c. H₂O at 100°; when heated to redness the blue oxide of W remains mixed with compounds containing N Marignac (l.c.) gave the formula (NH₄)₁₀W₁₂O₄₁.11H₂O to this salt; Berzelius, and Anthon (l.c.) gave the formula (NH₄)₂W₂O₇.H₂O; and Riche (v. Marignac, l.c.) the formula $(NH_4)_2W_4O_{13}.5$ or $6H_2O$.

Various double salts have been isolated; with Na₆W,O₂₁ (v. Knorre, B. 19, 821; Gibbs, Am. 7, 236); with K₆W,O₂₄ (v. Laurent, A. Ch. [3] 21, 59; Marignac, A. Ch. [3] 69, 55); with $Mg_3W_7O_{24}$ v. M. (l.c.), Lotz (A. 91, 61); with $Zn_3W_7O_{24}$ v. Lotz (l.c.); with $Cd_3W_7O_{24}$ v. L. (l.c.); with Hg₃W₇O₂₄ v. Anthon (J. pr. 8, 399;

9, 337).

Barium paratungstate Ba₃W₇O₂₄.8H₂O. A white pp. formed by adding (NH₄)₆W₇O₂₄Aq to excess of BaCl₂Aq, washing and drying over H₂SO₄ (Lotz, A. 93, 60; cf. Knorre, B. 18, 327; 19, 820). Forms a double salt with Na, W,O, (Scheibler, J. pr. 83, 314).

Cadmium paratungstate Cd, W,O24.16H2O; white needles, obtained by adding Na, W,O,,Aq to excess of solution of a salt of Cd (Gonzalez,

J. pr. [2] 36, 44).

Calcium paratungstate Ca, W,O2.18H2O; a crystalline pp. formed by ppg. excess of CaCl,Aq by Na_eW,O₂₄Aq, and drying in the air (Knorre, B. 18, 328). Forms a double salt with Na_eW,O₂₄ (Gonzalez, l.c.).

Chromium paratungstate Cr₂W₇O₂₄.9H₂(); a grey powder, obtained by heating CrCl, with (NH₄)₈W₂O₂₄; insoluble in water, but dissolves in CrCl₂Aq (Lotz, A. 91, 66).

Cobalt paratungstate Co, W,O,, 25H,O (Gonzalez, l.c.); forms a double salt with Na, W, O24

(G., l.c.).

Copper paratungstate Cu, W,O,4.19H,O; a pale-green pp. formed by adding Na, W,O,Aq to

excess of CuSO, Aq (Knorre, B. 19, 826; v. also Gonzalez, J. pr. [2] 36, 44). Forms a double salt with Na₈W₂O₂₁ (K., l.c.).

Lead paratungstate Pb, W,O24; a white pp. formed by adding (NH₁)_BW₂O₂₄Aq to Pb(NO₃)₂Aq (Lotz, A. 91, 49). Forms a double salt with $Na_6W_7O_{24}$ (Gonzalez, *l.c.*).

Lithium paratungstate Li₆W₇O₂₁.19H₂O. Large prisms, unchanged in air; obtained by boiling the proper quantities of WOs and LizCOs in water, and allowing to crystallise (Scheibler, J. pr. 83, 321).

Magnesium paratungstate Mg3W7O21.24H2O; obtained by ppn. (Knorre, B. 19, 824). Forms a double salt with Na, W,O24 (K., l.c.).

Manganese paratungstate Mn₃W₇O₂₄,34H₂O (Gonzalez, J. pr. [2] 36, 44). Forms a double salt with Na, W,O24 (K., l.c.).

Nickel paratungstate Ni₃W₂O₂₁.14H₂O (Anthon, J. pr. 9, 344).

Potassium paratungstate K₆W,O₂₁.6H₂O (Marignac, A. Ch. [3] 69, 33, gave the formula K₁₀W₁₂O₄₁.11H₂O). Prepared by saturating hot KOHAq with WO₃.xH₂O, and allowing to cool; by boiling K2WO, with H2O; or by fusing wolframite with 3 of its weight of K2CO3, boiling the product with water, saturating the aqueous solution with CO2, and crystallising (M., White rhombic crystals, isomorphous with the NH, salt (M., l.c.). Decomposed, by melting, to K₂WO, and 5K₂O.14WO, without forming any meta-salt (Knorre, J. pr. [2] 27, 91). Solubility = 2.15 cold water, 6.6 boiling water.

Various hydrates have been obtained (v. Zettnow, P. 130, 241; cf. Lefort, A. Ch. [5] 9, 93; 15, 321; 17, 470; 25, 200). Forms double salts with Na, W, O24 (v. Marignac, A. Ch. [3] 69, 55; Laurent, A. Ch. [3] 21, 54).

Silver paratungstate Ag, W,O,,; a yellow, crystalline pp. formed by adding Na, W,O, Aq to excess of AgNO₃Aq (Gonzalez, J. pr. [2] 36, 44).

Na. W, O24.16H2O. Sodium paratungstate This formula is given by Scheibler (J. pr. 83, 285); the formula Na₁₀W₁₂O₄₁ is given by Laurent (l.c.), Marignac (l.c.), and by Friedheim a. Meyer, (Zeit. f. anorg. Chemie, 1, 81). Prepared by satu-rating hot NaOHAq, or Na₂CO₃Aq, with WO, xH,O and evaporating to the crystallising point; also by passing CO2 into Na2WO4Aq until the reaction is only faintly alkaline. Large, well-formed, white, triclinic crystals (Scheibler, L.c.; Marignac, A. Ch. [3] 69, 39). Effloresces in air; loses c. \(\frac{3}{4}\) of its H₂O over H₂SO₄; is dehydrated without decomposition at 300° (S., l.c.); heated to dull redness leaves a residue, insoluble in water, containing more than 5WO, to Na₂O (M., l.c.; also Knorre, J. pr. [2] 27, 71). Melts above dull redness, giving Na₂WO, and Na₂W,O₁₈ (meta-salt). Heated with water to 150° gives much meta-salt (K., l.c.). Solubility 8 to 9 at 35° to 40°; for the salts obtainable from an aqueous solution v. M. (l.c.), and also K. (B. 18, 2362). By boiling a solution with Na₂CO₃ transformation to Na₂WO₄ is complete (v. Schmidt, Am. 8, 16). A hydrate with 21H,0 has been obtained (M., l.c.). Forms double salts with Sr. W.O. (Gonzalez, J. pr. [2] 36, 44), and with Zn, W,O₃₄ (Knorre, B. 19, 823). Strontium paratungstate Sr₂W,O₃₄.8H₂O

(Knorre, B. 18, 827).

Zinc paratungstate Zn, W,O21.xH2O (Gonzalez, J. pr. [2] 36, 44).

METATUNGSTATES or TETRATUNGSTATES. Ruo.4WO, or RuW4O18. Salts of the acid H₂W₄O₁₃ (v. METATUNGSTIC ACIDS, p. 802). The metatungstates may be obtained by causing metatungstic acid H₂W₄O₁₃.xH₂O (v. p. 802) to interact with carbonates, chlorides, or nitrates; also by the reaction of a weak acid (H,PO,Aq is the best) with orthotungstates, the acid being added as long as the pp. of WO.xH.O which is at first formed re-dissolves: also from the alkali salts by double decomposition, or better from the Ba salt by interacting with sulphates; the alkali metatungstates are prepared by boiling solutions of the ortho-tungstates for some time with WO₃.xH₂O.

Most of the metatungstates are very soluble in water and are crystallisable from aqueous solutions; aqueous solutions are not ppd. by acids in the cold, but pps. are obtained after long standing or boiling. Metatungstates are changed to ortho- salts by excess of alkali; when strongly heated, orthotungstates are formed. Solutions of metatungstates are not ppd. by H₂S; (NH₄)₂SAq gives a blue pp.; pps. are not obtained with salts of the heavy metals except Pb and Hg1; K, FeCy, Aq does not form a

Ammonium metatungstate

(NH₁)₂W₄O₁₂.8H₂O. This formula is given by Scheibler (J. pr. 83, 303); more complicated formulæ are given by Margueritte (A. Ch. Lotz (A. 91, 55). Obtained by boiling the para- salt (NH₄)₆W₄O₁₂.xH₂O with WO₂.H₂O (Margueritte, A. Ch. [3] 17, 477; also by boiling the para-salt with a little HNO, Aq (Laurent, A. Ch. [3] 21, 62); also by heating the dry parasalt to 250°-300° (Scheibler, J. pr. 83, 304; cf. Persoz, A. Ch. [4] 1, 101; and Marignac, A. Ch. [4] 3, 71). White octahedra; effloresces in air; loses 7H2O at 100°. Solubility in cold water = 120 (Lotz, l.c.), = 286 (Riche, A. Ch. [8] 50, 45). The solution is optically refractive. Insoluble in alcohol or ether. Forms a double salt with NH, NO, (Marignac, A. Ch. [3] 69, 61).

Barium metatungstate BaW, O18.9H2O. Obtained by adding BaCl₂Aq to a hot conc. solution of Na₂W₄O₁₈ acidified by HClAq (Scheibler, J. pr. 83, 304). Crystallises in large, white, tetragonal octahedra; S.G. 4298 at 14°; loses 6H₂O at 100°; easily soluble in hot water, decomposed by much cold water to WO, and BaW,O10.6H2O, which dissolve again on heating.

Cadmium paratungstate CdW,O18.10H2O. Lustrous, white octahedra; unchanged in air; obtained by decomposing BaW₄O₁Aq by CdSO₄Aq (S., *l.c.*, p. 273).

Calcium metatungstate CaW.O13.10H2O. Quadratic octahedra; obtained by dissolving CaCO, in H₂W₄O₁₂Aq and crystallising (S., l.c., p. 314).

Cobalt metatungstate CoW4O12.9H2O (S., *l.c.* p. 317).

Copper mctatungstate CuW,O,,.11H,O (S., l.c p. 317).

Magnesium metatungstate MgW,O,, 8H,O

(S., l.c.).
Manganese metatungstate MnW₄O₁₁,10H₂O (S., l.c. p. 273).

Mercurous metatungstate Hg₂W₄O₁₃.25H₂O. Obtained, as a white pp., by adding HgNO, Aq to solution of H₂W₄O₁₈ or a meta-salt (S., *i.c.*, p. 319).

Nickel metatungstate NiW4O12.8H2O (S., l.c.,

p. 273).

Potassium metatungstate K.W.O.s.8H.O. Lustrous octahedra; obtained by boiling K, W, O24Aq with WO3.xH2O; very soluble in hot water, considerably less soluble in cold water (S., l.c., p. 303). A pentahydrate was obtained by Marignac (A. Ch. [4] 3, 71).

Silver metatungstate Ag₂W₄O₁₃. Obtained, as a crystalline crust, by adding AgNO.Aq to Na, W.O. Aq, evaporating, separating from Ag2WO4, and evaporating again (S., Lc. p. 318). metatungstate Na₂W₄O₁₈.10H₂O. Sodium Obtained by boiling Na, W, O, Aq with WO, xH2O and allowing to evaporate; white, lustrous octahedra; S.G. 38647 at 13°; very soluble in

hot, but less in cold, water (S., l.c. p. 303) Strontium metatungstate SrW,O18.8H2O (S.,

l.c. p. 314). Zinc metatungstate ZnW₄O₁₃.10H₂O (S., l.c. p. 273).

TUNGSTATES OTHER THAN ORTHO-, PARA-, AND META- SALTS.

Ditungstates. Salts οf the form $R_{2}^{1}O.2WO_{2} = R_{2}^{1}W_{2}O_{7}$ Salts of K and Na of this composition were said by Lefort (A. Ch. [5] 17, 470) to be formed by neutralising cold solutions of K₂WO₄ and Na₂WO₄ by acetic acid; Knorre (J. pr. [2] 27, 83) obtained only metatungstates, R₂W₄O₁₃, by this method.

Two to five tungstates. The existence of salts of the class 2R, 10.5WO, = R, 1W, O,, is doubtful. Marignac obtained a salt to which he gave the formula Na₄W₅O₁₇.11H₂O along with the para-salt $Na_6W_7O_{24}.16H_2O(A.Ch.[3] 69, 50)$. The same salt seems to have been obtained by Lefort (l.c.; cf. Forcher a. Gibbs, J. 1880. 341) by adding acetic acid and alkali to Na2WO.Aq and allowing to stand for some time (cf. also Schmidt, Am. 8, 16).

Three to eight tungstates. A salt (NH₄)₆W₂O₂₇.8H₂O = 3(NH₄)₂O.8WO₃.8H₂O seems to have been prepared by Marignac (A. Ch. [3]

 $R^{11}O.3WO_3 = R^{11}W_3O_{10}$. Tritungstates Lefort (A. Ch. [5] 17, 470) prepared a salt to which he gave the formula Na₂W₂O₁₀.6H₂O, by pouring a boiling solution of Na₂W₂O₇.6H₂O (obtained by adding acetic acid to Na2WO4Aq until the liquid shows an acid reaction) into boiling acetic acid, allowing to cool, separating the lower syrupy layer, and crystallising it. A corresponding salt $K_2W_2O_{10}.2H_2O$ was obtained by a similar method; and a series of tritungstates was prepared by mixing equivalent quantities of the Na salt and metallic acetates, in solution, and adding alcohol. L. describes the tritung states as easily decomposing in aqueous solutions to diand tetra-salts. The following salts were formed by L.:—BaW₂O₁₀.4H₂O; CdW₃O₁₀.4H₂O; CdW₃O₁₀.4H₂O; CoW₂O₁₀.4H₂O; FeW₂O₁₀.4H₂O; PbW₃O₁₀.2H₂O; MgW₃O₁₀.4H₂O; MnW₃O₁₀.5H₂O; NiW₃O₁₀.2H₂O; MgW₃O₁₀.4H₂O; Na₂W₃O₁₀.4H₂O; SrW₃O₁₀.5H₂O. Knorre's experiments led him to regard Lefort's tritungstates as mixtures (J. pr. [2] 27, 83).

Penta- and octo- tungstates.

of K and Na of the composition R2W8O16 and R₂W₈O₂₅ are said to have been isolated, the former by fusing mixtures of R₂WO₄ and WO₅, the latter by fusing R₂W₄O₁₅ w th R₂WO₄ (Knorre, J. pr. [2] 27, 81, 91).

A few tungstates that do not find places in any of the foregoing classes have been described by different observers (v. especially Lefort, A. Ch. [5] 9, 93; 15, 324; 17, 470; 25, 200; also Scheibler, J. pr. 83, 237; Gonzalez, J. pr. [2] 36, 44; Knorre, J. pr. [2] 27, 93; Lotz, A. 91, 49; Feit, B. 21, 133; Cleve, Bl. [2] 43, 170; Högborn,

Bl. [2] 42, 2).

PERTUNGSTATES. By boiling a solution of the paratungstate Na, W,O₂₄, 16H₂O with H₂O₂Aq, and then evaporating in vacuo, Péchard (C. R. 112, 1060) obtained small white crystals of a salt to which he gave the formula Na.O. $W_2O_{12}H_2O =$ NaWO₄.H₂O; and by treating (NH₄)₈W,O₂₄.6H₂O in a similar way he isolated the corresponding NH, pertungstate. According to P. these salts are decomposed by alkalis with evolution of O; solutions of them set free I from KIAq, and

react with HClAq, giving off Cl.

TUNGSTEN BRONZES. Bright-coloure, lustrous, metal-like, crystalline solids, obtained by the action of reducers—such as H, coal-gas, Sn, or Fe-on tungstates of Li, K, or Na; also by the electrolysis of these tungstates when molten. The composition of these compounds, which are known as bronzes and are used as pigments, is represented by the formula $M_x(WO_s)_y$, where M = Li, K, or Na. The constitutions of the compounds are not known; they may be represented as compounds of tungstates with WO₂, by the general formula $xM_2O.yWO_2.sWO_2$; they may also be represented as compounds of M2O with a radicle composed of W and O in a ratio greater than W:20 and less than W:30—that is, as compounds of M₂O with oxides intermediate between WO₂ and WO₃. The tungsten bronzes are insoluble in most acids, also in alkali solutions.

Sodium tungsten bronzes. These compounds are formed by the interaction of Na tungstates and H (Wöhler, P. 2, 350), Sn (Wright, A. 79, 221), coal-gas (Schnitzler, D. P. J. 211, 484), Zn or Fe (Zettnow, P. 130, 261), or by electrolysing Na tungstates (Scheibler, J. pr. 83, 321). According to Philipp (B. 15, 499), the different processes yield the same bronze if the same tungstate is used, and the more WO, there is in the tungstate employed the richer in WO, is the bronze produced. All the compounds are insoluble in acids or alkalis; they dissolve in NaClOAq; heated with NHAq and AgNO₃Aq they give Ag and WO₃ (cf. P. a. Schwebel, B. 12, 2234; 15, 500; Knorre, J. pr. [2] 27, 51). These compounds are slowly oxidised to tungstates by heating to redness in air (K., l.c.). Philipp (i.c.) gave S.G. of all the sodium bronzes as 7.2 to 7.8 at 16° to 18° .

Purple-red bronze Na, W₂O₅ = Na₂O.2WO₂.WO₂ = Na₂O.W₃O₅ = Na₂WO₄.W₂O₅. Prepared by fusing for c. half an hour a mixture of 10.9 g. Na₂CO₃, 71.7 g. WO₃ and 20 g. tinfoil, and boiling successively with H₂O₃ NaOHAq, and HClAq. Red cubes; the powder transmits green light

when suspended in water (P., l.c.).

Red-yellow bronze Na, W, O₁₅ =

2Na, O.8WO, 2WO, = 2Na, O.W, O₁₈. Prepared by

melting 60 to 80 g. of a mixture of Na₂WO₄ and WO, in the ratio 2Na2WO4:WO, adding 30 g. tinfoil, and keeping molten for 1 to 2 hours. Red-yellow cuber; the powder is brownish yellow, and transitits blue light when suspended in water (P., l.c.).

Gold-yellow bronse Na, W.O. = $5Na_2O.7WO_3.5WO_3 = 5Na_2O.W_{12}O_{31}$. Prepared by heating Na, W,O, to dull redness in H, powdering and again heating in H, and then boiling with H₂O, HClAq, and Na₂CO₃Aq successively. Golden yellow, crystalline powder (P., l.c.; cf. Wöhler, P. 2, 350; Wright, A. 79, 221).

Blue bronze Na W O 15 = Na O.4 WO 3. WO 2 = Na₂O.W₃O₁₄. Prepared by fusing Na₂WO₄ with more than 2WO₃, and adding tinfoil; also by melting Na₂W₂O₂₄ (paratungstate) and electrolysing with 6 Zn-Pt elements (P., l.c.; Scheibler, J. pr. 83, 321; cf. Knorre, J. pr. [2] 27, 49; and Zettnow, P. 130, 261). Dark-blue cubes, with a

red sheen; S.G. 7.28 at 17°.

Potassium tungsten bronzes. The com- $K_2W_4O_{12} = K_2O.3WO_8.WO_2 = K_2O.W_4O_{11}$ pound seems to be the only one of this class that has been isolated. Prepared by fusing K₂WO₄ with WO, and adding tinfoil; also by fusing K,CO, with from 3WO, to 4WO, and reducing by H or coal gas; also by electrolysing a molten mixture of K2WO4 and WO3. Reddish-violet prisms, giving a blue powder, which transmits greenish light when suspended in water; S.G. c. 7.1 (Laurent, A. Ch. [2] 67, 219; Zettnow, P. 130, 262; Knorre, J. pr. [2] 27, 63).

Lithium tungsten bronzes. A compound of this class, probably Li₂W₅O₁₅, is obtained by fusing Li₂W₇O₂₄ (paratungstate) with tin. Darkblue crystals (Scheibler, J. pr. 83, 321; Knorre,

l.c., p. 69; Feit, B. 21, 133).

A potassium sodium bronze and a potassium lithium bronze have been obtained (Knorre, l.c.

p. 66; Feit, l.c.).

COMPLEX TUNGSTIC ACIDS SALTS. Tungstic oxide WOs combines with several anhydrides, e.g. B₂O₃, P₂O₅, As₂O₅, SiO₂, &c., and water, to form acidic compounds, and also with these anhydrides and basic oxides to form salt-like compounds; the whole of these compounds are usually grouped together as complex tungstic acids and complex tungstates, and they are divided into such classes as arsenotungstates, phosphotungstates, &c.

ANTIMONOSOTUNGSTIC ACIDS AND SALTS. ACcording to Lefort (A. Ch. [5] 17, 487), the compounds Sb₂O₃.5WO₃.4H₂O and Sb₂O₃.6WO₃.8H₂O are formed by dissolving tartar emetic in Na,O.8WO,Aq and Na,O.2WO,Aq respectively. Gibbs (Am. 7, 892) obtained

4BaO.6Sb,O,22WO,.36H,O and 6K,O.4Sb,O,.12WO,.25H,O.

Arsenotungstic acids and salts. Compounds of the form As₂O₅16WO₅.xH₂O have been obtained by Kehrmann (A. 245, 45; cf. Fremery, B. 17, 296); and series of compounds of As₂O₃ with WO, and besic oxides (Na₂O, K₂O, BaO, &c.), in which the ratio of As₂O₃. WO, is 1:6 and 1:8, have been described by Gibbs (Am. 7, 813). Compounds of WO, with As,O, and bases, and also with As,O, and As,O, and bases, and finally with As,O, and P,O, and bases, are described by Bibbs (l.c.).

· Borotungstic acids and sairs. Two com-

pounds of B₂O₂ and WO₂ have probably been isolated, in which the ratios of the oxides are 1:14 and 1:9 respectively; compounds of each of these with bases are known (Klein, A. Ch. [5] 28, 350).

Borotungstic acid and salts

B₂O₃.14WO₃.xH₂O. A solution containing B₂O₃ and WO, in the ratio 1:14 is obtained by boiling Na₂WO₄Aq with B₂O₂Aq, oppg. by HgNO₂Aq, decomposing the pp. by H₂S, and expelling H₂S from the filtrate by warming (for details v. K., l.c.). Compounds of the form xMO.B₂O₂.14WO₂.yH₂O have been formed, where x = 2, 3, and 4, and M is Ba, K_2 , Ag_2 , Na_2 and Sr.

Tungstoboric acid and salts. The compound B₂O₃.9WO₃.xH₂O, usually known as tungstoboric acid, is obtained in yellowish octahedra by evaporating the solution of borotungstic acid. A considerable number of compounds of bases with B₂O₃ and WO₃, in the ratio B₂O₃:9WO₃, has been obtained; most of them are of the corn 2MO.B₂O₂.9WO₃.xH₂O, M=(NH₂)₂, Ba, Cd, Ca, Co, Cu, Li₂, Mg, Mn, Hg₂, Ni, K₂, Na₂, Tl₂; a few salts 2M₂O₃.B₂O₃.9WO₃.xH₂O (M=Al or Cr) are also described by Klein (l.c.).

FLUOTUNGSTATES and allied compounds; v.

Tungstoxyfluorides (p. 799).

IODOTUNGSTATES. Blomstrand (J. pr. 40, 327) described a compound 2K₂O.I₂O₃.2WO₅.3H₂O, obtained by adding the calculated quantity of HIO, to K, WO, Aq.

PHOSPHOTUNGSTIC ACIDS AND SALTS. The oxides WO, and P2O, combine in several proportions, in presence of H₂O, to form complex acids which contain large quantities of WO, relatively to the P₂O₅. A great many compounds are known containing WO₃ and P₂O₅ combined with basic oxides.

Phosphotungstic acids. Compounds of the form P₂O₃.xWO₃.yH₂O; compounds wherein r=24, 21, 20, 16, and 12 seem to have been isolated. The existence of these compounds was made known by Scheibler (B. 5, 802); they have been investigated chiefly by Gibbs (Am. 2,217, 281; 4, 377; 5, 361, 391; 7, 313, 392; Péchard, C. R. 109, 301; 110, 754; and Kehrmann, B. 20, 1805, 1811; 24, 2326; 25, 1966; A. 245, 45; Zeit. f. anorg. Chemie, 1, 428; also Drechsel, B. 20, 1452; and Brandhorst a. Kraut, A. 249, 373). The phosphotungstic acids are obtained by boiling H₂PO₄Aq with H₂W₄O₁,xH₂O (meta-acid); by decomposing the mercurous salts by HClAq; and by boiling Na₄W₄O₂₄xH₂O (para-salt) with Na₂HPO₄Aq, decomposing by HClAq, and dissolving the acids thus formed in ether. The phosphotungstic acids are unchanged in solutions in dilute acids; these solutions are scarcely acted on by H2S, and very slightly by zinc; they give phosphates and tungstates when boiled with alkalis; characteristic pps. are produced with alkaloids, urea, albumen, &c.

acid PhospLoduodecitungstic and salts P.O. 24WO. xH2O and yMO.P.O. 24WO. xH2O. The acid is obtained by evaporating a solution of H.PO, and H.W.O, xH.O in the proper proportion (Péchard, l.c.); by boiling a solution of tungstate and phosphate of Na, in the ratio 24Na, WO,:2Na, HPO, acidulating with HNO,Aq, ppg. by HgNO,Aq, decomposing the Hg salt by HClAq, filtering, and evaporating in vacue

(Gibbs, Am. 2, 217); by boiling H,PO,Aq with BaWO, in the proper proportion, decomposing by a small excess of H₂SO, Aq, removing H₂SO, by BaOAq, filtering, and evaporating (Sprenger, J. pr. [2] 22, 418). The acid crystallises in white, regular octahedra, or cubes. The value for x in the formula $P_2O_8.24WO_8.xH_2O$ varies according to different observers, from 40 to 53, 59, and 61. The salts yMO.P₂O₅.24WO₈.xH₂O are obtained by mixing tungstates (normal or para-salts) with H₃PO₄Aq or a phosphate, and decomposing by slight excess of HClAq, HNO₅Aq, or H2SOAAq; in most cases the salts ppt. as fine, white powders; in some cases it is advisable to evaporate to dryness, extract with alcohol and ether, and evaporate. The salts have been described by Gibbs (l.c.); Kehrmann (Zeit. f. anorg. Chemie, 1, 430); K. a. Freinkel (B. 24, 2327); Sprenger (J. pr. [2] 122, 418); and Brandhorst a. Kraut (A. 249, 373). Salts have been obtained wherein $MO = (NH_4)_2O$, BaO, CuO, K_2O , Ag₂O, and Na₂O, and y = 1, 2, and 3.

Phospholuteotungstic acid salts. and P₂O₅.16WO₃.xH₂O and yMO.P₂O₅.16WO₃.xH₂O. The acid is obtained by boiling Na₂WO₄Aq with H,PO,Aq, acidulating with HNO,Aq, adding NH₄Cl and boiling, crystallising the pp. from NH, ClAq, evaporating the NH, salt thus produced with aqua regia, and crystallising the acid so formed from water (for details v. K., l.c.). To an acid obtained by decomposing the K salt by H₂SiF_sAq, or the Ag salt by HClAq, K gives the formula P₂O_s.18WO_s.xH₂O; and he says that the salts are yMO.P₂O₃.18WO₃.xH₂O. The acid forms citron-yellow, six-sided tablets; it melts by the heat of the hand, and is very soluble in water. According to K. (B. 20, 1805; A. 245, 45), salts of this series are obtained by boiling any phosphotungstates with excess of H,PO,Aq in presence of an alkali salt. It is probable that two series of salts exist—one with 16WO, and another with 18WO, (cf. Gibbs, l.c.; and Pechard, l.c.). Salts have been obtained wherein $MO = (NH_4)_2O$, BaO, CaO, CuO, PbO, and K_2O , and y = 1, 3, 4,and 6.

Other phosphotungstic acids and salts.

I. Salts of the form yMO.P₂O₅.22WO₃.xH₂O have been obtained, wherein $MO = (NH_{\star})_2O$, BaO, and $K_{\star}O$, and y = 2, 3, 4, and 7 (v. K. a. F., B. 24, 2327; 25, 1966; K., Zeit. f. anorg. Chemie, 1,435; Sprenger, l.c.; Gibbs, l.c.).

II. An acid of the form $P_2O_3.21WO_3.xH_2O$, and salts $yMO.P_2O_3.21WO_3.xH_2O$, where $MO = (NH_4)_2O$, K_2O , and Ag_2O and y = 3, have been isolated (K. a. F., l.c.; K., l.c.).

III. The acid P2O5.20WO3.xH2O and a salt $6\text{BaO.P}_2\text{O}_3.20\text{WO}_3.x\text{H}_2\text{O}$ we Péchard (l.c.) and Gibbs (l.c.). were obtained

IV. For the acid P2Os.12WOs.xH2O and the salts yMO.P.O₅.12WO₃.xH₂O, where MO = (NH₁)₂O, BaO, CaO, CuO, PbO, Li₂O, MgO, Hg₂O, K₂O, Ag₂O, Na₂O, Tl₂O, and ZnO, v. Pechard (C. R. 110, 754).

of few salts the yMO.P₂O₅.7WO₃.xH₂O yMO. P_2 O_a.7WO_a.xH₂O have been obtained (MO = (NH₄)₂O, BaO, CaO, Hg₂O, Na₂O; ybeen doubtful) (Gibbs, l.c.; K., l.c.).

VI. Compounds of alkali phosphates with WO, and with phosphoric acids and WO, have been isolated; also compounds of alkalis with H₂PO₂ and WO₂, and with H₂PO₂ and WO₃, are described by Gibbs (Am. 5, 861; 7, 818,

PLATINITUNGSTATES, v. this vol., p. 283.

Silicotunestic acros and salts. Three compounds of SiO, with WO, and $\mathbf{H}_2\mathbf{O}$, which react as acids, have been isolated, and many compounds of bases with SiO2 and WO2 have been obtained (Marignac, A. Ch. [4] 3, 5).

Silicoduodecitungstic acid. and SiO_12WO_xH_O; as most of the salts contain 4M_O (or 2MO) the acid is usually written 4H_O.SiO_12WO_xH_O (=H_8SiW_{12}O_{42}xH_2O). According to Drechsel (B. 20, 1452), the acid is most easily obtained by dissolving Na_WO_4 in boiling water, nearly neutralising with HNO, Aq, dissolving the crystals that separate in cold water, boiling this solution with gelatinous SiO_{2.x}H₂O until HClAq ceases to give a pp., filtering, evaporating, adding a large excess of H2SO, Aq, extracting with ether, separating the lowest layer and evaporating (for details v. D., Large, colourless, dimetric octahedra (with x=29); loses $25H_2O$ at 100° , and all H_2O at 850°. Very soluble in water and alkali solutions. Most of the salts of this acid have the composition 4M₂O.SiO₂12WO₃.xH₂O, or 2MO.SiO₂.12WO₃.xH₂O; M₂O = (NH₃)₂O, Hg₂O, K₂O, Ag₂O, Na₂O; MO = BaO, CaO, MgO; xvaries from 7 to 28 (Marignac, I.c.).

Tungstosilicic acid, and salts. This name is generally given to an acid containing SiO2 and WO, in the same ratio (1:12) as silicoduodecitungstic acid. The formula of the acid is generally written 4H2O.SiO2.12WO3.20H2O. The acid is obtained by evaporating a solution of silicodecitungstic acid to dryness, taking up with water, filtering from separated SiO2, evaporating to a syrup, and allowing to crystallise. Forms short, white, triclinic prisms; melts under 100°, giving off 16H2O and again solidifying. The salts of this acid are obtained by dissolving carbonates in a solution of the acid, and evaporating; they are more soluble and less easily crystallised than the salts of the duodeci-acid: the salts are of the forms

(1) 2MO(4M₂O).SiO₂.12WO₈.xH₂O, where

M = Ca, or $M_0 = K_0$;

(2) $2M_2O.2H_2O.SiO_2.12WO_3.xH_2O$, where $M_2 = K_2$ or Na_2 ; some more complex salts have also been isolated (Marignac, l.c.).

Silico-decitungstic acid, and selts. The acid $4H_2O.SiO_2.10WO_3.xH_2O$ (x probably = 3) is obtained by decomposing the Ag salt by HClAq, filtering, and evaporating in vacuo; the Ag salt is prepared by adding AgNO, Aq to the NH4 salt which is obtained by boiling an acid tungstate of NH, with SiO₂.xH₂O, filtering, and evaporating. The acid forms a clear, yellowish, glassy solid, which does not wholly dissolve in water; loses 3H₂O at 100°; on boiling with water most of the SiO2 separates, and the filtrate contains tungstosilicic acid. A few salts have been obtained; they are difficult to purify from silicotungstates and tungstosilicates (Marignac, Lc.).

STANNIPHOSPHOTUNGSTATES. Gibbs (Am. 7, 892) obtained the salt 2(NH₄)₂O.P₂O₃.2SnO₃.22WO₃.15E₂O by the reaction of SnOl₄.2NH₄Cl and 2Na,O.P.O..24WO.xH2O.

TITANOTUNGSTIC ACIDS. Two acids, 4H2O.TiO2.12WO8.xH2O and

4H₂O.TiO₂.10WO₃.xH₂O₄, are described by Lecarne (v. Klein, Bl. [2] 36, 17).

Vanadotunes c acids and salts. Several compounds of WD, with V2O, and H2O have been described by Gibbs (4m. 4, 377; 5, 361, 391) and Rosenheim (A. 251, 197); they belong to the forms V2O, 10WO, xH2O, $V_2O_3.18WO_3.xH_2O$, and $V_2O_3.4WO_3.xH_2O$. Compounds of these with yM_2O and yMO are also described; and a great variety of compounds coming under the general formula $mMO.nV_2O_1.pWO_3.xH_2O$ have been obtained (cf. Friedheim, B. 23, 1505; and Rothenbach, B. 23, 3050).

Tungsten, oxybromides of. Two oxybromides, WO Br and WOBr, are formed by the reaction of Br on WO₂, and in other ways. The formula WOBr, is very probably molecular, from the

analogy of WOCl.

Tungsten dioxydibromide WO2Br2. tained by reating WO2 in a stream of Br vapour, also by passing vapour of WBr, over heated WO, (Roscoe, C. N. 25, 73); also by heating WS, or e mixture of WO, and C in Br (Borck, J. pr. 54, 254). Red, transparent, prismatic crystals; appearing black when hot, and giving a yellow powder. Volatilised at red heat with partial decomposition to WO3 and WOBr4.

TUNGSTEN OXYTETRABROMIDE WOB,. Obtained by passing Br vapour over a heated mixture of equal parts of WO, and W, and cautiously distilling from less volatile WO₂Br₂ (Roscoe, C. N. 25, 73). Also formed by passing Br mixed with some Ooverheated W, or by heating a mixture of WO, and C in Br vapour (Blomstrand, J. pr. 82, 430). Brownish black, lustrous, crystalline needles; melts at 277° and boils at 327°. Heated in moist air gives WO, and HBr, and is decomposed to the same products by water.

Tungsten, oxychlorides of. Two oxy-

chlorides, WO2Cl2 and WOCl4, are formed by heating W in Cl mixed with some O, and in other

reactions.

TUNGSTEN DIOXYDICHLORIDE WO2Cl2. tained by heating WO₂ in a stream of Cl, and subliming (Roscoe, C. N. 25, 63); also by passing CO2 over a mixture of WO, and CaCl2 heated to redness (Schultze, J. pr. [2] 21, 439). Also formed, along with WOOl,, by heating W in Cl mixed with O; by heating WO₃ and C in Cl; by heating WO₂ in CCl₄ (Watts a. Bell, C. J. 38, 442); and by heating WO₂ with PCl₂ (Schiff, A. 197, 185; cf. Tungsten trioxide, Reactions, No. 10, p. 801). Yellow, four-sided tablets; sublimes without melting at c. 266°, with partial world. decomposition to WO, and WOCl. Not decomposed in moist air, nor by cold water (R., l.c.). Heated in dry NH, gives off NH, Cl and leaves WO, (Rideal, C. J. 55, 43); in this reaction WO, Cl, behaves like CrO, Cl, (R., C. J. 49, 367). According to Smith a. Shinn (Zeit. f. anorg. Chemie, 4, 881), a black compound W.O.N.H. is

formed by heating WO₂Cl₂ in dry NH₃.

TUNGSTEN OXYTETRACHLORIDE WOCL. Mol.

w. 341-04. Formed by heating W in Cl containing a little O; also by passing Cl over a hot mixture of WC, and C; by heating WO, in CCl. (Watts a. Bell, C. J. 33, 442); by heating WCl, or WCl, in O and Cl; by heating WCl, with H₂C₂O₄; and by heating WO₂Ol₂. Prepared by passing vapour of WOl₆ over heated WO₂ in a current of Cl (Roscoe, C. N. 25, 63). Also by heating WO, and PCl, in the ratio WO,:PCl, or WO_s:2PCl_s, distilling off POCl_s and the small quantities of WCl, and WCl, that are formed, washing the reddish residue with a little cold CS, (to remove WCl,), then dissolving in a larger quantity of warm CS2, evaporating, and carefully heating the residue in a stream of CO2, whereby the more volatile WOCl, is separated from WO₂Cl₂ (Schiff, A. 197, 185).

WOCL forms ruby-red, transparent needles; melts at 210.4°, and volatilises at 227.5°, forming a yellow vapour (R., l.c.). V.D. 170.2 to 175.8 at 350°, 171.5 at 440° (R., l.c.; of. Debray, C. R. 60, 820). Exposure to air produces a crust of yellow WO₂Cl₂. Heated in NH₃ probably gives W₂N₃ (Rideal, C. J. 55, 43). Decomposed by water, with a hissing sound, to WO, and HClAq

(Roscoe, l.c.).

Tungsten, exyfluorides of. No compound of W with O and F has been isolated, but compounds are known which may be regarded as containing WO.F₂. These compounds, WO.F₂.2MF and WO.F₂.MF, and also a compound WO₃F₂.2KF, are described under Tungsr-

OXYFLUORIDES (p. 799).

Tungsten, phosphides of. By heating W in vapour of P, a dark-grey powder was obtained by Wöhler a. Wright (A. 79, 244), to which they gave the composition W₃P₄. By heating a mixture of WO₃ and P₂O₅ in a graphite crucible, W. a. W. (l.c.) obtained large, dark-grey, lustrous, six-sided prisms; S.G. 5.207; insoluble in acids, including aqua regia, soluble in a molten mixture of Na₂CO, and NaNO,; unchanged by heating in air, but burns brilliantly in O. From an estimation of the P, W. a. W. gave the formula W₂P.

Tungsten, salts of. No salts have been isolated by replacing the H of oxyacids by W. Except the halides, the only salts of W that are known are those wherein W forms part of the

negative radicle.

Tungsten, selenides of. By saturating a solution of Na, W,O, with H,Se, and then adding dilute H₂SO₄Aq, Uelsmann (A. 116, 125) obtained a black pp., to which he gave the formula WSe, (one estimation of W is given); by heating this in a tube a grey solid was obtained, said by U. to be WSe2 (no analyses). WSe2 is said to be easily soluble in solutions of alkalis, alkali sulphides, or alkali selenides.

Tungsten, sulphides of. Two compounds of W and S are known, WS2 and WS3; the latter is acidic, forming salts M2WS. The V.D. of

neither has been determined.

Tungsten disulphide WS2. Obtained by heating WO, to whiteness in vapour of S or in H2S (Berzelius); also by heating W and S; also by heating WO, with 6 pts. HgS out of contact with air (Borck, J. pr. 54, 254). According to Carnot (Bt. [2] 32, 164), WS₂ is usually formed when a compound of W is heated in a stream of dry H₂S (v. also von Uslar, A. 94, 256; Corleis, A. 232, 262). A dark-grey, graphite-like, lustrous, crystalline powder (v. Riche, A. Ch. [8] 50, 26). Is said to decompose steam at a full red heat; reduced to W by very long sontinued heating TUNGSTEN TRISULPHIDE WS. Obtained by tusing powdered wolframite with 2 pts. C, 3 pts. S, and 3 pts. Na₂CO₃, dissolving in water, decomposing the Na, WS, Aq thus produced by HClAq, washing the pp. completely out of contact with O, and drying at 100° (von Uslar, A. 94, 256; Corleis, A. 232, 264). A black powder; slightly soluble in water, more soluble on boiling, probably with partial decomposition. Soluble in alkali sulphide solutions, forming thio-salts (v. TUNGSTEN, THIO- ACIDS AND SALTS OF, infra). By adding to Na₂WS₄Aq a little more HClAq than was needed to combine with the Na, Winssinger (Bl. [2] 49, 452) obtained a colloidal form of WS.

Tungsten, sulpho- acids and salts of, v. next article.

Tungsten, thio-acids and salts of. Rv saturating (NH₄)₂WO₄Aq and K₂WO₄Aq with H.S. Berzelius obtained (NH₄)₂WS, and K₂WS, (P. 8, 267); these salts were more fully examined by Corleis (A. 232, 258), who also obtained Na, WS,, and also some thio-oxy-tungstates. All the thio-tungstates that have been isolated are ortho- salts, i.e. salts of H2WS4. Very dilute solutions of thiotungstates can be titrated with IAq in presence of KHCO, (C., l.c.).

Ammonium thiotungstate (NH₄)₂WS₄. Obtained by passing H2S for four or five hours into a solution of 10 g. WO_s.H₂O in 100 c.c. NH₃Aq S.G. 94, +20 c.c. water, allowing to stand for some hours in a closed vessel, and washing the crystals that separate with alcohol and ether. Forms orange-yellow prisms, isomorphous with (NH₄)₂MoS₄; very easily decomposed in moist air; easily soluble in water, slightly soluble in alcohol. Heated in CO2 gives WS2 (B., L.c.;

C., l.c.).

Ammonium dithio-oxy-tungstate

(NH₄)₂WS₂O₂. Obtained by passing H₂S into a solution of 10 g. WO₂.H₂O in 40 c.c. NH₃Aq S.G. 90, +10 c.c. water, until the liquid becomes turbid, and washing the crystals that form with alcohol and ether. Forms yellow, prismatic crystals. When dry, the salt is unchanged in air. Decomposed by re-crystallising from water, giving paratungstate (NH₄)₆W₇O₂₄.6H₂O (C., l.c.).

Potassium thiotungstate K. WS. Obtained by warming (NH₄)₂WS₄ with KHSAq, adding alcohol, and crystallising from conc. KHSAq. Forms yellow, prismatic needles; easily soluble in water. By continued boiling with KHSAq paratungstate is formed, K₆W₁O₂₁.6H₂O (C., l.c.). Forms a double salt with KNO₃, with the composition K, WS, KNO, (B., Lc.).

Potassium trithio-oxy-tungstate

K.WS.O.H.O. Obtained in citron-yellow, quadratic tablets, by passing H.S for three or four hours into a solution of 10 g. K2WO4 in 10 c.c. water, and evaporating in vacuo, or precipitating by alcohol (C., l.c.).

Potassium monothio-oxy-tungstat.

K, WSO, H,O. Obtained, in almost colourless, very hygroscopic, crystalline masses, by passing H.S into a solution of 10 g. K.WO, in 5 c.c. water until the liquid is turbid, filtering, adding 4 to 5 vols. alcohol, separating the under layer of liquid, allowing to crystallise, and washing with alcohol and ether (C., l.c.).

Sodium thiotungstate Na WB. Obtained,

but not pure, by Corleis (l.c.), by decomposing (NH₄)₂WS, by NaHSAq.

Tungsten, thiochloride of. By heating W to redness with S₂Cl₂, Smith a Oberholtzer (Zeit. f. anorg. Chemie, 5, 63) obtained a red, crystalline sublimate, unstable in air, probably W.S.Cl_s. M. M. P. M.

TUNGSTEN BRONZES v. under Tungsten OXYACIDS, p. 806.

TUNGSTIC ACIDS v. Tungsten oxyacids, p. 802; also Tungsten thio-Acids, supra.

TUNGSTOXYFLUORIDES v. under Tung-

STEN FLUORIDES, p. 799.
TUNGSTEN, ORGANIC COMPOUNDS OF. Tungsten, heated with MeI at 240°, forms WMe,I, [110°], which crystallises in tables, and is converted by Ag₂O into WMe₂O, which dissolves in acids (Riche, C. R. 42, 203; Uahours, A. 122, 70).

TUNICIN v. Animal cellulose, vol. i. p. 718. TURMERIC. The root of Amomum Curcuma. It contains cureumin (q. v.) and turmerol.

TURMERIC ACID C11H14O. [35°]. A product of oxidation of turmerol by cold KMnO4 (Jackson a. Menke, Am. 6, 77). Needles, sl. sol. water, v. sol. alcohol.—CaA', Baq. S. (of CaA', 1-28 at 16°. White needles.—AgA'.

TURMEROL C₁₉H₂₈O (?). (193°-198° at 60 mm.). S.G. 17.902. $[a]_D = 33.5^{\circ}$. An oil occurring in turmeric. Distils with decomposition at 285°-290° (Jackson a. Menke, Am. 4, 368; 6, 77). Dextrorotatory. Does not combine with NaHSO₃. KMnO₄ oxidises it to acetic, terephthalic, turmeric, and apoturmeric acids. PCl, forms C1, H2, Cl, which is also got by heating turmerol with conc. HClAq at 150°. Sodium forms C19H27ONa, whence isobutyl iodide forms oily C19H27OC4H9.

Apoturmeric soid C.H.(CO.H), (?). [221].

Woolly mass, sol. boiling water.

TURPENTINE. Semi-fluid resins exuding from coniferous trees. They consist of resin dissolved in oil of turpentine. On distillation oil of turpentine passes over and colophony remains behind. The various oils of turpentine are described under TERPENES. The chief constituent of colophony is abietic anhydride (v. ABIETIO ACID and SYLVIO ACID). On oxidation by dilute HNO, colophony yields isophthalic, trimellitic, and terebic acids (Schreder, A. 172, 93). On the products of distillation of colophony v. Resins

TURPETHIN C. Had O. [c. 183]. Occurs in the root of Convolvulus Turpethum (Ipomæa Turpethum) (Boutron-Charlard, J. Ph. 8, 131; Spirgatis, J. pr. 92, 97; A. 139, 41). Purgative yellowish-brown resin, v. sol. alcohol, insol. ether (difference from jalapin). Conc. H₂SO₄ forms a red solution. Boiling alkalis convert is into turpethic acid. Boiling dilute mineral acids yield glucose (3 mols.) and turpetholic acid (1 mol.).

Turpethic acid C_{s.}H_{so}O_{1s}. Amorphous yellowish mass, v. sol. water. Split up by boiling HClAq into glucose and turpetholic acid.—BaA". BaH_AA", (dried at 100°).

acid.—BaA".- BaH,A", (dried at 100°).

Turpetholic acid C_{1.}H₃₂O₄. [c. 88°]. Minute needles (from dilute alcohol), al. sol. ether.— NaA' (dried at 100).—BaA's. Amorphous.—AgA'. Amorphous pp. A'. Amorphous pp. Ethyl ether EtA'. [73°]. Plates.

asthmatica (Hooper, Ph. [3] 21, 617). Crystalline, sl. sol. water, sol. alcohol and ether.

TYPE METAL. An alloy of 15-20 parts Sb with c. 70 parts Pl and 10 to 15 parts Sn; v.

LEAD ALLOYS, vol. iii. p. 124.

TYPES. The object of classification is to put together like things, and to put apart things that are unlike. A perfect system of chemical classification would place side by side those elements and compounds that are chemically similar, and it would also indicate the relations that exist between all the elements and all the compounds. Inasmuch as the object of chemistry is the study of the connections between composition and properties, and between changes of composition and changes of properties, a complete scheme of chemical classification must indicate the relations of the substances classified, both as regards composition and also as regards properties. To connect composition with properties necessitates a thorough knowledge of both, and this knowledge can be gained only by comparing one substance with other substances. But, because of the large amount of detailed investigation that is required before the chemical properties of an element or a compound are known, chemists have often forgotten the larger issues of their labours, and have busied themselves rather with the examination of individual bodies than with the comparative study of many bodies. And, because of the imperfection of chemical knowledge at any time, those who have attempted the classification of chemical substances have generally paid chief attention either to the composition or to the properties of the substances to be classified. Hence systems of classification have sometimes prevailed that were founded chiefly on similarities of composition, and at other times classificatory schemes have been in vogue that rested mainly on similarities of properties. But there has always been a desire, and generally an effort, to classify on the bases of composition and properties. To classify satisfactorily demands the recognition of a simple class-mark, which shall also be clear, definable, and invariably applicable. It is probably correct to say that none of the larger classes of chemical compounds has a class-mark of this description. Hence the main difficulty in chemical classification. Take, for instance, the great class of acids. What is the What must be known about the class-mark? composition of a compound before it is put into the class of acids? There is no single and sufficient class-mark to be gained by studying the compositions of acids. What, then, about the properties of acids? Here, too, no simple, definable, and applicable property has been found which serves to distinguish acids from all other compounds. Acids, it is usually said, are compounds containing replaceable hydrogen. But the expression replaceable hydrogen has not been, and cannot be, exactly defined. Hence, all that can be done is to set up an ideal or typical scid, and to place in the class acids those compounds which fairly closely approach this type as regards both properties and composition. And as with acids so with other classes of compounds. A classification by means of types becomes inevitable in such a science as chemistry,

TYLOPHORIN 3. An alkaloid in Tylophora! but a typical classification cannot be final in any exact science.

> Lavoisier's system of classification rested on oxygen; compounds were regarded as formed by the union of oxygen with other elements or groups of elements; the non-oxygenated part of a compound was called by Lavoisier the rest or radicle; compounds of oxygen with certain radicles were bases, compounds with other radicles were acids, and salts were formed by the union of bases with acids. Lavoisier used the term radicle to include elements and groups of elements.

> 'J'ai déjà fait observer, que dans le règne minéral presque tous les radicaux oxidables et acidifiables etaient simples; que dans le règne végétal au contraire, et surtout dans le règne animal, il n'en existait presque pas qui ne fussent composés au moins de deux substances, d'hydrogène et de carbone; que souvent l'azote et le phosphore s'y réunissaient, et qu'il en résultait des radiceux à quatre bases.'—Tratté élémentaire de Chimie, [1793], p. 251.

> About twenty-four years after Lavoisier made this statement Berzelius repeated it.

> 'Nachdem wir den Unterschied zwischen den Producten der organischen und der unorganischen Natur, und die verschiedene Art und Weise wie ihre entfernteren Bestandtheile untereinander verbunden sind, naher kennen gelernt, haben wir gefunden, dass dieser Unterschied eigentlich darin besteht, dass in der unorganischen Natur alle oxydirten Körper ein einfaches Radical haben, während dagegen alle organischen Substanzen aus Oxyden mit zusammengesetztem Radical bestehen. —Lehrbuch der Chemie, 2 Aufl. [Stockholm, 1817], vol. i. p. 544.

> Why do acids and bases unite to form salts? What is the cause of the formation of compounds by the union of radicles, either simple or compound? These questions were answered by Berzelius by appealing to the fact that the electric current very often resolves compounds into two parts, and in many cases also effects the combination of elements or groups of elements. Radicles combine, said Berzelius, because the negative electricity on one is thereby neutralised by the positive electricity on the other. Thus arose the electro-chemical conception of dualism (v. Dualism, vol. ii. p. 415; cf. RADICLE, vol. iv. p. 393).

> In 1832 Liebig and Wöhler made an exhaustive study of compounds obtained from bitteralmond oil; they showed that the relations of composition and properties of these compounds were brought together, and expressed in a consistent conception by supposing that all the compounds contained a radicle having the composition $C_1H_2O(C=12, O=16)$; this group of elements, common to all the derivatives of bitteralmond oil, they called benzoyl. The work of Liebig and Wöhler marked a great advance in chemical classification; it was the actual working out of the connections between composition and properties of a number of compounds, and the expression of these connections in clear and definite language. The term radicle became henceforth the expression of a vivifying concep-The compounds derived from bitteralmond oil had certain common properties, and they had also a common composition; they belonged to the same type. The compounds of benzoyl examined by Liebig and Wöhler included such compounds as these: C,H,O.OH, C,H,O.H, C,H,O.Cl, C,H,O.CN, C,H,O.NO. The radicle C.H.O is common to all; this radicle is united with H, Cl, NO, CN, or some other radicle, in the different compounds; neverthe-

less, the properties of the compounds are so similar that all are said to belong to one type. To what extent, then, it was asked, may the properties of one radicle differ from those of another before the replacement of one of these by the other carries with it a change of type? answer generally given to this question was: If the replacing radicle is chemically similar to the radicle replaced the type will not be destroyed. In 1834 Dumas, from the study of the action of chlorine on various organic compounds, announced the *empirical laws* of substitution as follows: (1) When a compound containing hydrogen is subjected to the dehydrogenating action of chlorine, bromine, iodine, oxygen, &c., one atom of chlorine, bromine, iodine, or oxygen, is taken up for each atom of hydrogen lost by the compound. (2) The same rule applies, without modifications, when the compound contains oxygen. (8) If a hydrogenised compound contains water the hydrogen of the water is given up without replacement, and then another quantity of hydrogen is absorbed, so that the final result is a replacement of hydrogen. A year or so later Laurent propounded a theory of substitution, which asserted that when equivalent substitution of hydrogen by chlorine or bromine occurs the chlorine or bromine takes the place of the hydrogen, and to a certain extent plays the same part as the hydrogen, and hence the chlorinated or brominated product must be similar to the compound from which it has been prepared. Laurent's views were much opposed, for a time by Dumas himself; but they gradually prevailed, and chemists became familiarised with the notion of the replacement of such a decidedly electro-positive element as hydrogen by an element so decidedly electro-negative as chlorine, resulting in the production of a compound of the same chemical type as the original substance.

About 1839 Dumas sought to distinguish mechanical types from chemical types. Compounds formed one from the other by the replacement of a certain number of equivalents of one radicle by the same number of equivalents of another radicle, and having their radicles similarly combined, and exhibiting similar properties, were said to belong to the same chemical type. Compounds were said to belong to the same mechanical type when they were composed of equal numbers of equivalents of radicles, but differed essentially in their properties. Dumas at this time regarded the properties of compounds as conditioned more by the arrangement than by the nature of their parts. He compared compounds to planetary systems, the planets being represented by the atoms of the compounds. One atom, he said, might be replaced by another atom, or an atom by a group of atoms, without destroying the system: if the number of replacing atoms, or radicles, were the same as the number of atoms, or radicles, replaced, and the relative arrangement of all the atoms or radicles were not altered, the compounds belonged to the same type. By some such development as this the theory of types came to include the older theory of radicles.

It is evident that the terms used by Dumas and others to express the conceptions of the theory of types are vague and incapable of

exact definition. The expression 'equivalent radicles' cannot be defined, nor can an exact and invariable connotation ! e given to the expressions 'radicles similar'y combined' and compounds having similar properties.' But if it had been possible to classify chemical substances in an ideally perfect way the theory of types would not have arisen. If there is to be a typical classification the language whereby that classification is expressed must be more or less vague. Whether a compound formed from another by replacing equivalents of one kind by an equal number of equivalents of another kind does or does not belong to the same type as the parent compound can be *determined only by a careful study of the properties of both compounds and by a comparison of the compounds one with another. The application of the theory of types was possible only when the properties and the compositions of compounds were exhaustively compared. The theory of types produced much fruit, because it suggested and demanded much inquiry into the fundamental problem of chemistry.

In 1849 Wurtz prepared two compounds which resembled ammonia in their prominent characters. These compounds were found to have the compositions C2H3N and C4H3N respectively (C=6). The similarities between the properties of these compounds and ammonialed Wurtz to regard them as substituted ammonias, and to express this conception by the formulæ C₂H₃.NH₂ and C₄H₅.NH₂. One method by which these compounds were prepared consisted in treating methyl and ethyl bromides (C2H3Br and C,H,Br; C=6) with ammonia, and then decomposing the products by potash. These reactions suggested the view that the new compounds were derived from methylic and ethylic ethers (C₂H₃O and C₄H₅O, according to the notation then used) by substituting NH₂ for O. Both views agreed in representing the new compounds as C₂H₃.NH₂ and C₄H₅.NH₂ respectively. But the properties of the compounds resembled those of ammonia; hence the view that Wurtz's compounds were derived from ammonia prevailed. This view was expressed by saying that these compounds belonged to the ammonia type, and by comparing the formula of the typical compound with the formulæ of the compounds formed after that exemplar; thus:

Type: Derivatives:

Ammonia NH₂.H. Methylamine NH₂.C₂H₃.

Ethylamine NH₂.C₄H₅.

Very soon after Wurtz's preparation of methylamine and ethylamine Hofmann argued that it should be possible to substitute each of the three atoms of hydrogen in NH, by the radicles C_xH_x and C_xH_y , and so to obtain the compounds $NH(C_2H_3)_2$, $N(C_2H_3)_3$, $NH(C_4H_5)_2$, and $N(C_4H_5)_3$ (C=6), all which compounds ought to resemble ammonia in their properties. Hofmann fulfilled his own prophecy by preparing diam tri-methylamine and diam tri-ethylamine; and, basing his methods on the conception of the ammonia type, he prepared a large series of ammonia-like compounds derived from NH₃, 2NH₃, &c., by substituting for H various radicles composed of carbon and hydrogen. The ammonia type was thus established.

About the year 1850 Williamson studied the about the year 1850 Williamson studied the relations of ether to alcohol. The formula of ether was then wri ten C_1H_5O (C=6, O=8), and that of alcohol $C_1H_5O_2$. Williamson heated potassium alcohol te $(C_1H_5KO_2)$ with ethylic iodide (C_1H_2I) , expecting to obtain ethylated alcohol $(C_4H_5(C_4H_5)O_2)$; but he obtained common ether. Williamson suggested that the formula then accepted for ether should be doubled and then accepted for ether should be doubled, and written $(C_4H_5)_2O_2$. He also found that by heating common alcohol with sulphuric acid, and adding methylic alcohol to the hot mixture, an etherlike compound was obtained, the simplest formula of which was C_3H_4O (C = 6, O = 8). Williamson argued that if common ether is C.H.O. then the ether of methylic alcohol must be C2HO, and that, in the reaction of sulphuric acid with ethylic and methylic alcohols simultaneously, a mixture of these two ethers might be expected to be produced. To account for the facts he had observed Williamson proposed to double the formulæ of the two ethers and to write them (C₄H₅)₂O₂ and (C₂H₃)₂O₂; and for the formula of the ether formed by the action of sulphuric acid on the two alcohols simultaneously be proposed the formula C4H5(C2H2)O2 $[=2C_xH_xO].$ Williamson compared the relations between the alcohols and ethers with the relations between water, potash, and potassium oxide. These relations are made clearer if the formulæ are expressed in terms of the atomic weights C = 12 and O = 16; thus:

Type: Derivatives:

Water H.H.O Alcohol C₂H₃,H.O Ether C₂H₃,C₂H₃,O Potash K.H.O Potassium oxide K.K.O.

Thus arose the water type. To this were referred such compounds as acetic acid C₂H₃O.H.O, anhydrous acetic acid C₂H₃O.C₂H₃O.O, and many others.

Gradually the greater number of compounds, both organic and inorganic, came to be referred to four fundamental types—the hydrochtoric acid type HCl, the water type HHO, the ammonia type NHHH, and the marsh gas type CHHHH. To these were added so-called condensed types and mixed types; sulphuric acid, for instance, SO_x OH.OH, was said to belong to the double water type.

Oxamic acid CO.NH₂.CO.OH was regarded by Wurtz as derived from the water-ammonia type.

Thus. $\begin{array}{c}
H \\ H \\ O \\
H \\
N
\end{array}$ gives C_2O_2 H N

The great danger attending the development of the theory of types was, that careful investigations into the reactions of compounds might easily be abandoned in favour of a superficial examination of a compound, followed by a relegation of it to this or that type. The theory began by being very vague and elastic; the adoption of a few definite types rendered it more exact; but the temptation to manipulate for-

mulæ on paper, and to invent condensed types and mixed types, gradually led chemists to see that the theory had done its work, and that it must give place to wider views, which should also be more accurate, regarding the connections between composition and properties.

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The conception of types carried with it, from the first, more or less vague notions regarding the arrangements of the atoms in those collocations of atoms that form the reacting units of compounds; at any rate, this conception was bound up with the general conception of atomic arrangement. The reacting units of compounds that showed similar properties were thought of as composed of atoms similarly arranged. Moreover, the type was said to be maintained when an atom, or a group of atoms, in the parent compound was replaced by an equivalent number of atoms, or groups of atoms. These two conceptions, that of equivalency, and that of atomic arrangement, led gradually to the representation of the reacting unit, or molecule, of a compound as a definitely arranged group of atoms held together by actions and reactions between individual atoms; these conceptions led to the wider conception of valency, and to a system of classification based on the notion that each kind of atom is capable of directly combining with a limited, and determinable, number of other atoms. The formula given to chlorosulphonic acid, with the object of connecting the composition with the properties of this compound, and of suggesting relations between this and other compounds, may be taken as an example of the merging of the notion of types into that of valency. This acid was regarded as derived both from water and hydrochloric acid; it was supposed to belong to the mixed water-hydrochloric acid type. Hence the formula, CISO,H, was written thus :-

The radicle SO₂ was regarded as equivalent to 2H; and it was said that the residues of the two parts of the mixed type, i.e. Cl and OH, were bound together by the radicle SO₂. Again, monochlorhydrin, C₃H,ClO₂, was regarded as a derivative of the triple type HHO.HCl.HHO, and was said to be formed by the binding together of the three residues OH, Cl, OH by the radicle C₂H₂ regarded as equivalent to 3H. Thus:—

$$\begin{array}{cccc} Type & HO & Monochlorhydrin & HO \\ H & & & Cl \\ HCl & & HO \\ \end{array} \right\} C_2H_6$$

The moment the conceptions underlying these formulæ, and formulæ like these, were expressed in the language of atoms and molecules, the notion of atomic valencies was gained. The group of atoms SO₂ replaces two atoms of hydrogen, one atom in the molecule HCl and one atom in the molecule HHO; and as the atomic group SO₂ is capable of directly combining with two atoms of hydrogen, or such a number of atoms as is equivalent to two atoms of hydrogen, this group binds together the atom Cl and the atomic group OH, and so the new molecule

SO. OH.Cl is produced. This statement contains the fundamental conception of atomic valency; it also contains the conception of types, and that of radicles.

The radicles were compared as regards their equivalency, generally in terms of hydrogen; then the elementary atoms were compared, and arranged in classes, such that all in one class were exchangeable, any number of one kind for the same number of another kind.

When the atoms had been classified in accordance with the number of atoms of hydrogen, or atoms equivalent thereto, with which each was capable of combining, the foundations had been laid of a system of classification which was more accurate than that based on the notion of types, and which at the same time included the essential characters of the typical system. article CLASSIFICATION in the first edition of this Dictionary represents the stage which had then (1863) been reached in the process of fusion of the theories of radicles, types, and valency. The system of classification developed in that article is based (1) on the compositions of compounds interpreted by help of the theory of valency, and (2) on the properties of the compounds interpreted by the help of the theory of types; the study of the properties of a compound is used to determine the radicles that the compound contains, and a knowledge of the valencies of these radicles determines the form of the compound, and, taken along with the reactions of the compound, the type to which it is to be referred.

Although the introduction of the principle of atomic equivalency widened the conception of chemical types, it also tended towards a method of classification which was based on too slight a study of the bodies to be classified. A superficial examination of a compound generally sufficed to bring out some similarities between it and a well-known typical substance; the new compound was at once referred to its type; the form of the type determined the form of the compound under examination; it was then only necessary to manipulate the empirical formula by arranging the elements in groups, or radicles, and to assign to each radicle such a valency as satisfied the general rules that had been deduced from the study of a few typical compounds. The conception of types became very mechanical in its applications, so mechanical indeed that it was in danger of becoming metaphysical. New radicles were easily invented, and their valencies were deduced by a priori arguments. Kekulé recalled chemists to the study of properties by insisting that the properties of a compound are conditioned by the properties of the atoms which compose the molecule of the compound; and at a later date Mendeléeff made the same demand by asserting that the properties and the compositions of all compounds are periodic functions of the atomic weights of the elements.

The notion of types plays an important part in the classification of elements an' compounds that has arisen from the practical examination of the meaning of Mendeléeff's periodic law. The periodic classification of the elements lays considerable stress on the study of the typical oxides, hydrides, hydroxides, &c., of each group of elements; for instance, the highest salt-forming oxide of Group I. belongs to the type R₂O,

that of Group II. to the type B,O, and so on. But the word type is not us I here with quite the same connotation as was given to it by the theory of types. All that i implied in statements such as those just nade is, that the elements of Group I. combin, with oxygen in the ratio of two atoms of element to a single atom of oxygen, that these oxides are salt-forming, and that they are characteristic of the elements The typical oxide-form of each of this group. group expresses the composition of that oxide which contains the greatest number atoms of oxygen relatively to one atom of the group-element, and which oxide reacts either with acids or with alkalis to form corresponding salts. The oxides that belong to the typical oxide-form of a group may be acidic or basic, or some of them may be acidic and some basic; all that is asserted of their properties is that they are salt-forming. The properties of the typical oxide of any individual element are conditioned by the general character of the group, and the general character of the series, wherein the element is placed; by the special character of the element itself; and by the position of the element in the whole periodic system of classification. The term type is used in the nomenclature arising from the comparative study of the elements and compounds based on the periodic law with a wider, and at the same time a more exact, meaning than that which was given to it when the notion of types was made the basis of chemical classification. In the older classification the conception of types was the basis of the system; the conception was of necessity vague, and hence it was necessary sometimes to widen, and sometimes to narrow, the applica-The modern system tion of the conception. of classification is based on the relative weights of the atoms of the elements, and the conception of types plays a subsidiary part; the form of the typical oxide, or hydroxide, or other compound, is determined, as the compositions of all the compounds are determined, by the relative masses of the atoms of the elements; and the general character of the typical oxide, hydroxide, &c., of each group is determined, as the special character of each compound of each element is determined, also by the relative masses of the atoms of the elements.

The periodic classification of the elements and compounds makes use of the notion of types in another way, by applying the conception to certain elements. The elements placed in series 1 and 2, viz. H, Li, Be, B, C, N, O, and F, exhibit almost the whole range of properties of all the elements. The properties of the succeeding elements may almost be said to be but variations on the theme announced in the change from hydrogen to fluorine. These eight elements are types of all the others. In a somewhat narrower sense the element lithium summarises the range of properties shown in Group I.; beryllium summarises the range of properties shown in Group II.; boron, the properties of Group III.; carbon, the properties of Group IV.; nitrogen, the properties of Group V.; oxygen, the properties of Group VI.; and fluorine, the properties of Group VII. Each of the seven elements is the typical element of its group. Objections have been raised to this use of the term typical, on the ground that each of the elements a question differs more from any member of is group than any other two members of the goup differ from one another. But if an element s to summarise the properties of some ten or el ven other elements, it must differ considerably rom each of these, while at the same time it resembles them all. The question of typical elements, and also that of typical oxide-forms, &c., are discussed in the article Periodic Law (vol. iii. p. 808); that article should be consulted.

The older classification by types prepared the way for the wider conception of valency, which took up and utilised the permanent features of the typical arrangements of elements and compounds. The study of valency led chemists to see the importance of examining the properties of the atoms of the elements, and thus made them rer ly to accept the more elastic, and more exact, generalisation of the periodic law. The theory of types developed into the theory of valency, and the theory of valency has been included in the theory of the periodicity of the connection between the atomic weights of the elements and the compositions and properties of the compounds of the elements. The general conception of types remains; we are still obliged to picture to ourselves a typical acid, a typical base, a typical salt, a typical alcohol, a typical amide, and so on. But this conception is no longer the basis of chemical classification. . We have gone deeper down, and laid the foundations of our system on the firmer basis of the atomic weights of the elements. M. M. P. M.

TYPHOTOXINE C₇H₁₇NO₂. An alkaloid obtained from pure cultures of the typhogen

bacillus (Gautier, Bl. [2] 48, 13).

TYROSINE C.H., NO. i.e.

C.B., (OH). CH2. CH(NH2). CO2H. p-Oxy-a-amidophenyl-propionic acid. Mol. w. 181. [235°].

S.G. 1.456 (Siber, B. 17, 2837). S. 04 at 20°;

65 at 100°. S. (90 p.c. alcohol) 0074 in the cold. H.F. 156,400. H.C.v. 1,070,800. H.C.p.

1,071,200 (Berthelot a. André, Bl. [3] 4, 227).

[a]_b = -8° in HClAq at 16°; -9° in KOHAq at 20° (Mauthner, M. 3, 343).

Occurrence.—Occurs abnormally in the liver and urine (e.g. in cases of poisoning by phosphorus (Frerichs a. Städeler, J. 1855, 729; 1856, 702; Blendermann, H. 6, 242), in the cutaneous scales in pellagra (Schmetzer, Dissert., Erlangen, 1862), in cochineal (De la Rue, A. 64, 35), in sprouting pumpkin seeds (Schulze a. Barbieri, J. pr. [2] 20, 401; 32, 457), in the roots of Stachys tuberifera (Planta, B. 23, 1699), in dahlia bulbs (Leitgeb, C. C. 1888, 1397), in alcoholic extract of woad leaves (Schunck, C. N. 37, 223), and in beetroot molasses (Lippmann, B. 17, 2835).

Fornation.—1. By potash fusion from casein (Liebig, A. 57, 127; 62, 269), globulin, feathers, hairs (Leyer a. Köller, A. 83, 332), and albumen (Nenoki, J. pr. [2] 17, 97).—2. By the action of boiling dilute H.SO, on ox-horn (Hinterberger, A. 71, 72), on fibrin (Städeler, A. 111, 12; 116, 57), and on silk (Weyl, B. 21, 1529).—3. By putrefaction of yeast (A. Müller, J. pr. 57, 162; Béchamp, C. R. 74, 115, 184), albumen and gelatin (Jeanneret, J. pr. [2] 15, 353).—4. In small quantity by heating conglutin with HClAq (Siegfried, B. 24, 419).—5. From p-amido-

phenyl-alanine and HNO₂ (Friedländer a. Mähly, B. 16, 854; Erlenmeyer a. Lipp, A. 219, 161).

Properties.—Stellate groups of slender silky needles (from water), insol. alcohol and ether, m. sol. NH₃Aq and KOHAq. Levorotatory. Its solution is not ppd. by lead acetate or subacetate until NH₄Aq is added. Boiling Hg(NO₂)₂ containing HNO₂ turns its solution red, forming a brownish-red pp. (R. Hoffmann, A. 87, 123; L. Meyer, A. 132, 156). After warming with conc. H₂SO₄, diluting with water, and neutralising with BaCO₂, the solution is turned violet by FeCl₂. Tyrosine gives off less N₂ when its solutions are decomposed by NaOH and Br in presence of

NH, than when the NH, is absent.

Reactions.—1. At 270° it splits up into CO. and CoH, (OH).CH2.CH2.NH2 (Schmitt a. Nasse, A. 133, 211).—2. Potash-fusion forms p-oxybenzoic and acetic acids (Barth, A. 136, 110; 3. KClO₃ and HClAq form tetra-chloro-quinone.—4. By putrefactive fermentation it is converted into hydro-p-coumaric acid, and finally into p-cresol and phenol (Weyl, B. 12, 1450).—5. HIAq and P at 150° from NH₃ and p-oxy-phenyl-propionic acid (Hüfner, Z. [2], 4, 391; 6, 113). Conc. HClAq and HBrAq at 240° have no action.—6. Conc. H₂SO₄ (4 pts.) at 100° forms crystalline C₂H₁₀(SO₃H)NO₃, v. sl. sol. cold water, which also occurs in a hydrated amorphous form (containing 2aq). H yields (NH,)HA" aq, Ba(HA"), 4aq, and Ca(HA"), 5aq (Städeler, A. 116, 57). BaC, H, NO, Saq is obtained from the product of the action of H.SO. (10 pts.) on tyrosine (1 pt.).—7. MeOH, potash, and MeI yield crystalline C₁₃H₁₀NIO₃K, sol. water, decomposed by warm KOHAq into NMe. and the methyl derivative of p-coumaric acid (Körner a. Menozzi, G. 11, 550).-8. Potassium cyanate added to boiling water containing tyrosine forms tyrosine-hydantoic acid C₆H₄(OH).CH₂.CH(NH.CO.NH₂).CO₂H crystallising from water in needles, insol. ether (Jaffé, H. 7, 310). This body begins to melt at 155°, being decomposed above 170°. It gives a red colour and pp. on warming with Millon's reagent. It forms KA'aq, crystallising from alcoholbenzene. -9. Tyrosine administered to a dog appears in the urine as tyrosine-hydantoin $C_0H_4(OH).CH_2.CH < NH.CO (275^\circ-280^\circ]$, which crystallises from water in needles, and is decomposed by heating in sealed tubes with barytawater into CO₂, ammonia and tyrosine. Other products found in the urine of a rabbit after a dose of tyrosine are hydro-p-coumaric and poxy-phenyl-acetic acids and phenols (Blender-

oxy-pienyr-aceuro acute and photosis (2-3-3-4) mann, H. 6, 251).

Salts. — Na₂C₂H₂NO₃. — CaC₂H₂NO₂. — BaC₂H₂NO₃ 2aq. Prisms, more sol. cold than hot water.—Cu(C₂H₁₀NO₃)₂. S. ·08 in the cold; '4 at 10C''. Got by adding Cu(OH)₂ to a boiling solution of tyrosine (Hofmeister, A. 189, 6). Insol. alcoh-1. Deposits black cupric oxide on boiling with water.—C₂H₁₁NO₂(HgO)₂ 2aq.—C₂H₁₁NO₃(HgO)₂ aq. — C₂H₁₁NO₃(HgO)₃ aq (Vintschgau, ·7. 1869, 985).—Ag₂O₄H₂NO₃ aq (Vintschgau, ·7. 1869, 985).—Ag₂O₄H₂NO₃ accomposed by water.—B'HOl 2aq: needles, decomposed by water.—B'HOl₂ (Gintl, Z. [2] 5, 704).—B'H₂SO₄. Slender needles, not coloured by FeCl₂

Di-bromo-tyrosine C, H, Br, NO, 2aq. S. .46 at 16°; 4 at 100°. Formed from dry tyrosine and Br vapour (Gorup-Besanez, A. 125, 281). Needles or tables (from water), sl. sol. alcohol. $Ag_2C_0H_2Br_2NO_3$ 2aq. — B'HCl 1 $\frac{1}{2}$ aq. — B'HBr. — B' $\frac{1}{2}H_2SO_4$. Prisms, sol. water and $B'HBr. - B'_2H_2SO_4$ alcohol.

Nitro-tyrosine C₉H₁₀N₂O₅ i.e. C₉H₁₀(NO₂)NO₃. Formed from tyrosine (1 pt.), water (4 pts.), and nitric acid (4 pts. of S.G. 1-3) in the cold (Strecker, A. 73, 70; Städeler, A. 116, 77). Paleyellow needles, sl. sol. cold water. — Salts: Ba(C,H₀N₂O₅)₂ (dried at 100°). Blood-red amorphous mass. Ag2C9H8N2O3 aq: orange pp. changing to a red powder.—B'HCl aq. Tufts of lemon-yellow needles.-B'HNO.

needles. -B' LSO. Yellow Lemon-yellow needles or granules.

Di-nitro-tyrosine C.H. (N) 12)2NO. [115°]. Formed by warming nitro-ty/osine with dilute HNO₃. Golden plates (from water).—Salts: CaC₂H,N₂O, Baq. Golden ix-sided tables.— BaC₃H,N₂O, 2aq. Ruby-red prisms with yellow reflex, exploding when heated.

Reference.—AMIDO-TYROSINE.

TYROTOXICON. A poisonous substance formed in milk, containing the butyric acid ferment, which has stood for some days. Occurs also sometimes in cheese (Vaughan, Ph. [3] 18, 479; H. 10, 146). When its alcoholic solution is mixed with PtCl, and evaporated violent explosion takes place.

U

ULEXINE C. H., N. O. [151°]. An alkaloid in the seeds of common furze (Ulex europæus) (Gerrard a. Symons, Ph. [3] 17, 101, 229; 19, 1029; 20, 978, 1017). Deliquescent crystals, v. sol. chloroform, insol. ether. Poison, paralysing respiration (Bradford, J. Physiol. 8,79). Possibly identical with cytisine. -B'HCl. Deliquescent.

B'H.PtCl. - B'2HAuCl.
ULTRAMARINE. The blue-coloured material contained in lapis-lazuli. Until 1828 ultramarine was obtained by powdering and washing lapis-lazuli; but since that year it has been manufactured by heating to bright redness mixtures of Al silicate (china clay), Na₂CO₈ (carbonated soda ash), S, and charcoal, and washing, powdering, drying, and sifting the product. (For details of the manufacture v. DICTIONARY OF APPLIED CHEMISTRY, vol. iii. p. 891.) The constituent elements of ultramarine are Al, Na, Si, S, and O. Treated with HClAq, ultramarine gives off H2S; the solution contains NaCl AlCl3, and SiO₂, and the residue consists of SiO₂, S, and some Al₂O₃; these reactions point to ultramarine being a compound of silicates and polysulphides. Much work has been done on the constitution of ultramarine, but the results obtained have not finally settled the question. From analyses and investigations made in 1891, Brögger a. Bäckström (W. J. 1891. 454) conclude that the composition of the essential colouring compound in natural ultramarine is represented by the formula Na,[Al(S,Na)]Al,(SiÔ,). They also conclude that artificial ultramarines may be expressed by the following formulæ:-

> $Na_3Al_3Si_3O_{12} = N$ $\begin{array}{l} Na_2Al_2Si_4O_{12}=A\\ Na_4[Al(SNa)]Al_2Si_2O_{12}=U(S_1)\\ Na_4[Al(SNa)]Al_2Si_3O_{12}=U(S_2)\\ Na_4[Al(S_2Na)]Al_2Si_3O_{12}=U(S_2)\\ Na_4[Al(S_3Na)]Al_2Si_3O_{12}=U(S_3). \end{array}$

The main constituents of the different varieties of blue ultramarine are N, A, and $U(S_n)$; of green ultramarine $U(S_n)$ and N; and $U(S_n)$ is the main constituent of white ultramarine, according to B. a. B. Formulæ more or less approaching those given above have been assigned to ultramarine by other observers; thus Silber 1880 represented blue ultramarine as Si.Al. Na.S.O. (B. 13, 1854). On the other hand, some chemists have regarded the egsential constituent of ultramarine as a sodium thiosilicate; for instance, Rickmann (B. 11, 2013 [1878]) gives S.Si(ONa), as the composition of 'ideal ultramarine blue'; and he regards artificial ultramarine as a mixture, the only essential part of which is sodium thiosilicate. Clarke (Am. 10, 126) suggested a formula for ultramarine which should indicate the relations of this body to various natural silicates; in this respect Clarke's formula is important (v. this vol. p. 451).

White ultramarine is obtained by completely cutting off air during the roasting of the materials; it is changed to blue by heating in O. SO₂, or Cl (v. Ritter, W. J. 1160. 226; cf. Philipp. B. 9, 1109; 10, 1227; Böttinger, A. 182, 311; R. Hoffmann, A. 194, 1).

Red ultramarine was observed by Scheffer, in 1873, to be formed in preparing the ordinary substance in a muffle furnace very strongly heated and freely exposed to the air; it contains less Na and more Al than the blue variety (B. 6, 1450; v. also Büchner, D. P. J. 231, 446;

Zettner, B. 8, 259, 353).

Yellow ultramarine is formed by heating the red variety in the air a little above 360°, for a short time (R. Hoffmann, A. 194, 1). According to Büchner (B. 7, 990), both the yellow and red forms are produced by heating blue ultramarine to 300°-400° in O or SO₂; the colour changes to red and then to yellow. If Cl is passed over ultramarine before the formation of the blue variety is completed, at 410°, the colour goes green, and then reddish-yellow (Zettner, B. 8, 259, 353); by heating the product with alkali, all Cl is removed and a violet ultramarine is produced. This variety goes to the red form when heated in vapour of HNO, or HCl at 1300-150° (Zettner, l.c.).

Green ultramarine is formed when a little Sn is used in the roasting process; it is converted into the blue substance by roasting with S. This form seems to be intermediate between white and blue ultramarine. According to Philipp (B. 9, 1109), S is not taken up in the change from green to blue; by heating the green form in sealed tubes with water, at 100°, he obtained the blue, the water removing a very small

quantity of sodium compounds.

Substituted ultramarines. Blue to green, and reddish-grey substances have been formed by heating ultra narine with AgNO, Aq, and by treating the pro uct of this reaction with KI, LiI, BaI2, ZnI2, & . These substances, known as silver ultramarine, potassium ultramarine, &c., probably contain Ag, K, Li, Ba, &c., in place of Na (v. Unger, D. P. J. 212, 224, 301; Philipp, B. 10, 1227; Heumann, A. 199, 253; 201, 262; Forcrand a. Ballin, Bl. [2] 30, 112). By heating silver ultramarine with various alcoholic iodides, Forcrand (C. R. 88, 30) obtained ultramarines containing alcoholic radicles, such as ethyl ultramarine, amyl ultramarine, &c. Selenion and tellurium ultramarines have also been produced, wherein S is substituted by Se or Te (v. Leykauf, W. J. 1876. 555; Guimet, A. Ch.,[5] 13, 102; Plicque, Bl. [2] 28, 518; Morel, Bl. [2] 28, 522). Various substituted ultramarines have been examined by Wunder (Chem. Zeit. 1890. 1119). M. M. P. M.

ULMIC ACID. C. 66.4-63.9; H. 4.6-4.5; O. 29-31-5. C₁₈H₁₄O₈ or C₁₈H₁₆O₇. H.C. 1983, 200. H.F. 266,200. Formed by action of HCl on cane sugar. Is an anhydride or mixture of anhydrides $C_{18}H_{14}O_{e}$ (humic anhydride) (Berthelot a. André, Bl. [3] 7, 441, 451). Turns yellow in air and sunlight, giving off CO_{2} . The same thing happens with the humic acid extracted from vegetable mould by KOHAq and ppd. by HCl. Dilute alkalis cause it to swell up, forming insoluble salts and a small quantity of a soluble basic salt. When conc. KOHAq is used a salt is formed, from which two-thirds of the alkali can be removed by washing, leaving KC₁₈H₁₅O, aq, and is reconverted into the acid C₁₈H₁₅O, by HClAq. Ammonia converts humic anhydride (ulmic acid) into an amido acid $C_{54}H_{47}NO_{19}NH_8$, whence HCl sets free $C_{54}H_{47}NO_{19}$.

Salts.—KÖ₁₈H₁₀O, aq. Hard, black, insoluble mass, yielding a brown powder. The 'ulmin' of Malaguti. Formed by shaking the Hard, black, inanhydride with KOH (1 pt.) and water (120 pts.). -NaC₁₈ \mathbf{H}_{13} O, aq.—Na, $\mathbf{C}_{18}\mathbf{H}_{13}$ O, xaq.

References.—Humic, Saccharumic, and Sac-

CULMIC ACIDS.

UMBELLIC ACID C.H.O. i.e.

[4:2:1] C₆H₃(OH)₂·CH:CH.CO₂H. Mol. w. 182. DI-OXY-CINNAMIC ACID. Formed by warming its lactone, umbelliferone with KOHAq (Tiemann a. Reimer, B. 12, 994; Posen, B. 14, 2745). Yellowish powder. Turns brown at 240°, and decomposes below 260°. Sol. alcohol and warm water, insol. ether. Resinified by boiling its aqueous solution. Reduces warm ammoniacal AgNO₂, forming a mirror. FeCl₃ gives a brown pp.

Methyl derivative C₁₀H₁₀O₄. [180°-185°]. Got by heating herniarin with alcoholic potash at 160° (Barth a. Herzig, M. 10, 165). Needles

(from water).

Vol. IV.

(a) - Di-methyl derivative C.H. (OMe), CH:CH.CO.H. [138°]. Formed by the action of MeI and KOH in MeOH on the methyl derivative of umbelliferone in the cold (W. Will, B. 16, 2115; 19, 1777). Needles, v. sol. alcohol and ether. Converted into the (\$)-isomeride by boiling alone or with HClAq. Both isomerides are reduced by sodium-amalgam to the same C.H. (OMe). CH. CH. CO.H, and are oxidised by KMnO, to the same C, H, (OMe), CO, H. CaA', 2aq: crystals.—BaA', 2aq: needles.

[184°]. (β) - Di - methylderivative. Formed as above, and also by saponification of its methyl ether C.H. (OMe)2.CH:CH.CO.Me [87°], which is got by evaporating methyl-umbelliferone (1 mol.) with NaOH (2 mols.) nearly to dryness, and digesting the residue with MeI and MeOH (Tiemann a. Will, B. 15, 2079). Needles, sol. alcohol and ether. Sl. sol. cold water.

(a)-Di-ethyl derivative [4:2:1] C₂H₃(OEt)₂C₂H₂CO₂H. Di-ethyl-umbellic acid. [107°]. Formed by mixing ethylumbelliferone (5 g.) with EtI (10 g.) and a solution of KOH (4 g.) in alcohol (Will a. Beck, B. 19, 1780). Small plates, v. sol. alcohol, ether,

and benzene.

(β)-Di-ethyl derivative. [200°]. Formed by boiling the (a)-isomeride for some time, and also by heating ethyl-umbelliferone (10 g.) with EtI (9 g.) and a solution of Na $(2\frac{1}{2}$ g.) in alcohol for six hours at 150°. Crystals, m. sol. alcohol, sl. sol. water.

UMBELLIFERONE C,H,O, i.e.

 $C_6H_3(OH) < \begin{array}{c} CH:CH \\ O-CO \end{array}$ Mol. w. 162. [224]. 1 at 100°. Formed by the dry distillation of various resins, chiefly those derived from umbelliferous plants, e.g. galbanum, sagapenum, and asafætida (Sommer a. Zwenger, A. 115, 15; Mössmer, A. 119, 260; Hlasiwetz a. Grabowski, A. 139, 100; Hirschsohn, C. C. 1877, 182). Formed by heating malic acid (1 mol.) with resorcin (1 mol.) and H2SO4 (Pechmann, B. 17, 932). Prepared by distilling the alcoholic extract of gum galbanum (Tiemann a. Reimer, B. 12, 993). Needles, sl. sol. cold water and ether, v. sol. alcohol. Dissolves in acids without change. Alkalis above 50° form umbellic acid. Its solution in dilute alkalis shows blue fluorescence. H2SO4 forms a solution with green fluorescence. Reduces AgNO₃ on boiling. Yields (4,2,1)-di-oxy-benzoic acid on oxidation (Tiemann a. Parrisius, B. 13, 2354). Fuming HNO, and H₂SO₄ form C₂H₂(NO₂)₂O₃ [216°] (Posen, B. 14,

Acetyl derivative C₁₁H₈O₄. [140°]. Formed by acetylation, and also by boiling di-oxy-benzoic aldehyde with Ac2O and NaOAc (Tiemann, B. 10, 2216; 12, 995). Needles, sl.

sol. water.

Methyl derivative C, H,O, i.s.

 $C_6H_3(OMe) < \begin{array}{c} CH:CH \\ O-CO \end{array}$. [114°]. Formed by heat-

ing umbelliferone with KOMe and MeI (T. a. R.). Leaflets, nearly insol. water. Appears to be identical with herniarin [118°], which is extracted by alcohol from Herniaria hirsuta (Barth a. Herzig, M. 10, 161).

Ethyl derivative C₁₁H₁₀O₂. [88°]. Plates, v. sol. alcohol (Will a. Beck, B. 19, 1779). Oxim of the methyl derivative

[4:2:1] C_eH_s(OMe) < CH:CH O—C:NOH. [138°]. Formed

from the methyl derivative of thio-umbelliferone and hydroxylamine (Aldringen, B. 24, 3465). Needles, v. sol. hot water and alcohol. FeCl. colours its solation brownish-red.

Phenyl-hydraside of the methyl derivative C.H.(OMe) CH:CH O-C:N,HPh. Yellow needles, giving a blue colour with H.SO.

Thio-umbelliferone. Methyl derivative $C_eH_s(OMe)<_{O-CS}^{CH:CH}$. [114°]. Formed by heating the methyl derivative of umbelliferone with P.S. (Aldringen, B. 24, 8465). Needles, sol. alcohol, v. sl. sol. water.

References .- Bromo- and METHYL- UMBEL-

LIFERONE

UMBELLOL C. H. O. (216°). V.D. 4.29. Occurs in the volatile oil of the Californian laurel (Stillmann, B. 13, 629). Oil, sol. H2SO4 with red colour, becoming black. Attacked by

UMBELLULIC ACID v. HENDECOÏC ACID. UNDECANE v. HENDECANE.

UNDECOIC ACID v. HENDECOIC ACID. UNDECOLIC ACID v. HENDECINOIC ACID.

UNDECYLENE v. HENDECYLENE. UNDECYLENIC ACID v. HENDECENOIC ACID. UNDECYLIC ACID v. HENDECOIC ACID.

URACIL CO<NH.CH CH. Di-oxy-pyr-Derivatives of this compound are formed by the action of acetoacetic ether on urea, thio-urea, and guanidine (Behrend, A. 229, 1; 240, 6; List, A. 236, 1; Jäger, A. 262, 365;

Warmington, J. pr. [2] 47, 201).

Amido-uracil C₄H₄N₂O₂. Formed, together with oxyuracil C₄H₄N₂O₂, by reducing nitrouracil (Behrend), and by boiling di-bromo-pyruvuramide with baryta-water (Fischer, A. 239, 193). Yellow needles, sol. water. Converted by cyanic acid into oxyxanthine $C_3H_aN_4Q_3$, which crystallises in needles, sl. sol. water, sol. alkalis, gives with Cl the murexide reaction, may be oxidised to alloxantin, and gives some alloxan with bromine-water.

Acetyl derivative C, H, N,O, 1 aq. Prisms. Nitro-uracil v. NITRO-DI-OXY-PYRIMIDINE. Bromo-nitro-oxy-uracil. Dihydride

C4H4N4O5Br i.e. CO NH.CH(OH) CBr(NO2). Formed by adding Br to nitro-uracil suspended in water (Behrend, A. 240, 11). Crystalline mass, sl. sol. cold water and alcohol. Boiling with water produces brominated nitro-methanes and a crystalline compound C₅H₇N₅O₅, formed by the action of urea upon nitro-uracil. Tin and HCl reduce it to amido-uracil.

Methyl-uracil v. DI-OXY-METHYL-PYRIMIDINE. Nitro-methyl-uracil C.H.N.O. Prepared by nitrating methyl-uracil with HNOs (S.G. 1.5) mixed with an equal volume of H2SO4 (S.G. 1.84) (Behrend, A. 240, 3). Prisms or tables (from water), sl. sol. hot alcohol.

Isomeride v. NITRO-DI-OXY-METHYL-PYRIM-

Nitro-di-methyl-uracil v. NITRO-DI-OXY-DI-METHYL-PYRIMIDINE.

Di-chloro-oxy-methyl-uracil. Dihydride CCL CMe(OH).NH CO. C,H,Cl,N,O, i.e. Formed by the action of chlorine on methyluracil suspended in water (Behrend, A. 236, 59). Tables (from water); slowly decomposed by boiling water. M. sol. alcohol; not decomposed by boiling alcohol. M. sol. ether. A solution of SnCl₂ in HClAq at 100° reduces it to chloromethyl-uracil O.H.ClN2O, which crystallises from water in needles. Fuming HNO, gives dichloro-barbituric acid.

Di-bromo-oxy-methyl-uracil J.H.N.Br.O. i.e. CBr₂ CMe(OH).NH . From lethyl-uracil (or bromo-methyl-uracil), and bromine water (Behrend, A. 229, 18; 236, 57). Jubes (from hot water). Boiling alcohol conve ts it into bromomethyl-uracil C,H,N,BrO2. Fuming HNO, forms di-bromo-barbiturio acid.

Di-chloro-oxy-tri-methyl-uracil. Dihydride $CO < NMe.CO \longrightarrow COl_2$. [144°]. by passing Cl into an aqueous solution of trimethyl-uracil (Hagen, A. 244, 14). Crystals, sl. sol. hot water. Reduced by boiling with SnCl, to chloro-tri-methyl-uracil, which crystallises in prisms, m. sol. hot water.

Nitro-ethyl-uracil v. NITRO-DI-OXY-ETHYL-

PYRIMIDINE.

Methyl-ethyl-uracil v. DI-OXY-METHYL-ETHYL-PYRIMIDINE.

Nitro-methyl-ethyl-uracil v. NITRO-DI-OXX-METHYL-ETHYL-PYRIMIDINE.

Amido-uracil carboxylic acid v. Dr-oxy-AMIDO-PYRIMIDINE CARBOXYLIC ACID. References .- THIO-METHYL-URACIL and UR-

AMIDO-CROTONIC ACID.

URAMIDO - ACETIC ACID v. HYDANTOÏO ACTD.

p-URAMIDO-BENZENE SULPHONIC ACID NH2.CO.NH.C8H4.SO8H. Formed by evaporating a solution of p-amido-benzene sulphonic acid and potassium cyanate (Pellizzari, A. 248, 156), and also by heating dry p-amido-benzene sulphonic acid (1 mol.) with urea (11 mols.) (Friedel, C. R. 112, 868). Pearly plates (from dilute alcohol), v. sol. water. - BaA', 3aq; monoclinic prisms, sol. Aq. o-URAMIDO-BENZOIC ACID C₈H₈N₂O₂ i.e.

NH2.CO.NH.C8H4.CO2H. Formed from potassium cyanate and the hydrochloride of o-amidobenzoic acid (Griess, J. pr. [2] 5, 871; B. 11, 1730). HNO, yields only one di-nitro-derivative.

m-Uramido-benzoic acid C₈H₈N₂O₉. S. 1 at 100°. S. (96 p.c. alcohol) '72 at 16°. S. (ether) '127 at 16°. Occurs in urine after a dose of m-amido-benzoic acid (E. Salkowski, H. 7, 113).

Formation.—1. By adding potassium cyanate to a boiling saturated solution of the sulphate of m-amido-benzoic acid (Menschutkin, A. 153, 84; Z. [2] 4, 275).—2. By fusing m-amido-benzoic acid with urea (Griess, Z. [2] 5, 812; B. 2, 47). 3. By boiling m-cyanamido-benzoic acid with HClAq (Traube, B. 15, 2122).

Properties.—Small prisms (containing aq), v. sl. sol. hot water. At 200° it yields V. 81. 801. 10t water. As a condition of the condition of benzoic acid, CO2, and NH3.

Salts. NH, A'aq. - KA'. - CaA', 4aq. -PbA', 2aq. -AgA': curdy pp., becoming crystalline.

Ethyl ether EtA'. [176°]. Formed from m-amido-benzoic ether and KCyO (Griese, J. pr.

[2] 4, 293). Plates, sl. sol. hot water. Amide C.H.N.O. Formed from m-amido-benzamide and KONO (Menschutkin). Scales (from water), decomposed by fusion, al. sol.

alcohol.

p-Uramido-be agoic acid C_sH_sN₂O_s. Formed in like manner (1 riess, J. pr. [2] 5, 369). Plates, almost insol. colc water, v. sl. sol. hot water, m. sol. hot alcohol. Yields some p-amido-benzo-nitrile [86°] on listilling with P₂O₅.—BaA'₂. Plates, v. e. sol. cold water.

Di-uramido-benzoic acid C.H.O.N.O. (NH2.CO.NH)2C6H3.CO2H. Formed by fusing diamido-benzoic acid with urea (Griess, B. 2, 47). Small granules, v. sl. sol. hot water.—BaA'₂ (dried at 130°). Nodules, v. sol. hot water.

References .- AMIDO-, NITRO-, and OXY- UR-

AMIDO-BENZOIC ACIDS.

URAMIDO-BENZOYL v. OXY-QUINAZOLINE.

p-URAMIDO-BENZYL-UREA C₀H₁₂N₁O₂ i.e. NH₂.CO.NH.C₆H₄.CH₂.NH.CO.NH₂. [197°]. p-ω-Di-uramido-toluene. Formed by heating together p-amido-benzyl-amine, hydrochloride (1 mol.), and KCNO (2 mols.) in aqueous solution (Amsel a. Hofmann, B. 19, 1289). Small needles. Sol. hot water.

URAMIDO-ISOBUTYRIC ACID v. ACETON-

URAMIC ACID.

o-URAMIDO-CINNAMIC ACID $C_{10}H_{10}N_2O_3$ i.e. NH₂.CO.NH.C₆H₄.CH:CH.CO₂H. Formed from o-amido-cinnamic acid and aqueous KCNO (Rothschild, B. 23, 3341). Sol. NH3Aq and hot

HClAq.

(β)-URAMIDO-CROTONIC ACID C₅H₈N₂O₃ i.e. CH₃.C(NH.CO.NH₂):CH.CO₂H. The ether is converted by NaOHAq into the salt NaA', which crystallises in needles, v. sol. water; but on adding acids to this salt di-oxy-methyl-pyrimidine C₅H₆N₂O₂ is produced. Methyl-uracil is reconverted by NaOHAq into sodium uramidocrotonate (Behrend, A. 229, 5)

Ethyl ether EtA'. [166°]. Formed from acetoacetic ether and urea. Needles, v. sol. ether, m. sol. alcohol. Decomposed by warm HClAq into urea, acetone, CO₂, and alcohol. Alcohol forms CH₃.C(NH.CO₂Et):CH.CO₂Et [29°] (Meister, A. 244, 234).

CH3.C(NH.CO.NH2):CH.CO.NH2. Amide [131°]. Crystals (containing EtOH) (Meister). Amido-uramido-crotonic acid. Lactam.

 $C_3H_3(NH_2)N_2O_3$ i.e. $CO < NH.CMe > C.NH_2$ (?). Amido-methyl-uracil. Methyl-uracil (v. DI-oxx-METHYL-PYRIMIDINE) is converted by Br in CS2 into crystalline bromo-methyl-uracil C₅H₅BrN₂O₂, whence conc. NH₂Aq at 150° forms amidomethyl-uracil (Behrend, A. 231, 250). Satiny plates (from water), melting near 250°. Sol. alcohol. The hydrochloride is converted by potassium cyanate into oxy-methyl-xanthine $C_0H_0N_4\tilde{O}_2$ 2aq, orystallising from water in prisms.—B'HCl aq. Monoclinic crystals; $a:b:c=1\cdot629:1:2\cdot345$; $\beta=52^\circ$ 57'.—B'₂H₂PtCl₆ 2aq: needles, v. sol. water.

Acetyl derivative C,H,(NHAc)N,O, 1aq. Formed from oxy-methyl-xanthine and Ac.O at 170°. Thick trimetric prisms (from water). Decomposes near 210° without melting. Sol.

alkalis and reppd. by acids.

URAMIDO-ETHANE SULPHONIC ACID

NH2.CO.NH.CH2.CH2.SO3H. Taurocarbamic acid. Occurs in small quantity in urine after a dose of taurine (E. Salkowski, B. 6, 744, 1191). Formed by evaporating a solution of taurine with KCyO. Formed also by the action of bromine-water on ethylene-\psi-thio-urea (Gabriel, B. 22, 1142). Dimetric plates, v. sol. water, sl. sol. alcohol, insol. ether. Decomposed by baryta-water at 140° into CO2, taurine, and NH3.—BaA'2: tables (from alcohol).—AgA'. Radiating tufts of crys-

URAMIDO - HEXOIC ACID C,H14N2Os i.e. NH₂.CO.NH.CH(C,H₂).CO₂H. [200°]. Formed by boiling isobutyl-hydantoin with baryta-water (Pinner a. Spilker, B. 22, 696). Needles, v. sol. alcohol and hot water.—BaA'2: nodular groups of prisms.

Ethyl ether EtA'. [135°]. Formed by heating isobutyl-hydantoin with KOH and EtBr.

Needles, v. sol. alcohol and hot water.

Nitrile. NH2.CO.NH.CH(C4H9).CN. Formed by heating the cyanhydrin of isovaleric aldehyde with urea (Pinner a. Lifschütz, B. 20, 2351). Oil, sol. alcohol and ether. On warming with HClAq it yields isobutyl-hydantoin

 $C_4H_9.CH < \stackrel{CO.NH}{NH.CO} [210^\circ].$

Amide NH2.CO.NH.CH(C4H4).CO.NH2. [170°]. Formed from the nitrile (1 pt.) and conc. H₂SO₄ (8 pts.) in the cold. Crystals, v. sol. water and alcohol.

URAMIDO-HIPPURIC ACID C, H, IN, O, Formed by fusing m-amido-hippuric acid with urea (Griess, J. pr. [2] 1, 235). Nodules, v. e. sol. hot water. Decomposed by boiling conc. HClAq into glycocoll and m-uramido-benzoic acid.—AgA'. Crystalline pp.
URAMIDO-NITRO-AMIDO-PHENOL

C₂H₈N₄O₄ i.e. C₆H₂(NO₂)(NH₂)(NH.CO.NH₂).OH. Formed by reducing di-nitro-oxy-phenyl-urea. with ammonium sulphide (Griess, J. pr. [2] 5, 1). Reddish-brown needles, sl. sol. hot water and alcohol.-B'HCl: white scales.-BaA', 31 aq: steel-blue needles, v. sol. hot. water. Boiling HClAq converts it into amidocarboxamidonitrophenol C, H, N, O,, which yields BaA' 2 xaq and B'HClaq, and is reduced by tin and HClAq to diamidocarboxamidophenol C,H,N,O, which yields B'HCl, crystallising in plates.

URAMIDO-DI-NITRO-PHENOL v. DI-NITRO-

OXY-PHENYL-UREA

p-URAMIDO-PHENYL-ACETIC ACID
NH., CO.NH.C., H., CH., CO., H. [174°]. Formed
by boiling p-cyanamido-phenyl-acetic acid with HClAq (Traube, B. 15, 2122). Small crystals (containing 1 aq), sol. water, alcohol, and ether.

acid. a-Uramido-phenyl-acetic Ethylether NH2.CO.NH.CHPh.CO2Et. [139°]. Formed by the action of aqueous KCyO on the hydrochloride of α-amido-phenyl-acetic ether (Kossel, B. 24, 4150). V. sol. alcohol and hot water.
Nitrile NH₂CO.NH.CHPh.CN. [17

[1709] Formed by heating the cyanhydrin of benzoic aldehyde with urea (Pinner a. Lifschütz, B. 20, 2351). Converted by acids into C.H.N.O. [1820]. crystallising in scales, v. sol. alcohol and hot water.

a-URAMIDO-PHENYL-CROTONIC ACID. Nitrile CHPh:CH.CH(NH.CO.NH2).CN.

[160°]. Formed by heating ures with the cyanhydrin of cinnamic aldehyde (Pinner a. Lifschütz, B. 20, 2351). Needles, v. sol. hot alcohol. Boiling HClAq converts it into di-oxy-styryl-pyrazole

Amids CHPh:CH.CH(NH.CO.NH.).CO.NH. Formed from the nitrile and cold conc. H.SO. (Pinner a. Spilker, B. 22, 692). Microcrystalline

3 e 2

powder, decomposing at 210°-220° with evolution of NH_a. Sol. alcohol, sl. sol. hot water.

a-URAMIDO-PROPIONIC ACID C₄H₈N₂O₈ i.e. CH₂-CH(NH.CO.NH₂).CO₂H. Lacturamic acid. [155°]. Formed by evaporating a solution of alanine sulphate mixed with KCyO (Urech, A. 165, 99). Formed also by boiling lactyl-urea with baryta-water (Heintz, A. 169, 128). Prisms, sl. sol. cold water and alcohol, insol. ether.—BaA'₂ aq (dried at 100°).—PbA'2aq.—AgA': needles, m. sol. water.

Nitrile CH, CH(NH.CO.NH,).CO.H.

[106°] (Franchimont a. Klobbie, R. T. C. 7, 16). Amide CH₂.CH(NH.CO.NH₂).CO.NH₄.

[196°]. Small needles (from alcohol), sol. water. URAMIDO.SUCCINIC ACID. Amic acid NH. CO.CH., CH(NH.CO.NH.).CO.H. [188°] (G.); [157°] (P.). Succinuramide. Amidosuccinuric acid. S. 5·4 at 23°. Formed by evaporating the mixed solutions of asparagine KCyO (Guareschi, G. 7, 404; B. 10, 1747). Dextro-asparagine gives a lævo- compound, while lævo-asparagine gives a dextro- compound (Piutti, B. 19, 1693). Prisms, nearly insol. alcohol and ether. According to Piutti, the dextro- and lævo- compounds both melt at 157°.

Reactions.—1. Decomposed by fusion, with formation of malyl-ureide C₅H,N₅O₅ [230°–235°], which is also formed by heating urea with asparagine at 125° (Grimaux, A. Ch. [5] 11, 400) and crystallises from water in pointed rhombohedra.—2. Boiling HClAq (S.G. 1·12) forms malyl-ureidic acid C₅H₅N₂O₄, which is also got by heating aspartic acid with urea at 130°. Malyl-ureidic acid crystallises in prisms [215°–220°], S.25 at 100°, which yields BaA'₂aq and is converted by heating with bromine-water into C₅H₂Br₈N₂O₅, S. 3 at 100°, C₅H₄Br₄N₄O₅, S. 25 at 100°, C₅H₄Br₂N₄O₅, which is v. sol. water, C₅H₂BrN₄O₄, S. 6. 6 at 100°, and C₄H₄Br₂N₂O₅, S. 10 in the cold and c. 22 at 100°.

URAMIDO-THIO-FORMIC ACID v. THIO-ALLOPHANIC ACID.

DI-URAMIDO-TOLUENE v. URAMIDO-BENZYL-REA.

DI-p-URAMIDO-DI-o-TOLYL-SULPHIDE [2:1:4] S(C_sH₃Me.NH.CO.NH₂)₂. Thio-p-tolyl-di-urea. [151°] (when containing benzene of crystallisation). Formed by heating the hydrochloride of di-p-amido-di-tolyl-sulphide with KCNO (Truhlar, B. 20, 669). Crystallises from benzene in white silky needles (containing C_sH_s).

URAMIL O.H.N.O. i.s.

CO NH.CO CH.NH. Dialuramide. Amido-

barbituric acid. Amido-malonyl-urea. Murexan. Mol. w. 143. Formed by boiling alloxantin with NH₂Cl (Liebig a. Wöhler, A. 26, 310; Grimaux, C. R. 87, 752) and by boiling ammonium thionurate with HClAq. Formed also by reducing nitro- or nitroso- barbituric acid with HIAq (Baeyer, A. 127, 223). It is also a product of the action of acids on murexide (Beilstein, A. 107, 183).

Properties.—Silky needles, al. sol. hot water, insol. alcohol and ether. Sol. conc. H_2SO_4 and re-ppd. on dilution. Its alkaline solution turns red in air, forming purpurates; thus it is converted in ammoniacal solution into murexide by boiling with air or HgO. Conc. HNO_g forms alloxan.

Bromine at 85° forms orange C₄H₄BrN₅O₅ (Mulder, B. 14, 1060).

URAMILIC ACID C₂H₂N₂O (?). S. 14 in the cold; 33 at 100°. Formed by heating uramil with H₂SO₄ and water, or by evaporating ammonium thionurate with som H₂SO₄ (Liebig a. Wöhler, A. 26, 314; Gregory, P. M. 24, 187). Transparent, four-sided prisms, sl. sol. cold water. Converted by boiling conc. HNO₃ into nitrobarbituric acid.

URANATES v. URANIUM, OXYAGIDS AND SALTS OF, p. 826.

URANIUM. U. At. w. 239 (not determined with very great accuracy), Mol. w. unknown. M.P. not determined accurately; between red heat and white heat. S.G. of fused U at $\frac{150}{100} = 18.685$ (Zimmermann, B. 15, 851). S.H. at $100^{\circ} = 0276$ (Z., l.c.). S.V.S. 12.8.

Historical.—In 1789 Klaproth prepared some compounds of a new element from pitchblende; to the characteristic element of these compounds he gave the name uranium, from gupavos = the heavens, in allusion to the planet Uranus which had been discovered a few years before (Crell's A. 1789 [2] 400). The compounds of U were examined by Richter (Gehlen's A. 4, 402), Buchholz (Gehlen's A. 4, 17, 134), Lecanu (J. Ph. 9, 141; 11, 279), Brande (S. 44, 1), Berzelius (P. 1, 359), and Artvedson (P. 1, 245). In 1840, Péligot showed that the substance supposed till then to be U was really an oxide of U (A. Ch. [3] 5, 5); P. isolated the metal, and determined the at. w. to be 120. Mendeléeff in 1872 (A. Suppl. 8, 133; cf. C. N. 41, 39 [1880]) proposed to double the at. w. of U, making it 240; this proposal was completely justified by the determination of S.H. of pure U by Zimmermann in 1882 (B. 15, 851), and by the determination of the V.D. of UCl, and UBr, (Z., A. 216, 2 [1883]).

Occurrence.—Never uncombined. A few minerals containing compounds of U are found sparsely distributed and not in large quantities; the chief of these is pitchblende, which consists of U₃O₈ along with SiO₂ and oxides of Sb, As, Ca, Cu, Fe, Mg, V, &c. Carbonate, sulphate, phosphate, niobate, tantalate, and various other compounds of U are also found in varying quantities in several rare minerals (v. Zimmermann, A. 232, 300; Hidden a. Mackintosh, Am. S. [3] 38, 474; Hillebrand, Am. S. [3] 38, 329; 40, 384). Lockyer (Pr. 27, 49) stated that some of the lines in the emission-spectrum of U coincide with dark lines in the solar spectrum. Hutchins a. Holden (P. M. [5] 24, 325) think there is no trustworthy evidence in favour of the existence of U in the solar atmosphere.

Preparation.—1. A mixture of UCl₄, NaCl, and Na is strongly heated in a closed iron cylinder. Zimmermann (A. 116, 14) used a cylinder of soft iron with walls c. 3 centim. thick, bored from a larger cylinder (similar to that employed by Nilson a. Pettersson for preparing Be; B. 11, 883) fitted with a cap which could be screwed on tightly. He placed a layer of pure fused NaCl in the cylinder, then small pieces of Na (c. 3 to 4 pts. for 10° pts. UCl₄), then UCl₄ (which must be put into the cylinder as quickly as possible, as it is chygroscopic), and covered the whole with a layer of fused NaCl. Z. raised the cylinder to a white heat, kept it at

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that temperature for some time, cooled, and washed the con ents with dilute alkali solution. and then with vater. According to Z., the metal as thus prepare it is quite pure (v. infra).—2. Moissan (C. R. 116, 347) obtained U, containing from 5 to 13 5 p. C, by mixing the oxide prepared by strongly heating UO2(NO3)2 with excess of charcoal, compressing the mixture in a carbon crucible, imbedding this in MgO, and heating for 10-12 minutes in an electric furnace, using a current of 60 volts and 450 ampères.

UCl4 is obtained by heating the nitrate UO2(NO3)2 in air, mixing the oxide thus obtained (chiefly U_sO_s) with C, and heating in a stream of Cl (v. Uranium tetrachloride, p. 823). The nitrate is obtained from pitchblende by powdering the mineral, washing with water, drying, and washing to remove S and As, dissolving in HNO3Aq, evaporating to dryness, dissolving in water, filtering from Fe₂O₃, PbSO₄, &c., evaporating a little, and allowing to crystallise. The crystals of UO2(NO3)2 thus obtained are purified by draining in a funnel, washing with a little cold water, drying in the air, dissolving in ether in a wide-mouthed bottle, crystallising by spontantous evaporation, and re-crystallising from hot water; the crystals of UO2(NOs)2 are then dissolved in water so as to form a conc. solution, UO₂.C₂O₄ is ppd. by adding H₂C₂O₄Aq to this solution, the pp. is washed with boiling water, and then decomposed by heat, and the UO2 thus produced is digested with conc. HClAq, washed, dissolved in HNO, Aq, and the solution is crystallised (Péligot, A. Ch. [3] 5, 5; 12, 258; cf. Ebelmen, A. 43, 286; Wertheim, J. pr. 29, 209).

The oxide U₃O₈ used by Zimmermann (v. supra) for the preparation of UCl, from which pure U was obtained, was prepared by him as follows. A warm solution of commercial uranium hydroxide in HClAq was saturated with H.S; after standing for some time the liquid was filtered off, NH₃Aq and (NH₄)₂CO₃Aq were added in excess, the liquid was warmed and (NH₄)₂SAq was added; after standing the liquid was filtered, saturated with HClAq, and boiled till CO2 was removed, then mixed with NH3Aq and (NH₄)₂SAq and heated on the water-bath till most of the (NH₄)₂S was removed; the pp. of UO2 was collected, washed, and heated over the blowpipe; the U₂O₈ thus formed was dissolved in HNO₂Aq and the solution was crystallised; the crystals of UO₂(NO₃)₂ were dissolved in ether, the solution was poured off from insoluble substances, evaporated to dryness, and the residue was strongly heated; the U₂O₈ which remained was again dissolved in HNO, Aq, and the crystals that formed on evaporation were dissolved in ether, and the ethereal solution was evaporated to dryness and the residue decomposed by heating.

Properties. - A lustrous, hard, silver-white metal; S.G. 18.685 at $\frac{18^{\circ}}{46}$ (Zimmermann, B. 15, 851); if UCl, is reduced by Na at a red heat, U is obtained as a greyish-black powder. S.H. at $100^{\circ} = 0.0276$ (Z., l.c.). U is somewhat malleable, but cannot be hammered into thin plates. The emission-spectrum, obtained by the aid of electric sparks, shows only a few lines, the most marked being one in the yellow (Thalen, A. Ch. [4] 18, 239). For the fluorescence, phosphorescence, and absorption-spectra of U compounds v. Becquerel (A. Ch. [4] 27, 539), also Morton a. Bolton (Am. Ch. 3, 361, 401). U melts towards a white heat; according to Moissan (C. R. 116, 1429), the metal is volatilised by heating in an electric furnace using a current of 350 ampères and 75 volts. After long standing in the air the surface of U becomes covered with a steel-blue to black film of oxide; U that has been fused burns brilliantly when heated on Pt foil in the air; pulverulent U burns at 150°-170° in the air. U combines directly with Cl or Br, also with S; the metal dissolves in dilute H2SO.Aq or HClAq; it is not acted on by acetic acid, nor by KOHAq, NaOHAq, or NH₈Aq.

The at. w. of U has been determined (1) by determining U and Cl in UCl, (Péligot, A. Ch. [3] 5, 12 [1842]; Rammelsberg, P. 55, 318; 56, 125 [1842]); (2) by analyses of UO₂,C₂O₄,H₂O (Ebelmen, A. Ch. [3] 5, 191 [1842]; Peligot, A. Ch. [3] 20, 341 [1846]); (3) by analyses of UO₂(C,H₃O₂)₂,2H₂O (Peligot, Lc.; Wertheim, J. pr. 29, 207; (4) by analyses of the double salt J. p_7 , z_9 , $z_0(f)$; (4) by analyses of the double sale UO_2 .Na(C_2 H₃O₂)₃ (Wertheim, l.c.); (5) by transforming UO_2 into U_3 O₈, and UO_2 .Na(C_2 H₃O₂)₃ into Na₂UO₄ (Zimmermann, A. 232, 117, 273 [1886]); (6) by determining S.H. of U (Zimmermann, B. 15, 851 [1882]); (7) by determining V.D.s of UBr₄ and UCl₄ (Z., A. 216, 2 [1882])

[1883]).

The at. w. of U was formerly taken as c. 120; in 1872 Mendeléeff adduced reasons for doubling this value, based on a comparative study of the properties of compounds of U to which he was led by considering the periodic arrangement of the elements (A. Suppl. 8, 178; C. N. 41, 39). When the at. w. of U was taken as 120, the formulæ given to the chief oxides, chlorides, and salts of this element were UO, UCl2, USO4, &c.; U₃O₄; U₂O₃, UOCl, UO₂(NO₃), &c.; and U was classed with the iron metals. Mendeléeff pointed out that U was very much heavier than the iron metals, that the oxide U2O3 did not form salts similar to those derived from Fe₂O₂, that the salts of UO, i.e. USO, U(NO₃)₂, &c., were not isomorphous with the salts of MgO, FeO, NiO, &c. Mendeléeff also drew attention to the existence of a chloride which would have to be represented as U_2Cl_5 if U=120, and that this chloride was obtained by heating UCl, in H. Mendeléeff proposed to double the value then accepted for the at. w. of U, and, following on this, to write the formulæ of the chlorides, oxides, and salts as UCl₂, UO₂, U(SO₄)₂, &c.; U₅O₈; UO₃, UO₂Cl₂, UO₂(NO₃)₂, &c.; and UCl₅. It was pointed out that many of these formulæ are analogous to those of the corresponding compounds of Cr, Mo, and W; and it was shown that the properties of compounds of U more nearly resemble those of compounds of Cr, Mo, and W than of any other elements. Especial stress was laid by Mendeléeff on the acidic character of the highest oxide of U. Finally, Mendeléeff showed that an element with the properties of U, and an at. w. equal to c. 240, finds its proper place in Group VI., series 12, of the periodic arrangement of the elements. Mendeleeff suggested that the S.H. of pure U should be determined; that the V.D. of the volatile chloride should be found; that the action of H

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on the chloride then taken to be UCL should be studied; that selts of the lower oxide (UO if U=120, UO_2 if U=240) should be examined, especially as regards their isomorphism with salts of ThO₂, SnO₂, ZrO₂, and TeO₂; that the crystalline forms of corresponding compounds of MoO₃, WO₃, and UO₄ (U=240) should be studied, and that an especially thorough crystallographic examination should be made of the compounds R₂UO₂F₄; and that the physical properties of compounds of Cr, Mo, W, and U should be compared. Since the publication of Mendeléeff's memoir, the S.H. of U, and the V.D.s of UBr, and UCl, have been determined, the action of H on UCl, has been examined, and something has been done in the study of the crystalline forms of compounds of U, including those of the form R2UO2F4; the results of all these investigations fully confirm the justness of Mendeléeff's view that the at. w. of U is c. 240, and that this element must be classed with Cr, Mo, and W. The exact value to be given to the st. w. of U has not been yet determined; Zimmermann's measurements gave 238.75 from the ratio UO2:U8O8, and 238.67 from the ratio $UO_2Na(C_2H_3O_2)_3:Na_2UO_4$ (0=15.96). The mol. w. of U is not known.

U is the last member of the even-series family of Group VI.; this family also contains, besides O, the elements Cr, Mo, —, and W. The chromium elements are both metallic and non-metallic in their chemical relations; they form oxides MO₂ that are acidic, and some of the lower oxides are basic. The oxide UO₂ forms some corresponding salts; a series of salts of the form UO₂X is also known, where X = SO₄ &c. Several series of uranates M₂UO₄, M₂U₂O₇, &c., are known. The haloid compounds are UX₂, UX₄, and UX₅. (For a comparison of the metals of the Cr family v. Chromium group of elements, vol. ii. p. 168; cf. Classification, vol. ii. p. 207.)

ii. p. 207.)

The atom of U is tetravalent in the gaseous

molecules UBr, and UCl,

Reactions and Combinations.—1. Heated in air U is burnt to a voluminous deposit of U,O, (perhaps containing also UO₂), which stops the further oxidation of the metal. Pulverulent U burns when heated in air to 150°-170°.-2. U burns brilliantly when heated in chlorine, forming UCl4; heated in bromine to c. 240° part of the metal is changed to UBr.—3. Heated in sulphur vapour US₂ is produced.—4. It is generally said that U does not react with water; but, according to Moissan (C. R. 116, 347), the metal obtained by him by reducing the oxide by charcoal in an electric furnace decomposed water at the ordinary temperature; this specimen of U contained from 5 to 13.5 p.c. C.—5. U dissolves in dilute sulphuric acid, rapidly on warming, forming USO,, and giving off H; pulverulent U dissolves in hot conc. H₂SO₄, giving off SO₂; the compact metal is scarcely acted on by conc. H.SO. -6. U dissolves in hydrochloric acid, dilute or conc., rapidly on warming, forming a hyacinth-red solution of UCl_s, which goes green on shaking in air, and then contains UCl .-7. Conc. nitric acid, even when hot, scarcely acts on U that has been fused; the pulverulent metal is quickly oxidised by hot conc. HNO, Aq to U.O. -8. Several metallic salts in solution are reduced by U, with ppn. of he metals, e.g. HgNO₂, AgNO₂, CuSO₄, SnCl₂, Pl.Cl₄, AuCl₂.

Detection and Estimation. - Uranyl salts (UO2.SO4, &c.) are formed by leating uranates with acids, also by the reactions of acids with UO, and also by oxidising uranous salts (salts of UO). K4FeCyeAq gives a brownish-red pp. with uranyl salts, soluble in HClAq or in (NH₄)₂CO₂Aq; this reaction is said to be extremely delicate; H.O.Aq and alcohol give a brown pp. (Fairley, C. N. 62, 227); carbonates give yellow pps., soluble in excess, forming yellow liquids; NaOHAq or KOHAq forms yellow pps., NH, ClAq does not interfere with the ppn., but in presence of sugar or tartaric acid pps. are not produced. Uranous salts are produced by reducing uranyl salts in solution by Zn and HClAq. After reduction the solutions give green pps. with KOHAq or NaOHAq, the pps. becoming brown and then yellow in the air; with (NH₄)₂SAq green pps. are formed, quickly becoming dark green, and black on boiling; K₄FeCy_eAq produces yellow-green pps. becoming red in the air (v. Zimmermann, A. 213, 285).

U is generally estimated as UO_2 or U_3O_8 (v. Zimmermann, A. 232, 209). Solutions may be reduced by Zn and H_2SO_4Aq , and then titrated with KMnO₄Aq, the quantity of O used corresponding with the change of UO_2 to UO_3 (Z., l.c.; v. also Bělohoubek, J. pr. 99, 231).

Uranium, arsenates of. A compound, probably UO₂.As₂O₃. 4aq, is described by Rammelsberg (P. 59, 26). Werther (J. pr. 43, 321) describes UO₂H₂AsO₄. 3aq, and various double salts of this compound with arsenates of Ca, Cu, and Ne.

Uranium, bromides of. UBr₄ is formed by heating U₃O₈ mixed with C in Br vapour; and UBr₅ by heating UBr₄ in H. Zimmermann (A. 216, 5) failed to obtain a compound with more Br by heating UBr₄ with Br to 230° in a sealed tube.

URANIUM TETEABROMIDE UBr. (Uranous bromide.) Mol. w. 558. Prepared by strongly heating a mixture of U₃O₈ and sugar, mixing with sugar charcoal that has been heated to redness, placing the mixture in a tube, and heating it first in a stream of thoroughly dried CO₂, and then in Br vapour, and allowing to cool in CO₂; the Br used must be perfectly freed from Cl. HBr, and H₂O by distillation from KBr, MnO₂, and P₂O₅ (Zimmermann, A. 216, 2; Alibegoff, A. 232, 132; cf. Hermann, J. 1861. 260). UBr, forms black, fusible leaflets; volatile at red heat, giving a vapour with V.D. 281 (Z., l.c.); very hygroscopic; dissolves in water, with a hissing sound, forming a green liquid.

By evaporating a solution of UO₂.xH₂O in HBrAq, Rammelsberg (P. 59, 12) obtained green crystals of hydrated UBr₄, probably UBr₁,8H₂O.

Uranoxybromides. (Bromoxy-uranates.) UO₂Br₂2MBr or M₂UO₂Br₃. By the reaction of HBrAq with K₂UO₄ and $(NH_4)_2$ UO₄ respectively. Sendtner (4. 195, 825) obtained yellowish, hygroscopic tablets of the composition M₂UO₂Br₄.2H₂O, where M = NH₄ and K.

URANIUM TRIBROMIDE UBr. Mol. w. not determined. Obtained by Alibegoff (A. 233, 119; cf. Zimmermann, A. 216, 5) by passing dry H over molten UBr. Dark-brown needles; easily,

volatilised; not changed by continued heating in H; very hygros opic; warmed in H2S, in complete absence of nir, forms U.S.

Uranium, chlorides of. By heating U in Cl both UCl, and UCl, are produced; and by heating UCl, in H, UCl, is formed.

URANIUM PENTACHLORIDE UCla. Mol. w. not determined; formula probably molecular, from analogy of MoCl, and WCl,

Preparation .- 1. By heating U in excess of Cl.-2. A slow stream of dry Cl is passed over a mixture of charcoal and any oxide of U, or UO2Cl2, heated to dull redness in a glass tube; dark-green octahedral crystals of UCl, are formed near the heated part of the tube, and in front of these needle-shaped crystals of UCI, are deposited, mixed with a loose brown powder, which as also UCl. If the stream of Cl is rapid, most of the UCl, is produced as a light-brown, mobile powder (Roscoe, C. J. [2] 12, 933).

Properties and Reactions .- UCl, crystallises in long, dark, needle-shaped forms; the crystals are lustrous, and reflect greenish light, in transmitted light they are dark ruby red. Both the crystals and the brown powder are very hygroscopic. Dissolves in water, with a hissing sound, forming a yellow-green liquid and giving off HCl. Decomposed to UCl, and Cl when heated, either alone or in CO, or Cl; decomposition begins at 120° in CO₂, and is complete at 235°. Heating in NH, probably produces a nitride (Roscoe, Lc.; no analyses given).

Combination.—A compound UCl, PCl, is described by Cronander (J. 1873. 222) as an amorphous, yellow solid; formed by heating UO, with excess of PCl, in a sealed tube for a long time.

URANIUM TETRACHLORIDE UCI,. (Uranous chloride.) Mol. w. 380.48.

Preparation.-1. U is heated in Cl.-2. A mixture of charcoal and an oxide of U, or UO2Cl2, is heated in a slow stream of dry Cl (Péligot, A. 43, 258; cf. Roscoe, C. J. [2] 12, 933). The crystals that form nearest to the heated part of the tube are again heated in a stream of dry CO2, whereby traces of UCl, are volatilised and removed (Zimmermann, A. 216, 8).-3. A solution of ammonium uranate in HClAq is reduced by Zn, and the solution is evaporated and crystallised. Arendt a. Knop (C. C. 1857. 164) reduced, by boiling the solution for 15 min. with Cu and a little PtCl,Aq; they then filtered, saturated with H2S, filtered, boiled off H2S, and crystallised.

Properties and Reactions. - Dark - green, lustrous, regular octahedral crystals; volatilises at red heat, in complete absence of moisture, giving a red vapour (Péligot, l.c.) with V.D. 192 (Zimmermann, l.c.). Very deliquescent; fumes much in air, giving off HCl; dissolves in water, with a hissing sound, forming a green solution that is unchanged in air; gives green, amorphous UCl, by evaporation in vacuo, but is decomposed by evaporation in the air. When UCl, Aq is dropped into boiling water all the U is ppd. as UO2.xH2O. Solution of UCl, reduces salts of Au and Ag to the metals, and FeCl, to FeCl., &c. By heating in dry H one-fourth of the Cl is removed, and UCl, remains (Péligot, A. 48, 266). According to Péligot (l.c.), UCl, combines with several metallic chlorides to form double salts.

Uranoxychlorides. (Chloroxyuranates.) UO₂Cl₂·2MCl.2aq, or M₂UO₂Cl₄·2aq. Salts of this composition, M = NH₄ and K, are produced by the reaction of large excess of HClAq with uranates of NH, and K (Péligot, A. 43, 279). The K compound is also formed by acting on UO2Cl2 with excess of KClAq, and separating the KCl that crystallises with the K2UO2Cl4 mechanically (Berzelius, P. 1, 366). The K compound forms thick plates (for crystalline measurements v. de la Provostaye, A. Ch. [3] 6, 165); easily loses water; gives off HCl at a little above 100°; melts at red heat, giving off Cl, and leaving UO2 and KCl. The ammonium compound forms very deliquescent rhombohedra.

URANIUM TRICHLORIDE UCl. Mol. w. not determined. Obtained by Péligot (l.c.) by heating UCl, in dry H as long as HCl was given off. Also formed in solution by long-continued reduction of uranyl salts by Zn and HClAq (Zimmermann, A. 213, 300). A brownish-red solid; dissolves very readily in water, forming a red solution, which gives off H and becomes green (v. P., l.c.; Z., l.c.; also A. 216, 12).

Uranium, ferrocyanides of. Various compounds are described by Wyruboff (A. Ch. [5] 8, 444) and Atterberg (Bl. [2] 24, 355) as obtained by ppg. K, FeCy, Aq by solutions of salts of U. The following formulæ are given:

 U₂FeCy₆. 10aq ; (2) U₂K₂FeCy₈. 6aq; (3) $(UO_2)_8K_2(FeCy_6)_2.6aq$; (4) $(UO_2)_8K_8(FeCy_6)_4.12aq$.

Uranium, fluorides of. Only one compound, UF4, has been isolated with certainty.

URANIUM TETRAFLUORIDE UF. (Uranous fluoride.) Mol. w. not determined; formula probably molecular, from analogy of UBr, and UCl. Prepared by adding HFAq to U_sO_s, boiling, pouring off the yellow solution (which contains UO2F2), washing the green powder that remains, first on a filter and then by decantation, until free from HFAq, and drying at 100° (Smithells, C. J. 43, 125; cf. Bolton, Z. [2] 2, 353). The process of washing is very tedious, and occupies many days. UF, is a green powder; insol. in water and dilute acids; slowly dissolved by conc. acids; boiling NaOHAq produces UO2. Heated in a Pt crucible with the lid on, a small quantity of a white, bulky sublimate is obtained; this sublimate is UO₂F₂ (Smithells, l.c.); the residue in the crucible is chiefly UO₂. Heated on Pt foil, UF₄ leaves U,O, (S., l.c.).

Double salts.—Bolton (l.c.) obtained the compounds UF, KF and UF, NaF—as green powders, insoluble in water and dilute acids, giving UO2, KF (or NaF), and HF when heated to reduess, in closed vessels—by the action of formic or oxalic acid on UO₂F₂. 3KF or UO₂F₂. NaF (v. infra) in sunlight. These compounds may be called uranofluorides, and written

KUF, and NaUF,.

Uranium hexafluoride UF, was said by Ditte (C. R. 91, 115) to be formed by heating U_sO_s with HFAq, evaporating the yellow liquid, and heating the crystals of UF_s.8HF thus obtained; but Smithells (C. J. 48, 125) has shown that the substance obtained by evaporating the

solution formed by heating U.O. with HFAq is UO2F2 (v. URANIUM OXYFLUORIDE, p. 828).

Uranoxyfluorides. (Fluoxy-uranates.) These compounds may be regarded either as double salts of UO2F2 with alkali fluorides, or as salts derived from hypothetical acids H_xUO₂F The compounds are formed by adding alkali fluorides to solutions of uranous nitrate, and crystallising under different conditions (v. Bolton, Z. [2] 2, 353; Baker, C. J. 35, 763). The compounds that have been isolated belong to several series: (1) MF.UO₂F₂.xaq = MUO₂F₃.xaq, where M = Na and x = 2 and 4 (Bolton); (2) $3MF.UO_2F_2 = M_3UO_2F_3$, where $M = NH_4$ and K (Bolton; Baker); (3) $3KF.2UO_2F_2.2aq$ =K₃U₂O₄F₂, 2aq (Baker); (4) 5KF.2UO₄F₂.2aq (Baker); (5) 3BaF₂5UO₄F₂ 2aq (Bolton). Two other series were described by Ditte (C. R. 91, 166), 4MF.UO₄F₂ and 4MF.UOF₄.xaq; but the experiments of Smithelle (C. J. 42, 131) have shown that these Smithells (C. \hat{J} . 43, 131) have shown that these compounds do not exist.

Uranium, haloid compounds of. U combines directly with Br and Cl, but not with I; no iodide, indeed, has been isolated. The haloid compounds of U belong to the form UX, where X = Br or Cl; UX_4 , where X = Br, Cl or F; and UX_5 , where X = Cl. The compounds UBr_4 and UCl_4 have been gasified, and the formulæ are molecular. From the analogy of WCl, and MoCl, the formula UCl, is probably molecular; this compound decomposes to UCl, and Cl when heated, either alone or in CO2 or Cl. From the analogy of CrCl, it may be supposed that UCl, and UBr, are molecular formulæ. No haloid compounds of U have been isolated corresponding with the chlorides of Cr, Mo, and W, M_xCl_{2x}. No haloid compounds of U are known containing each more than one halogen. Various oxyhaloid compounds, chiefly UO2X2, and compounds of these with alkali haloids, are known.

Uranium, hydroxides of, v. URANIUM, OXIDES AND HYDRATED OXIDES OF (infra); also URANIUM

OXYACIDS, AND SALTS OF (p. 826).

Uranium, iodides of. No compound of U and I has been isolated. Hermann (J. 1861. 260) and Sendter (Verbind. des Uraniums, Erlangen, 1877) failed to obtain an iodide by heating a mixture of UO, and C in I vapour or in HI.

Uranium, nitride of, U₂N₄. A greyish-black powder; obtained by heating UCl₄ in NH₃, then mixing with NH₄Cl and heating for a long time in a stream of NH, (Uhrlaub, Verbind. einiger Metalle mit Stickstoff, Göttingen, 1859). Roscoe (C. J. [2] 12, 933) states that a nitride is formed by heating UCl, in NH, but no analyses are

Uranium, oxides and hydrated oxides of. When U is burnt in air the product is U,O,; by heating this oxide with reducers, or for a long time in a stream of N or CO₃, UO₂ is formed; UO₂ is produced by fusing U₃O₃ with KClO₃, also by heating UO₂(NO₃)₂; by adding H₂O₂Aq to UO₂(C₂H₃O₂)₂Aq or UO₂(NO₃)₂Aq a hydrate of UO₄ is obtained. Besides these four oxides, there intermediate between VIII. others, intermediate between UU2 and UO2, perhaps exist. No definite hydrate of UO, has been isolated; the pp. formed by adding NH₂Aq to UCl₂Aq, or to solution of a uraneus salt (U(SO₂)₂, &c.), is either a hydrate, or a

mixture of hydrates, of $\mathbf{UO_2}$. A hexahydrate of $\mathbf{U_3O_6}$ probably exists. Uranic a id is the mono-UO, dissolves in some acids hydrate of UO, to form corresponding salts; J₃O₈ is said to react with conc. H_2SO_4 or HOlAq to form both uranous salts UX_2 , and uranyl salts UO_2X , where X = a divalent acidic radiole. UO_2 interacts with several acids to form uranyl salts UO₂X¹¹; UO₃ also reacts as an acidic oxide with strong bases, forming uranates, di-uranates, &c., M₂UO₄, M₂U₂O₇, &c.

According to Guyard (Bl. [2]1, 89), two oxides having the compositions UO and U,O, exist; but the more accurate investigations of Zimmermann (A. 213, 301) showed that the substances examined by Guyard were not definite compounds. Péligot described a black oxide U₂O₅; Zimmermann (A. 232, 273) has shown that the substance is a mixture, in variable proportions, of UO2

and U.O.

URANIUM DIOXIDE UO₂. (Uranous oxide.) Mol. w. not known. This oxide was supposed to be the element U until 1840, when Péligot proved the presence of O in the substance

(A. Ch. [3] 5, 5).

Formation.—1. By heating U₃O₈ with such reducing agents as H (Arfvedson, P. 1, 245), NH₄Cl and S (Hermann, J. 1861. 258), a little C (Buchholz, Gehlen's Ann. 4, 17, 134), oxalic acid (Wertheim, J. pr. 29, 211), &c.-2. By heating U,O, for a long time in a stream of N or CO₂ (Zimmermann, A. 232, 283).—3. By strongly heating UO_2 .C₂O₄ in absence of air (Berzelius, P. 1, 359), or in a stream of H (Péligot, A. Ch. [3] 5, 5; Ebelmen, A. Ch. [3] 5, 189).

Preparation .- A mixture of UO2012 and excess of NaCl and NH Cl is strongly heated in absence of air; and the product is washed thoroughly with water, and dried (Wöhler, A. 41, 345; cf. Hillebrand, Zeit. f. anorg. Chemie, 3, 243). If air is not excluded during heating, some UO, is obtained (H., l.c.). This process gives UO2 as a black crystalline powder. In place of using UO2Cl2, a solution of ammonium uranate in HClAq may be mixed with excess of NaCl and NH₄Cl, and evaporated to dryness, and the residue heated. By fusing any oxide of U with borax for 24 hours, and washing with water and then with dilute acetic acid, or dilute HClAq, Hillebrand (l.c. p. 249) obtained jet-black octahedra of UO₂ isomorphous with ThO₂.

Properties.—Prepared by heating UO₂.C₂O₄, UO₂ is a dark-brown, pyrophoric powder; if the temperature has been kept high the oxide is lustrous, reddish-black, and not pyrophoric; prepared by heating UO₂Cl₂ with NaCl and NH₂Cl, UO₂ forms black, non-pyrophoric, octahedral crystals. Péligot gave S.G. 10.15; according to Hillebrand (l.c.), the S.G. is nearly 11. UO2 is insoluble in water, HClAq, and dilute H.SO.Aq; it dissolves in HNO.Aq, and in conc. H₂SO₂ in presence of a little water. UO₂ is oxidised to U₂O₂ by heating in air.

Reactions.-1. Heated in air or oxygen U,O, is produced; UO, burns brilliantly in the O-H flame (Clarke, G. A. 62, 853).—2. Heating in chlorine produces UO2Cl2; when mixed with carbon and heated in chlorine, UCl, and UCl, are formed. UBr, is produced by heating UO, mixed with carbon in bromine vapour.—8. U,O, is formed by heating UO, in water vapour

(Regnault, A. Ch. [2] 62, 358).—4. Heating in vapour of carbon disulphide produces U.O.S. (Hermann, J. 1861. 258; H. Rose, G. A. 73, 139). UO2 is said to be unchanged by heating in hydrogen sulphide (Arfvedson, P. 1, 245).— 5. A solution of UO₂ in conc. sulphuric acid gives green crystals of U(SO₄)₂ on evaporation.

HYDRATES OF URANIUM DIOXIDE. No definite hydrate of UO2 has been isolated; NHAq produces a reddish-brown pp. when added to UCl, Aq or solution of a uranous salt, this pp. becomes black on boiling; when washed with air-free water and dried in vacuo it forms black, coherent lumps. A black pp. is also produced by dropping UCl4Aq into boiling water. These black pps. contain UO₂; they are probably hydrates of this oxide. The name uranous hydroxide is often given to the pp. obtained by either of the processes described; the pp. dissolves in dilute H₂SO,Aq, HNO,Aq, and ĤClAq.

URANOSO-URANIC OXIDE U3O8. (Olive-green Uranyl uranate UO2.2UO3.) oxide of uranium. Mol. w. not known.

Occurrence.-In combination with SiO2, and oxides of Sb, As, Ca, Cu, Fe, Mg, &c., in pitch-

Preparation. - U, or a decomposable U compound, e.g. UO₂(NO₃)₂, is heated in the air and then in a stream of O, and is allowed to cool in O (v. Zimmermann, A. 232, 283). For a method of preparing UsOs from pitchblends v. URANIUM,

Preparation of (p. 821).

Properties. — A dark olive-green powder, sometimes almost black, but always showing a green streak when rubbed on unglazed porcelain (Z., l.c.). S.G. 7.2 (Karsten, S. 65, 394); 7.31 (Ebelmen, J. pr. 27, 385). S.H. 07979 (Donath, B. 12, 742). Insoluble in water; slightly soluble in dilute HClAq or H₂SO,Aq; soluble in HNO, Aq, also in H, SO, with a little water.

Reactions.—1. Gives off O when heated in air or in an indifferent gas; the final product of heating in N or CO₂ is UO₂ (Zimmermann, A. 232, 283).—2. Reduced to UO₂ by heating with hydrogen, carbon, sulphur, potassium, or sodium. 3. Heated in vapour of carbon disulphide U₃O₂S₄ is formed.—4. Heated with conc. sulphuric acid, or with hydrochloric acid to 180°-200°, gives both uranous and uranyl sulphates, or chlorides; reacting as UO2.2UO3 (Z., A. 232, 287).-5. Silver nitrate solution is slowly reduced to Ag (Isambert, C. R. 80, 1087).

HYDRATE OF URANOSO-URANIC OXIDE. adding NH, Aq to UCl, Aq and allowing the pp. to oxidise in the air, and drying in vacuo over H₂SO₄, Rammelsberg (P. 55, 319) obtained a greenish-black solid which may have been a hydrate of U,O,; one determination of water only is given which agrees fairly with UsOs. 6aq.

URANIUM TRIOXIDE UO. (Uranic oxide. Uranic anhydride. Uranyl oxide (UO2)O.) Mol. w. not known. This oxide is obtained by carefully heating the hydrate UO₂.H₂O (v. infra) to 800° (Ebelmen, A. Ch. [2] 5, 199); to 250°, temperature being gradually raised (Jacquelin, Ph. C. 1845. 193). The oxide is also formed by heating UO₂.CO₃.2(NH₂)₂CO₃ to 300° (E., Lc.). The preparation of UO, may be conducted by heating UO, (NO,), in a Pt basin until decomposition begins, then transferring to glass tubes and heating these in an oil-bath to 250° so long as acid fumes are given off.

UO_s is a chamois-yellow powder. S.G. 5.02 to 5.26 (Brauner a. Watts, P. M. [5] 11, 60). Ebelmen (l.c.) described UO, obtained by heating UO, CO, 2(NH,), CO, as a sealingwax-red powder. When UO, is heated to redness it gives off O and U,O, remains (E., l.c.; cf. Read, C. J. 65, 313 [1894]); the oxide seems to be somewhat volatile at the temperature of a porcelain oven (Elsner, J. 1866. 35).

UO, reacts with many acids as a basic oxide, but the salts formed are always basic salts of the forms $UX_g, 2UO_g$, where X = the radiole of a monobasic acid, and $UX_g, 2UO_g$ where X = the radiole of a dibasic acid. It is customary to write the formulæ of these basic salts as UO, XI, and UO2-XII, and to call them uranyl salts

$$\begin{split} &(\text{UO}_2 = \text{uranyl}) \; ; \; \; \text{thus} \frac{\text{U(NO}_3)_{6}, \text{2UO}_3}{3} = \text{UO}_2(\text{NO}_4)_{2}, \\ &\text{and} \; \; \frac{\text{U(SO}_4)_3, \text{2UO}_3}{3} = \text{UO}_2(\text{SO}_4). \end{split}$$

UO, behaves towards strong bases as an acidic oxide, forming uranates M2UO4, and diuranates $M_2U_2O_7$, &c.; these salts are generally formed by fusing U compounds with bases and O_7 and by ppg. uranyl salt solutions by strong bases. When KOHAq, NaOHAq, NH, Aq, BaOAq, or CaOAq is added to a solution of a uranyl salt, e.g. to UO₂(NO₃)₂Aq, the pp. consists of a uranate of the metal of the base. Alkali carbonates ppt. alkali uranates, but the pps. contain also CO2; the pps. dissolve in excess of the alkali carbonates forming double carbonates of uranyl and the alkali metal, e.g. UO2.CO3.2(NH4)2CO3. Addition of alkali or alkali carbonate to solutions of uranyl salts containing other metallic salts causes ppn. of a uranate of the metal of the salt present, generally mixed with alkali uranate (cf. Uranium oxyacios, AND SALTS THEREOF, p. 826).

By adding KOHAq to a cold solution of UCl.

or UO2(NO3)2, and dialysing, Graham (T. 1861. 213) obtained a solution, free from acid and alkali, which he regarded as a loose compound of sugar and UO, and which he called sucrate

of peroxide of uranium.

HYDRATES OF URANIUM TRIORIDE. Pure hydrates of UO, are not obtained by ppg. uranyl salt solutions by alkalis (v. supra); but according to Ebelmen (A. Ch. [3] 5, 199) UO₂.xH₂O is produced by boiling an aqueous solution of UO₂.CO₃.2(NH₂)₂CO₃ until a yellow pp. is formed, and allowing this pp. to stand for a long time. By the continued action of sunlight on UO₂(C₂O₄) in presence of water, UO3.xH2O is said to be formed; after washing and drying in the air, the solid has the composition UO₂.2H₂O; and after drying in vacuo the composition is UO₂.H₂O (E., Lc.). Berzelius (B. J. 24, 118) obtained UOs.H2O by heating UO2(NOs)2 on a sand-bath until acid fumes ceased to be given off, washing the residue with boiling water, and drying in air. The same hydrate is said to be formed by heating U₂O₂ with KClO₂ until fusion begins, boiling out with water, and drying in air (Drenkmann, J. 1861. 256). The best method of preparing UO₂.H₂O seems to be that given by Malaguti (A. Ch. [B] 9, 463), viz. boiling a solution of UO₂(NO₂)₂ in absolute alcohol as long as a reaction proceeds, washing the yellow solid that separates with water, and drying in air or in vacuo; EtNO₅, N₂O₅, C₂H₄O, and H.CO₂H are

formed in the reaction.

UO₂·H₂O is described as a yellow powder; S.G. 5·93 at 15° (Malaguti, *l.c.*); reddens litmus paper; when hot it absorbs CO₂ from the air; heated to 250°-300° water is given off and UO₂ remains (Ebelmen, *l.c.*; Jacquelin, A. Ch. [3]5, 199); heated to redness gives off O and leaves U₂O₂· UO₃·H₂O is generally called *uranic acid*; v. URANIUM OXYAGIDS, AND SALTS THEREOF (*infra*).

URANIUM PEROXIDE. By adding UO₂(NO₃)₂Aq to a mixture of H₂O₂Aq and a large excess of H₂SO₄Aq, and allowing to stand for a considerable time, Fairley (C. J. 31, 133 [1877]) obtained a small quantity of a heavy, crystalline, almost white pp., which when dried (? in air, or at 100°) had the composition UO₄. By adding dilute H₂O₂Aq to solution of UO₂(NO₃)₂ or UO₂(C₂H₂O₂)₂, free from acids, Fairley (l.c. p. 127) obtained a yellowish-white pp., which when dried by pressure between filter paper had the composition UO₄.4H₂O, and when dried at 100° the composition UO₄.2H₂O.

Zimmermann (A. 232, 273 [1886]) failed to obtain the tetrahydrate UO, 4H₂O; according to him the dihydrate UO, 2H₂O is very hygroscopic, and begins to decompose at 115°. Alibegoff also obtained only UO, 2H₂O; he failed to prepare the anhydrous oxide UO₄ (A. 233, 123

[1886]).

Hydrated uranium peroxide, dried at 100°, is a yellowish-white powder; decomposed by heat, giving off O and leaving much U₂O₅; reacts with alkali solutions to give UO₂.xH₂O, and as solution from which alkali peruranate 2M₂O.UO₆. Saq, crystallises (Fairley, L.c.; v. Peruranates under URANIUM OYYACIDS, AND SALTS

тневеог, р. 827).

Uranium oxyacids, and salts thereof. UO₂.H₂O₃ = H₂UO₄ reacts with strong bases as an acid, forming uranates M^r₂UO₄; di-uranates M^r₂U₂O₇ are also known, and a few tri- and hexa-uranates. A few peruranates are derived from hydrated uranium peroxide UO₄.xH₂O. The compounds of UO₂X₂ with MX (X = Br, Cl, or F) may be regarded as bromoxy-, chloroxy-, and fluoxy-uranates. The substance known as uranium red is perhaps UO₂.ONH₄.SNH₄, i.e. the ammonium salt of an oxythio-uranic acid (v. URANIUM OXYBULPHDES, D. 828).

(v. Uranium Oxysulphides, p. 828).

Uranic acid H₂UO₄. This compound is obtained as a yellow powder, which reddens litmus paper, and is decomposed to UO₂ and H₂O at 250°-300°, and to U₃O₃ and O at a higher temperature, by boiling a solution of UO₂(NO₃)₂ in absolute alcohol, washing the solid that separates with water, and drying in air or in vacuo. (For other methods of preparation, and more details v. Hydrates of Uranium Trioxide, p. 825.) The salts of H₂UO₄ are prepared indi-

rectly.

URANATES, DI-URANATES, &c. Salts derived from H₂UO₄; salts derived from the hypothetical acids H₂U₂O₇, H₂U₂O₁₀, H₂U₄O₁₉ are also

knowr

URANATES. Salts of the composition $M_2^1UO_4$ and $M_1^1UO_4$. These salts are prepared by ppg. solutions of uranyl salts by bases in solution, or by adding alkali solution to a mix-

ture of a uranyl salt and a metallic oxide in solution; also, in many cases, by oxidising U_1O_0 or UO_2 in presence of a base or a salt of a base. The uranates are insoluble in water, and are not readily decomposed by heat except the ammonium salt. Carson a. Norton $(Am. \ 10, 219)$ have pointed out that analyses of uranates always show considerable differences between the percentages of U found and calculated; they have found the same anomaly in analyses of uranates of ammonium and several ammonium derivatives, such as NEtH. NEt. H. &c.

derivatives, such as NEtH₄, NEt₂H₂, &c.

Ammonium uranates. The pure salt does not seem to have been isolated. An impure uranate containing c. 90 p.c. UO₃ ((NH₁)₂UO₄ requires 84·66 p.c. UO₃) is obtained by adding NH₃Aq to a solution of a uranyl salt (Péligot); it is a yellow powder, which loses H₂O and NH₄ above 100°, very slightly soluble in water, and insol. NH₃Aq (Arfvedson, P. 1, 245). An impure salt is also obtained by adding NH₄ClAq or (NH₄)₂SO₄Aq to boiling Na₂UO₃Aq, as long as NH₃ and CO₂ are given off, washing the pp., and drying at a low temperature (Anthon, D. P. J. 156, 211). Heated to redness gives U₃O₄; many of the compounds of U are prepared from ammonium uranate.

Potassium uranate K₂UO₄. Prepared by heating 6 g. U₃O₈ with HClAq and HNO₃Aq, adding 4 g. KCl and 16 g. NH₄Cl, evaporating to dryness, heating very strongly until the fused mass is orange-yellow, and washing with water (Zimmermann, A. 213, 290). The salt is said to be formed also by strongly heating UO₂HPO₄ (obtained by ppg. a uranyl salt by a phosphate) with K₂SO₄ (Grandeau, A. Ch. [6] 8, 223). Orange-yellow, rhombic tablets; insol. water, cold or hot; easily sol. acids (Zimmermann, Lc.).

Sodium uranate Na, UO4. Prepared by fusing U2O8 with NaCl and Na, CO3, or with NaCl with gradual addition of NaClO3, and washing with water; resembles K2UO4 (Zimmermann,

Z.c.)

Lithium uranate Li₂UO₄ has been prepared similarly to K₂UO₄; it is said to be decomposed by hot water (Z., l.c.). Uranates of Ba, Ca, and Sr—MUO₄—are said by Ditte (C. R. 95, 988) to be formed by heating the chlorides of these metals with U₂O₈.

DI-URANATES. Salts of the composition $M_2^{\text{I}}U_2O_7$ and $M^{\text{II}}U_2O_7$. The formulæ may also

be written M'2UO4.UO3 and MIIUO4.UO3.

Barium di-uranate BaU₂O₇. Formed by ppg. UO₂(C₂H₂O₂)₂Aq by BaOAq (Berzelius, P. 1, 359); also by heating to redness UO₂.Ba(C₂H₂O₂)₄ (Wertheim, J. pr. 29, 207). A yellowish-red powder.

Calcium di-uranate CaU₂O₂. Formed, according to Ditte (C. R. 95, 988), by strongly heating U₃O₈ with Ca(ClO₃)₂; a yellowish-green solid.

Copper di-uranate CuU₂O₂. This salt is said to be formed, as a green crystalline powder, by fusing Cu(UO₂.PO₄)₂. Eaq with Na₂CO₃, and washing with water (Debray, A. Ch. [3] 61, 451).

Lead di-uranate PbU₂O₇. Formed by digesting UO₂(C₂H₂O₂)₂Aq with freshly ppd. PbCO₃ (Wertheim, J. pr. 29, 207); also by adding NH₂Aq to a mixture of UO₂(NO₃)₂Aq and Pb(NO₃)₂Aq (Arfvedson, P. 1, 258). A yellowish.

red powder, becoming green when heated; strongly heated in H, gives a pyrophoric mixture of Pb and UO2.

Potassium di-uranate K₂U₂O₇. Formed by melting together UO₃ and K₂CO₃ and washing with water; also by strongly heating $UO_2.K_2(CO_3)_2$ or $UO_2.K(C_2H_2O_2)_3$ (Berzelius, P. 1, 359). A

yellowish-red powder.

Silver di-uranate Ag₂U₂O₇. Obtained by adding amorphous K2UO4 to molten AgNO3, and washing with ice-cold water (Alibegoff, A. 232, 129; 233, 117, 143). Also by strongly heating UO₂.Ag(C₂H₂O₂)₃ (Wertheim, J. pr. 29, 221; of. Rammelsberg, P. 59, 10). An amorphous, yellowred solid; easily sol. acids; decomposed by hot water (Alibegoff, l.c.).

Sodium di-uranate Na₂U₂O₇. 6aq. Obtained by ppg. uranyl salts by NaOHAq (Patera, J. pr. 51, 125; cf. Stolba, Fr. 3, 74). Loses water of crystallisation over HSQ. A light-vellow to crystallisation over H.SO. A light-yellow to darkish-yellow powder. Known commercially as uranium yellow (v. Dictionary of Applied

CHEMISTRY, vol. iii. p. 893).

Di-uranate of strontium SrU2O, is said also

to exist (Ditte, C. R. 95, 988).

TRI-URANATES. The only salt that has been isolated is sodium tri-uranate Na₂U₂O₁₀; it is prepared by fusing UO2(SO4) with NaCl, and boiling out with water. This salt forms yellow leaflets resembling mosaic gold; S.G. 6.912; hygroscopic; insol. water, sol. acids (Drenkmann, J. 1861, 255).

HEXA-URANATES. The potassium salt, K₂U₆O₁₉. 6aq, is described as a yellow microscopically crystalline powder, which loses all water at 300°-400°; prepared by melting $UO_2(SO_4)$ with KCl (Drenkmann, J. 1861. 255).

PER-URANATES. By adding excess alkali and H2O2Aq to solutions of uranyl salts, Fairley (C. J. 31, 134) obtained salts which may be formulated as MI4UOs. xaq and MI2RIIUOs. xaq where $M = NH_4$, K, or Na, and $R = UO_2$.

Ammonium-uranyl peruranate (NH₄)₂(UO₂)UO₈. Saq. The formula may also be written (NH₄)₂O₄O₂O₈. Saq, or

(NH₄)₂O.UO₃.UO₄. 8aq, or, regarding the compound as a double peroxide of U and NH₄, as 2UO₄.(NH₄)₂O₂.8aq. The salt is prepared by adding excess of NH₂Aq and H₂O₂Aq to a solution of a uranyl salt and ppg. by alcohol; it is an orange-yellow solid, readily soluble in water. The quantity of KMnO4 in solution decolourised shows that the ratio of U to 'peroxide oxygen' in the salt is U:30.

Potassium per-uranate K₄UO₈. 10aq. salt may also be looked on as a double peroxide of U and K (UO4.K4O4.10aq), or as 2K2O.UO6.10aq, It is prepared by adding alcohol to a solution obtained by treating UO₂(NO₂)₂Aq with KOHAq and excess of H2O2Aq, and rapidly drying the orange-yellow pp. by pressure in filter paper. The salt rapidly absorbs CO₂ from the air and loses O; it is easily decomposed by heat, giving off H2O and O. Experiments showed that the

ratio of U to 'peroxide oxygen' is U:30.
sodium peruranate Na₄UO₆. 8aq. The
formula may also be written UO₄.Na₄O₄. 8aq or
2Na₄O.UO₆. 8aq. The salt is prepared by dissolving 'ordinary uranic hydrate,' or UO4. xaq, in excess of NaOHAq along with excess of H₂O₂Aq, allowing to crystallise if conc. solutions

are used, or ppg. by a little alcohol if dilute solutions are used, and drying by pressure in filter paper. Poleck (B. 27, 1051 [1894]) obtained this salt by adding sodium superoxide to solution of a salt of U. The salt is golden-yellow and lustrous; it slowly absorbs CO₂ from the air, and gives off O. When heated, alone or in CO₂, it gives off 30; it also loses 30 in contact with acidified KMnO, Aq or other unstable O compounds. The ratio of U to 'peroxide oxygen' is U:3O as measured by the quantity of KMnO. decolourised by a solution of the salt.

Sodium-uranyl per-uranate

Na₂(UO₂)UO₃. 6aq. The formula may also be written Na₂O.U₂O₃. 6aq or Na₂O.UO₃.UO₆. 6aq; or, regarding the compound as a double peroxide of U and Na, as 2UO4.Na2O2. 6aq. The salt is prepared similarly to Na, UO, 8aq, but using only the minimum quantity of NaOHAq required for solution, and adding alcohol, when it separates as a red oil which slowly crystallises to a red solid. The quantity of KMnO, in solution decolourised by a solution of the salt showed that the ratio of U to 'peroxide oxygen' is U:30.

Uranium, oxybromide of, UO2Br2. (Uranyl bromide.) In preparing UBr, by heating a mixture of U2O8 and C in Br vapour, Hermann (J. 1861. 260) obtained a yellow, easily volatile sublimate, which was probably an oxybromide. By heating UO₂ with Br and water, also by dissolving UOs in HBrAq, a colourless liquid is obtained, which on evaporation gives yellow needles of the hydrated oxybromide UO₂Br₂, 7aq (Sendtner, A. 195, 325). This compound combines with NH Br and KBr to form UO2Br.2MBr (v. Uranoxybromides, p. 822).

Uranium, oxychloride of, UO2Cl2. (Uranyl chloride.) Formed by strongly heating UO2 in Cl; a yellow, crystalline solid, easily fused, but volatilised only at a high temperature; heated with K forms UO₂ and KCl (Péligot, A. Ch. [3] 5, 5). By dissolving UO₃.H₂O in HClAq, also by oxidising UCl₄ by HNO₃Aq, and evaporating, yellowish-green crystals of UO₂Cl₂.xH₂O are observed to the constant of UO₂Cl₂.xH₂O are tained (Klaproth, Crell's A. 1789 [2]. 387).

By dissolving UO2Cl2in ether and evaporating, Regelsberger (A. 227, 119) obtained yellow needles of the compound UO₂Cl₂.2Et₂O.

UO2Cl2 forms compounds with NH4Cl and KCl of the form 2MCl.UO2Cl2 (v. URANOXYCHLORIDES,

p. 823).

 UO_2Cl_2 combines with NH, to form $UO_2Cl_2.xNH_3$, where x=2, 3, and 4. The first of these compounds, $UO_2Cl_2.2NH_3$, is formed by passing dry NH3 into UO2Cl2 dissolved in ether, drying the voluminous yellow pp. that forms, and then placing it in vacuo over H2SO4 until all ether is removed; the second compound, UO₂Cl₂.3NH₃, is formed by the action of NH, on UO₂Cl₂.2NH₃; and a mixture of UO₂Cl₂.3NH₃ and UO Cl₂4NH, is produced by passing NH, over solid UO₂Cl₂. When strongly heated in air these compounds give off all NH, and Cl and leave U₁O₂; heated out of contact with air they give UO₂ (Regelsberger, A. 227, 119). The three compounds are called by R. uranylammonium and their formulæ are written l)2, UO2(NH,Cl)(NH,NH,Cl), and chlorides, UO₂(NH₂Cl)₂, UC UO₂(NH₂NH₂Cl)₂. Water reacts thus:-(1) 3UO₂(NH₂Cl)₂ + 3H₂O = (NH₂)₂U₂O₇ + UO₂Cl₂ + 4NH₄Cl₃

(2) $UO_2(NH_2Cl)_2 + 2H_2O = UO_2(OH)_2 + 2NH_4Ol$; the second reaction takes place quickly with

Uranium, oxyfluoride of, UO2F2. (Uranyl fluoride.) This compound is produced by boiling U₃O₈ with HFAq, filtering off UF₄ (v. URANIUM TETRAFLUORIDE, p. 823), and evaporating the filtrate; it is also formed, in small quantities, by heating UF, in a closed Pt crucible. The compound differs according to the method of preparation (Smithells, C. J. 43, 125).

a-Oxyfluoride. About 1 gram UF, is heated in a Pt crucible with the lid on, temperature being raised as quickly as possible; after five minutes the crucible is allowed to cool, the light, bulky sublimate is removed, and the crucible is again heated for five minutes, and these processes are repeated several times (S., l.c. p. 129). This form of UO₂F₂ is a very bulky, light, white, crystalline solid; it is very soluble in water, forming a yellow liquid; heated in air it is

gradually changed to U,O,.

β-Oxyfluoride. U₃O₈ is treated with boiling HFAq in a leaden vessel, the yellow solution is poured off from green UF4, heated until the excess of HF is driven off, and allowed to evaporate; evaporation may be completed over H.SO, and CaO in vacuo; the residue is dried at 100° (S., l.c. p. 130). This form of UO,F, is a yellow saponaceous mass; in one instance the preparation had a semi-crystalline appearance, forming yellow, lustrous scales; it is very soluble in water or alcohol; heated on Pt foil U2O8 is formed; heated in a closed crucible it is slowly decomposed, giving UO₂. \$\textit{\beta}\$-UO₂F₂ probably forms a compound with HF; by β -UO₂F₂ evaporating a solution of U₂O₈ in HFAq over H₂SO₄ and CaO until the weight was constant, Smithells (l.c. p. 131) obtained a substance which gave results on analyses agreeing fairly with the formula UO₂F₂·HF·H₂O.

A solution in water of either form of UO₂F₂

mixed with KFAq and KHF2Aq, and evaporated, gives the quadratic fluoxy-uranate K, UO,F, (S., i.c. p. 130-1). For the compounds of UO₂F₂

with MF v. URANOXYFLUORIDES (p. 824).

Ditte (C. R. 91, 115) gave the formula UOF to the sublimate obtained by heating, in a closed Pt crucible, the solid that remained when U,O, was treated with boiling HFAq. Ditte said that this solid was UO₂F₂; Smithells (l.c.) has, however, shown that the solid obtained by the method used by Ditte is UF, and that the sublimate formed by heating this is a-UO₂F₂.
Uranium, oxysulphides of. Two compounds

probably exist.

URANIUM OXYSULPHIDE U₃O₂S₄. This composition was given by Hermann (J. 1861. 258) to a dark, greyish-black solid, obtained by heating UO., U.O., or NH, uranate to redness in vapour of CS. Dissolves in conc. HClAq; burns when heated in air (cf. H. Rose, G. A. 73, 139).

URANYL SULPHIDE (UO2)S. A brown solid, obtained by ppg. an ammoniacal solution of a uranyl salt by cold (NH₄)₂SAq, and quickly washing with conc. alcohol. Probably not obtained pure, as it quickly decomposes; water produces UO₂.wH₂O; at 40°-50° it is changed to UO₂ and S (Remelé, P. 124, 114; Zimmermann, A. 204, 204). (UO2)S is somewhat sol. water or dilute alcohol: dissolved by conc. HClAq, giving UCl,

and S; dissolved by (NH₄), SAq, only in presence of (NH₄)₂CO₂Aq (Zimmermann, l.c.), forming a

brown liquid.

Uranium black. By allowing (UO2)S to stand in contact with freshly prepared (NH.) SAq, in absence of air, a dull black amorphous powder is obtained. This solid dissolves in mineral acids, giving off traces of H₂S; it dissolves partly in warm (NH₄)₂CO₃Aq, but not in warm (NH4)2SAq, nor in caustic alkali solutions; heated in a tube to c. 270° it gives off a very little NH₃ and H₂O. Zimmermann (A. 204, 204) gives the formula U,O,0 to uranium black, neglecting (as impurities) the traces of NH, and

S which the substance contains.

Uranium red. By allowing (UO₂)S to stand for 24-48 hours in contact with cold (NH₄)₂SAq, exposed to air, a deep-red solid is obtained; soluble in dilute mineral acids, with separation of S and evolution of H2S; gives off NH_a, S, and H₂O when heated to 150° and, at above 200°, leaves U₃O₈ (Zimmermann, l.c.). According to Z., the formation of this red substance takes place only when $(NH_1)_2S_2O_3$ is present in the $(NH_4)_2SAq$ used; $(NH_4)_2S_2O_3$ is formed by the action of the air. 7. gives to uranium red the composition U,S(NH₄)2O, = UO.ONH.SNH.2U20. (cf. Patera, J. pr. 51, 122; Remelé, P. 124, 158; Hermann, J. 1861.

Uranium, salts of. Two classes of salts are formed by treating oxides of U with acids; UO, forms corresponding salts, UX2 when X = the radicle of a dibasic acid, and UX, when X = the radicle of a monobasic acid; UO₃ forms salts $(UO_2)X$ when X =the radicle of a dibasic acid, and $^2(UQ_2)X_2$ when X = the radicle of a monobasic acid. The salts UX^1 , and UX^{11}_2 are called uranous salts; and $(UQ_2)X^1_2$ and $(UQ_2)X^{11}$ are called uranyl salts. (For reactions of the two classes of salts v. Uranium, Detection and esti-mation of, p. 822.) Besides the normal salts, a large number of basic salts of both classes is known, and also many double salts.

The chief uranous salts are arsenates, phosphates, and sulphites. The chief urany i salts are arsenates, bromate, double carbonates, chromates, hypophosphite, iodate, nitrates, oxalate, phosphates, phosphite, selenates, selenites, sulphates, and sulphites (v. NITRATES, SUL-

PHATES, &c.).

Uranium, sulphides of. When U is burnt in vapour of S, the disulphide US2 is formed; by passing H2S over heated UBr3 the sesquisulphide U2S, is produced; and by heating U2S, in H the monosulphide US is obtained. The

mol. w. of none of these is known.

URANIUM DISULPHIDE US. Prepared by heating U in vapour of S (Péligot, P. 54, 122; cf. Zimmermann, A. 216, 18); also by heating UCl, to redness in H2S (Hermann, J. 1861. 258). A dark greyish-black powder; becomes crystalline when fused with borax; oxidises slowly in air, rapidly on heating. Easily soluble in conc. HClAq (Hermann, l.c.).

URANIUM SESQUISULPHIDE U2S. A stream of dry H2S, perfectly free from air, is passed over heated UBr, as long as HBr is given off. Greyblack solid, forming pseudomorphs of UBr.; decomposes in air, giving of H.S.; burns when heated in air; scarcely acted on by HClAq or

dilute HNO, Aq; decomposed violently by conc. HNO, (Alibegoff, A. 233, 117)

URANIUM MONOSULPHIDE US. A black, amorphous powder; obtained by passing dry H over U.S., kept at a red heat, for c. 40 hours, until H.S is no longer given off. Similar in properties to U2S3 (Alibegoff, A. 233, 135).

Uranium thio-acids, salts of. No thio- acid, nor salt of a thio- acid, of U has been definitely isolated. Uranium red (v. supra, p. 828) may perhaps be an ammonium oxythionate of U.

M. M. P. M. URANOXYBROMIDES p. 822, under URANIUM TETRABROMIDE

URANOXYCHLOP.IDES v. p. 823, under URANIUM TETRACHLORIDE

URANOXYFLUORIDES v. p. 824, under

URANIUM TETRAFLUORIDE.

URANYL COMPOUNDS, compounds of the radicle UO2; v. URANYL AMMONIUM CHLORIDES (p. 827), URANYL BROMIDE (p. 827), URANYL CHLORIDE (p. 827), URANYL FLUORIDE (p. 828), URANYL SALTS (p. 828), URANYL SULPHIDE (p.

URAZOLE. A name given by Pinner (B. 20, 2358) to di-oxy-triazole, which may be represented as NH.C(OH) N or NH.CO NH.CO NH.CO

UREA CH₁N₂O₁, i.e. CO(NH₂)₂. Carbamide. Mol. w. 60. [132°] (Lubavin, B. 3, 305). S.G. 1·323 (Schröder, B. 12, 562). S. (alcohol) 20 in the cold; 100 at 78°. H.C.v. 152,500. H.C.p. 152,200 (Stohmann a. Langbein, J. pr. [2] 44, 387); 160,900 (Berthelot a. Petit, C. R. 109,

759; 110, 887). H.F. 80,800.

Occurrence.—In the urine of mammalia, especially in that of flesh-eaters (Fourcroy a. Vauquelin, A. D. 1799). Occurs in small quantity in blood, muscle, chyle, and lymph of mammalia (Wurtz, C. R. 49, 52; Poisseuille a. Gobley, J. 1859, 612; Verdeil a. Dollfus, A. 74, 214; Munk, Pf. 11, 100; Pickard, C. R. 83, 1179; 87, 533; Gréhant a. Quinquand, C. R. 108, 1092; Garrod, Pr. 53, 478). Constitutes about 30 p.c. of the solid substance of the vitreous humour of the eye (Millon, C. R. 26, 119; A. 66, 128). Occurs also in saliva (Rabuteau, J. 1873, 877), in cow's milk (Lefort, Z. 1866, 190; Vogel, J. 1867, 932), and in other animal secretions.

Formation.—1. By evaporating a solution of ammonium cyanate (Liebig a. Wöhler [1828], B. J. 12, 266; P. 12, 253; 15, 619; A. 38, 108). 2. By adding a little NHO, to an ethereal solution of cyanamide (Cannizzaro a. Cloëz, A. 78, 230) or by the action of dilute (50 p.c.) H2SO4 on cyanamide (Baumann, B. 6, 1373).—3. From COCl. and dry NH, (Regnault, A. Ch. [2] 69, 180; Natanson, A. 98, 287), guanidine and NH,Cl being formed at the same time (Fenton, C. J. 35, 793).—4. By heating carbonic ether with NH₃Aq at 180°.—5. By heating ammonium carbamate at 140° (Basaroff, J. pr. [2] 1, 283).-6. By passing an alternating electric current through a solution of ammonium carbamate (Drechsel, J. pr. [2] 22, 481).-7. By electrolysis of ammonia solution, using carbon electrodes (Millot, Bl. [2] 46, 243).—8. By passing a mixture of NH₃ and CO₂ through a red-hot tube (Dexter, Am. 4, 35).—9. Formed by passing air charged with NH₃ and vapour of benzene over a red hot helix of platinum wire (Herroun, C. J. 39, 471). Acetylene, but not ethylene, may be substituted for benzene.—10. By heating ammonium thiocarbamate NH2.CO.SNH4, or by shaking it with water and PbCO3.-11. By the action of aqueous KMnO, on thio-urea (Maly, M. 11, 277).-12. By the action of KMnO, on KCy in acid solution (Baudrimont, J. 1880, 393) .-13. By the action of NH, Aq and H2S on copper fulminate (Gladstone, A. 66, 2).—14. By heating oxamide with HgO (Williamson, Mémoires du Congrès scientif. de Venise, A.D. 1847).—15. By oxidation of uric acid.-16. By the action of KClO, and HCl on guanine (Strecker, A. 118, 159).—17. By boiling guanidine with baryta (Baumann, B. 6, 1376).—18. By boiling biguanide sulphate or phenyl-biguanide sulphate with baryta (Emich, M. 12, 11).-19. By the action of baryta on arginin (Schulze a. Likiernik, B. 24, 2701).-20. By boiling creatin with baryta.-21. A product of oxidation of proteids by KMnO₄ (Béchamp, A. Ch. [3] 48, 348; C. R. 70, 866; Ritter, Bl. [2] 16, 32). Staedeler (J. pr. 72, 251) and Loew (J. pr. [2] 2, 289) failed to obtain urea in this manner. Dreschel (B. 23, 3097) obtained urea from albumen by electrolysis, but not by oxidation.

Preparation. - 1. Urine, concentrated by evaporation, is treated with nitric or oxalic acid, and the ppd. nitrate or oxalate of urea is decomposed by CaCO₂.—2. A solution of potassium cyanate mixed with ammonium sulphate is evaporated on a water-bath to dryness, and the residue extracted with alcohol.—3. A mixture of dry K4FeCy6 (28 pts.) and MnO2 (14 pts.) is heated till sticky on an iron plate. A. cold. aqueous extract of the mass is mixed with (NH₄)₂SO₄ (20.5 pts.), evaporated to dryness, and extracted with alcohol (Liebig).-4. Dry K, FeCy, (8 pts.) is fused with K₂CO₃ (3 pts.) and PbO (15 pts.) added to the melted mass. The cooled mixture is dissolved in water, mixed with in the basel of the water, in of (NH₄)₂SO₄ on KCyO obtained by heating dry K, FeCy, with K, Cr, O,; the yield being 25 p.c. of the weight of ferrocyanide (C. A. Bell, C. N. 32, 99).—8. A stream of dry NH, gas is passed through phenyl-carbonate (from phenol-sodium and COCL), melted in a water-bath, and the melt is then poured into hot water; the aqueous solution after separation from the phenol is evaporated to crystallisation (Hentschel, B. 17. 1286).

Properties. - Dimetric crystals, v. e. sol. water, m. sol. cold alcohol, insol. CHCl, nearly insol. ether. Sublimes at 130°. In vacuo it distils at 135° (Bourgeois, Bl. [3] 7, 45). Urea is not attacked by pure HNO, (Franchimont, R. T. C. 6, 217).

Reactions.-1. When heated alone at 150°-170° it yields NH3, cyanuric acid and biuret; at 140° it yields ammelide.—2. Decomposed into CO₂ and NH, by heating with water in sealed tubes above 100°, by boiling with potash, lime, or magnesia, by warming with conc. H2SO4, by evaporating with lead acetate, and by the action of ozone (Gorup-Besanez, A. 125, 207).

in presence of free alkali. CaO produces cyanamide (Emich, M. 10, 330). A 3 p.c. HCl solution decomposes 4 p.c. of the urea in 24 hours. NaOHAq of the same strength has less effect. Cold water does not decompose urea (Berthelot a. André, Bl. [2] 47, 840).—3. Alcoholic potash forms, on heating in sealed tubes, cyanic acid, NH_s, and water (Haller, Bl. [2] 45, 705).-4. Urea hydrochloride at 145° yields NH Cl and cyanuric acid (De Vry, A. 61, 249).-5. Chlorine passed over melted urea forms HCl, nitrogen, NH,Cl, and cyanuric acid (Wurtz, A. 64, 307).— 6. Hypochlorous acid, hypochlorites, chlorinewater, and hypobromites, set free nitrogen, e.g. $CON_2H_4 + 3HOCl = CO_2 + 3HCl + 2H_2O + N_2$ 7. Nitrous acid added to a hot solution forms CO_2 and nitrogen: $CON_2H_4 + N_2O_3 = CO_2 + 2H_2O + 2N_2$. Nitrous acid (2 mols.) when added to a cold solution of urea (1 mol.) reacts as follows: $2CON_2H_4 + N_2O_3 = (NH_4)_2CO_3 + 2N_2 + CO_2$ (Claus, B. 4, 140).—8. Neutral KMnO, has no action in the cold, and acts but slowly at 100°. In acid solutions KMnO₄ gives off nitrogen (1 vol.) and CO₂ (2 vols.) (Béchamp, J. 1856, 696; cf. Wanklyn a. Gamgee, C. J. 21, 25).—9. Bromine added gradually to an alcoholic solution of urea forms NH,Br, cyanuric acid, and nitrogen. In sealed tubes Br gives NH,Br and cyamelide (Smolka, M. 8, 64).—10. Heating with P₂O₅ forms cyanic and cyanuric acids, ammelide, cyamelide, and NH₃ (Weltzien, A. 107, 219).—11. Yields cyanamide on warming with Na (Fenton, C. J. 41, 262).—12. Conc. HNO₃ reacts thus: CON₂H₄ + 2HNO₃ = CO₂ + N₂O + NH₄NO₃ + H₂O (Franchimont, R. T. C. 2, 96; 3, 219).—13. P₂S₃ forms crystalline C₂H₁N₂S₂O₂, which yields CH A₂N₂S₂O₃ and CH A₂N₃S₂O₃ and CH A₂N₃S₃O₃ and CH A₃N₃S₃O₃ and C2H,AgN3PS2O2 and C2H3Ag3N3PS2O2, and gives off PH, on warming (Kutschig, M. 9, 406).-14. Zinc-dust at a red heat forms ZnCy2, ammonia, and hydrogen (Aufschläger, M. 13, 272). 15. CS₂ at 100° forms COS and ammonium sulphocyanide. With alcohol and CS2 the products are mercaptan, CO2, and ammonium sulphocyanide (Ladenburg, Z. [2] 5, 253; B. 1, 273; 2, 271; Fleury, A. 123, 144).—16. Alcohols in sealed tubes form carbanic ethers and alkyl allophanates (Cahours, C. R. 76, 1387).—17. Boiling Ac O forms acetamide and diacetamide (Hofmann, B.14, 2733).—18. ZnEt, forms CON, H, Zn, which is reconverted into urea by water (Gal, Bl. [2] 39, 648).—19. Oxalic ether at 135°-170° forms oxamide, allophanic ether, and alcohol (Grabowski, A. 134, 115). Oxamic ether forms, on fusion, oxaluramide (Carstanjen, J. pr. [2] 9, 143).--20. Methylamido-acetic acid (sarcosine) forms methyl-hydantoïn on fusion (Huppert, B. 6, 1278).—21. Aniline at 150°-170° gives NH. and di-phenyl-urea (Baeyer, A. 131, 251).— 22. Aniline sulphonic acid at 120° gives NH_CO.NH.C₈H₄.SO₅H (Ville, Bl. [3] 6, 6).— 23. Biguanide sulphate forms, on heating, ammeline, NH₄ and (NH₄)₂SO₄ (Smolka a. Friedreich, M. 10, 95).—24. Heated with acetone and ZnCl₂ or P₂O₅ at 110°-140° it gives s-t.:-methylpyridine and a base C_{1e}H₁₀N [119°]; (320°) (Riehm, A. 238, 22). This base forms hard monoclinic crystals, with metallic ring; its solutions fluoresce. It gives B'aHaPtOla: [225°].— 25. Heated with acetyl-acetone (1 mol.), ures (2 mols.) forms CH₂(CMe:N.CO.NH₂)₂, which crystallises from alcohol [200°] (Combes, Bl. [3]

The same body is formed, together **7**, 790). with CH₂ CMe:N CO [198°], by adding HClAq or H2SO, to an alcoholic solution of urea and acetyl-acetone (Evans, J. pr. [2] 46, 352; 48, 499). — 26. Benzoyl-acetone forms, in like manner, Bz.CH2.CMe:N.CC.NH2 [191°] and CH₂ CMe:N CO [227°] (Evans, J. pr. [2] 46, 352; 48, 509).—27. Acetoacetic ether (2 mols.) and NaOEt (2 mols.) form in the cold C₁₈H₂₂N₂O₇Na₂ [c. 165°] (Ernert, A. 258, 361). Acetoacetic ether in alcoholic solution forms uramido-crotonic acid (q. v.).—28. Urea reacts with aldehydes in In aqueous solution a diureïde the cold. R.CH(NH.CO.NH₂)₂ is usually formed, while solid urea forms triureïdes. On warming with excess of aldehyde, di- and tri-ureïdes yield tetraand hexa-ureïdes (H. Schiff, A. 151, 186). These ureides are sparingly soluble neutral solids, and are split up by warming with acids into the parent substances. Benzoic aldehyde forms benzylidene-di-urea (q.v.), while o-oxy-benzoic aldehyde forms C₀H₄(OH).CH(NH.CO.NH₂)₂aq, which is converted by acetoacetic ether in alcohol into C₅H₄(OH).CH:N.CO.N:CMe.CH₂.CO₂Et [200°], C_EH₄(OH).CH:N.CO.NH.CMe:CH.CO₂Et [204°], and C_eH. CH:N.CO NH, which decomposes between 260° and 270° (Biginelli, B. 24, 2963). Acrolein forms (CON₂H₄)₂3C₅H₄O, which chars at 250° (Lüdy, M. 10, 295). o-Nitrobenzoic aldehyde forms nitro-benzylidene-di-urea C₆H₄(NO₂).CH(NH.CO.NH₂)₂ aq [200°]. In like manner cuminic aldehyde in alcohol forms the di-ureïde C₃H₇.C₆H₄.CH(NH.CO.NH₂)₂ [176°] (Biginelli, B. 24, 2964), and cinnamic aldehyde forms Ph.CH:CH.CH(NH.CO.NH2); [172°], and compounds melting at 116° and 212°. 29. Chloral added to conc. aqueous solutions of urea forms (C₂HCl₃O)CON₂H₄ [150°], v. e. sol. hot water, and (C₂HCl₃O)₂CON₂H₄ [190°] nearly insol. hot water, both compounds being crystalline (Jacobsen, A. 157, 246). Chloral cyanhydrin at 105° forms the very stable crystalline C₃H₅Cl₂N₂O (Pinner a. Fuchs, B. 10, 1069).— 30. Benzene sulphonic chloride at 100° forms colourless crystals of C₆H₅SON₄H₇C₂O₂ aq (Elander, Bl. [2] 34, 207). Naphthalene (a)-sulphonic chloride acts similarly .- 31. Cinnamoyl chloride forms C₉H₈(NH.CO.NH.C₉H₈.NH.CO.NH₂)₂[184°] (Biginelli, B. 24, 2965).—32. ClCH₂.OH forms methylene-urea, which is split up by hot dilute H₂SO₄ into formic aldehyde and urea (Von Hemmelmayr, M. 12, 89).—33. Trichlorolactic acid and a little water form, on heating, acetyleneurea $C_4H_sN_4O_2$ (Pinner, B. 17, 1997).—34. Hexachloro-acetone (2 mols.) heated with urea (1 mol.) at 150° forms CO:N₂H₂(C₂Cl₂O)₂, crystallising from alcohol in hexagonal plates (Cloëz, A. Ch. [6] 9, 145).

Detection.—Urea gives, in conc. solution, crystalline pps. with nitric and oxalic acids. Urea dissolved in amyl alcohol is completely ppd. by an ethereal solution of oxalic acid (Brücke, M. 8, 195). Musculus (C. R. 78, 182) filters putrid urine, dries the filter at 35°, and employs it as a test for urea, which it converts into ammonium carbonate. A conc. aqueous solution of furfuraldehyde followed by a drop of HClAq (S.G. 1-10) colours a crystal of urea

violet (Schiff, B. 10, 774). o-Nitro-benzoic aldehyde added to an alcoholic solution of urea forms, on warming, a white pp. of o-nitro-benzylidene-di-urea [200°]. When present in small quantities this compound may be detected by resolving into the parent substances by boiling with very dilute H_2SO_4 , and then adding a little phenylhydrazine solution, which produces a red colour (Lüdy, M. 10, 295). Mercuric nitrate gives a white pp. with solutions of urea. The various methods of estimating urea may also be employed for its detection.

Estimation. — Hypobromite method. Urea is mixed with NaOH (100g. in 250 c.c. water) to which bromire (25 g.) has been added, and the nitrogen collected and measured (Knop, Fr. 9, 226; Hüfner, J. pr. [2] 3, 1; Dupré, C. J. 31, 534; Simpson a. O'Keefe, C. J. 31, 838; Russell a. West, C. J. 27, 749; Eykman, R. T. C. 3, 125; Schleich, J. pr. [2] 10, 263; Colquhoun, C. N. 67, 123; Camerer, Zeit. Biol. 29, 239). When this method is applied to urine it must be remembered that uric acid gives off 48 p.c. and creatinin 37 p.c. of its nitrogen in this way (Falck, Fr. 21, 300). Nevertheless, the quantity of nitrogen evolved is 8 p.c. less than the calculated amount (R. a. W.). Using Knop's solution, ammonia, urea, and oxamide give off 93, 92·3, and 75 p.c. of their nitrogen as gas (Foster, C. J. 33, 470; 35, 119). In the case of urea and oxamide the 'suppressed' nitrogen is present as cyanate and as nitrate (Fauconier; Luther, H. 13, 500). The amount of 'suppressed' nitrogen is less in dilute than in concentrated solutions (Hüfner, H. 1, 350; cf. Pflüger a. Bohland, Pf. 38, 325; 39, 1, 143), and is greatly diminished by the addition of 1 to 5 p.c. acetoacetic ether (Jacoby, Fr. 24, 318). According to Duggan (Am. 4, 47), if bromine is added to a solution of urea in NaOHAq the yield of nitrogen is 99.4 p.c. of the theoretical amount. In estimating urea in urine Pflüger (Fr. 26, 117) adds HClAq (1 c.c.) to urine (10 c.c.), ppts. various nitrogenous bodies by phosphotungstic acid, neutralises the filtrate with lime and then adds an equal volume of conc. NaOHAq and, after allowing to stand for some time, decomposes with Knop's hypobromite. Frothing of albuminous urine can be avoided by first shaking with a pilule of fat (Méhu, J. Ph. [5] 15, 607). Hamburger (R. T. C. 2, 181) introduced a volumetric method of employing hypobromite, but Pflüger a. Schenck (Pf. 37, 399) consider the method to be untrustworthy. E. Salkowski adds two drops of HCl to urine (2.5 c.c.) diluted with 5 or 10 volumes of water until the air is expelled from the flask, then adds the hypobromite, continues boiling, and collects the nitrogen. Fenton (C. J. 33, 300) recommends the use of sodium hypochlorite, which was first proposed by E. W. Ravy (A.D. 1854). In this case free NaOH must not be present, though excess of Na₂CO₃ may be present.

Mercuric nitrate method. Urea is completely ppd. by a solution of $Hg(NO_s)_{2^{-}}$. The white pp. is not decomposed, and therefore not turned yellow by Na_sCO_s . Urine (2 vols.) is mixed with a solution (1 vol.) prepared from saturated solutions of baryta (2 vols.) and $Ba(NO_s)_2$ (1 vol.); filtered through a dry filter

from the ppd. sulphate and phosphate and 15 c.c. (equivalent to 10 c.c. urine) of the filtrate are titrated with standard mercuric nitrate till a drop taken out gives a yellow pp. with Na₂CO₂ (Liebig, A. 85, 370). Assuming the compound CON₂H₄2HgO to be formed, 1 pt. ures should ppt. 7.2 pts. HgO, but in practice 7.72 pts. of the latter are required. A solution of 71.48 g. mercury in HNO, diluted to 1 litre is equivalent to 01 g. urea per c.c. If the urine contains more than 2 p.c. urea the titration gives too low results; in this case the urine must be diluted. When mercuric nitrate is added to a solution of urea nitric acid is set free. Liebig recommended the addition of Na₂CO₃ from time to time to keep the liquid neutral. Pflüger (Fr. 19, 375) proceeds as follows: During the titration, after each addition of Hg(NO₃)₂ a drop is placed in contact with a little pasty NaHCO₃. Long before the titration is ended, a yellow colour is seen between the two drops, but disappears on mixing them together. When the yellow colour is permanent, the titration is near its end; at this point the solution is neutralised by Na₂CO₂, and the titration continued till a permanent yellow colour is got on mixing the drops. Should the entire liquid become yellow on adding Na, CO, a fresh quantity of urea solution must be taken and the operation be repeated with greater speed. The presence of over 1 p.c. NaCl in urine interferes with the titration when Na₂CO₃ is used as indicator, the final reaction not being sharp. Since NaHCO₃ does not ppt. HgCl₂, when the bicarbonate is used as indicator the titration may proceed as if NaCl were absent, subtracting the amount of Hg(NO₃)₂ that is converted into HgCl₂ by the chloride (Rautenburg, A. 133, 55; Pfeiffer, Fr. 24, 475; Pflüger, Fr. 27, 120). It is, however, better to ppt. the chlorides by AgNO₃ and titrate in the usual way.

Barium carbonate method. The solution is heated with BaCl₂ and NH₃Aq in sealed tubes at 220°-240° and the ppd. BaCO₃ weighed (Bunsen, A. 65, 575). Before applying this method to urine other substances which might form BaCO₃ must be ppd. by phosphotungstic acid (Pfüger a. Bleibtreu, Fr. 28, 377).

Kjeldahl's method. Nitrogen in urea may be estimated by Kjeldahl's method. 5 c.c. urine are heated with H₂SO₂ (10 c.c.), and Nordhausen H₂SO₄ (10 c.c.) until no more water or gas comes off, and the liquid is clear yellow. After cooling, water (200 c.c.) and NaOH are added and NH₂ distilled off. In the case of urine it must be remembered that about 13 p.c. of the nitrogen is, on an average, combined in substances other than urea (Pfüger, Pf. 35, 454; 40, 533; Camerer, Z. B. 24, 306).

Other methods. Ures may also be estimated by fermentation and determination of ammonia set free (Miquel, C. R. 111, 501).

Campani (G. 17, 187) proposes to decompose urea by nitrous acid and pass the CO₃ into a solution of lime-water of known strength, and titrate with oxalic acid.

Cazeneuve and Hugouneng (Bl. [2] 48, 82) heat urea with a large excess of water and titrate the resulting ammonium carbonate.

Riegler (F. 83, 49) decomposes urea with

Riegler (F. 83, 49) decomposes ures with Millon's reagent and measures the mixture of N and CO₂ evolved.

Möiner a. Sjöqvist (Fr. 80, 888) add 5 c.c. of a saturated solution of BaCl₂ containing 5 p.c. baryta to 5 c.c. urine and then add 100 c.c. of a mixture of 97 p.c. alcohol (2 pts.) and ether (1 pt.). After 24 hours the liquid is filtered, the pp. washed with 50 c.c. alcoholether, and the filtrate and washings evaporated at 50° to 25 c.c., MgO and some water added, and the evaporation continued as long as NH₂ comes off. The urea is then determined by Bunsen or Kjeldahl's method (Bödtker, H. 17, 140).

Salts .- B'HCl. Very deliquescent crystalline mass, formed from urea and dry HCl. Decomposed by water into urea and HCl. At 140° it yields NH₄Cl and cyanuric acid.—B'HNO₃. Small plates, sl. sol. water and alcohol, v. sl. sol. \hat{HNO}_3 . $-\hat{B'}_2\hat{H}_2\hat{C}_2\hat{O}_4$. S. 4.4 at 15°. S. (alcohol of S.G. 833) 1.65. Monoclinic tables; a:b:c=1: 564: 491; a=82° 10' (Loschmidt, Sitz. W. 51, ii. 7, 384). V. sol. hot water. Ppd. from its aqueous solution by oxalic acid.—

B'H₂C₂O₄aq (Lubavin, A. Suppl. 8, 83).—

B'HAuCl₄aq: orange-red prisms or needles, v. sol. water, alcohol, and ether.—B'2HAuCl.: yellow needles, v. sol. hot water (Heintz, A. 202, 264). — B'₂H₂PtCl₆2aq. Yellow, deliquescent tables, v. sol. water and alcohol, insol. ether (Heintz, A. 198, 91).—B'H₂PO₄. Large crystals, v. sol. water and alcohol, sl. sol. ether (Lehmann, Buchn. Rep. 15, 224; Schmeltzer a. Birnbaum, Z. [2] 5, 206). Its solution does not ppt. MgSO₄. -B'₂2H₃PO₄. Its aqueous solution gives a crystalline pp. of cyanuric acid on heating.

Tri-chloro-acetate B'C₂HCl₂O₂. Plates (from alcohol) (De Clermont, J. 1873, 536).—
The fumarate B'₂C₄H₄O₄, maleates B'C₄H₄O₄, and B₂'C₄H₀O₄, maleates B'C₄H₄O₄, gallate B'C₇H₆O₅, and succinate B'₂C₄H₀O₄, gallate B'C₇H₆O₅, and succinate B'₂C₄H₆O₄ [145°], all crystallise in monoclinic forms (Loschmidt, Sitz. W. 52; ii. 238). The parabanate B'C₄H₂N₂O₅ and tartrate B'C₄H₄O₆ are trimetric.—The citrate B'₂C₄H₄O₄ are trimetric.—The citrate B'₂C₄H₆O₄ and an acid citrate B'C₄H₆O₆, have been prepared (Hlasiwetz, J. 1856, 698).—Cyanurate B'C₃N₄N₆O₅. Monoclinic crystals.—Picrate B'C₄H₃N₄O₇, [142°] (Smolka, M. 6, 920). S. 1.9 at 18·5°. S. (95 p.c. alcohol) 6 at 18°. Slender yellow needles (from alcohol). Tri-chloro methane sulphinate B'CCl₃SO₂H. [96°–100°]. Thin prisms (McGowan, J. pr. [2] 36,

220).
Compounds with metallic oxides.—
B'28Ag_O (Liebig). Formed by adding moist
Ag_O to a solution of urea. Grey powder composed of slender needles. According to Mülder

(B. 6, 1019), the yellow pp. got by adding NaOH to a solution of urea mixed with AgNO₃ is CON₂H₂Ag₂. It combines with iodine, forming CON₂H₂Ag₂I, a greenish mass, darkened by light (Tafel a. Enoch, B. 23, 1554).—B'₂HgO. Got by adding Hg(NO₃)₂ to a solution of urea mixed with KOH. White pp.—B'₂ShgO. White pp. formed by adding HgCl₂ to a solution of urea and HgCl₂, changes to a yellow granular powder on boiling with water.—B'HgO. Formed

by adding Hg(NO_s), to a warm solution of urea (Dessaignes, A. 82, 232; Liebig, A. 85, 289).

Compounds with metallic salts.—

B'AgNO_s. Prisms. Yields silver cyanate on

boiling with water.-B'.PdCl. Ppd. by adding a solution of PdCl₂ to one of urea (Drechsel, J. pr. [2] 20, 469).—B'NaClaq. Formed by evaporation of a solution of urea and NaCl. Deliquescent prisms. Alcohol extracts urea from the compound.—B'NaNO, aq. Prisms (from water).-B'NH,Cl. Leliquescent crystals. -B's(NH,Cl),HCl. Formed by dissolving urea in bleaching-powder solution (Beckmann, A. 91, 367). Large plates, sol. alcohol mixed with ether.—B',Mg(NO₃), (Werther, J. pr. 35, 5).—B',Ca(NO₃),—B',ZnCl₂. Very deliquescent crystals (Neubauer a. Kerner, A. 101, 337).— $B'CdCl_2$. Needles.— $B'_2Hg(NO_3)_2BHgO$. Granular powder formed by mix ng warm dilute solutions of urea and mercuric nitrate. At the moment of formation the pp. is flocculent.— B'2Hg(NO₃)2HgO. Formed by pouring mercuric nitrate into a solution of urea as long as a pp. is produced, and keeping the whole at 40° to 50° for some time (Liebig). Six-sided laminæ. —B'₂Hg(NO₂)₂HgO. Formed by adding an acid solution of mercuric nitrate to a solution of urea nitrate. Crusts of small tabular crystals (Liebig, A. 85, 296).—B'HgCl₂. Flat crystals, v. sl. sol. cold water (W.).—B'₁₂Cr₈O₂₁ 3aa. Formed by the action of water on the product of the action of CrO₂Cl₂ on urea (W. J. Sell, Pr. 33, 267; 45, 321). Olive-green needles (from hot water). PtCl₄ added to its solution in hot water ppts. silky green needles of B'12Cr2Cl6(PtCl4), 2aq. The chromate treated with PbCl, forms lead chromate and a liquid from which gaseous HCl ppts. green prisms of B'12Cr2O6 6aq, the aqueous solution of which heated with Ag2SO4 gives dark-green prisms of B'12Cr2(SO4), 10aq. similar way green prisms of B'₁₂Or₂(NO₃), may be got.—B'₁₂Or₆Cl₂O₁₄ 2aq. Green monoclinic Green monoclinic crystals, got from the product of the action of CrO.Cl. on urea by crystallising from conc. HClAq (1 vol.) diluted with water (9 vols.). By treatment with water it is split up into B'12 Cr8O21 and B'12Cr2Cl8. By treatment with conc. HClAq and B₁₂Cr₂Ot₃. By treatment with conc. HOIM
(1 vol.) and water (6 vols.) it is converted into
the salts B'₁₂Cr₂Cl₃O₁₂Saq. — B'₁₂Cr₂O₁₂4aq.
— B'₁₂Cr₄(C₂O₃)₆4aq. — B'₁₂Cr₂Cl₅6HgCl₂. —
B'₁₂Cr₄(C₂O₃)₆4aq. —B'₁₂Cr₂I₁₈. — B'₁₂Cr₂(SO₃)₃I₆.
—B'₁₂Cr₂(CO₃)₂I₄. — B'₁₂Cr₂(CO₃)₂I₆. —B'₁₂Cr₂Br₁₆.

Bronze-yellow tables, giving off bromine in

air.—B'₂CuCl₂. Small blue crystals.

Formy l derivative NH₂CO.NH.CHO.

[159°]. H.F. 13,400 (Matignon, C. R. 112, 1367). Formed by boiling urea with conc. formic acid (Geuther, Z. [2] 4, 300). Crystals, v. sol. water, being decomposed into urea and formic acid.

Acetyl derivative C₁H₆N₂O₂ i.e. NH₂CO.NHAc. Acetureide. Mol. w. 102. [214°]. H.F. -200 (M.). S. (alcohol) 1 in the cold; 10 at 78°. Formed by boiling ures with AcCl or Ac₂O (Zinin, A. 92, 40°; G.; Zande, R. T. C. 8, 235). Got also by the action of KMnO₄ on methyl-uracil (Behrend, A. 229, 29). Four-sided needles (from alcohol). Slowly but completely decomposed by pure HNO₂ with evolution of CO₂ (1 vol.) and N₂O (2 vols.) (Franchimont, R. T. C. 6, 215). It is not ppd. by Hg(NO₂)₂.

Di-acetyl derivative CO(NHAc),. [158°]. A product of the action of COCl, on acetamide at 50° (Schmidt, J. pr. [2] °5, 68).

Formed also from mercuric fulminate and AcCl (Scholl, B. 23, 3515). Needles (from alcohol). sl. sol. cold water.

Chloro-acetyl derivative

NH2.CO.NH.CO.CH2Cl. Formed from chloroacetyl chloride and urea (Jazukovitch, Z. 1868, 234; Tommasi, C. R. 76, 640). Thin needles (from alcohol). Begins to melt at 160°.

Tri-chloro-acetyl derivative NH₂.CO.NH.CO.CCl₃. [150°]. Crystals (from alcohol) (De Clermont, J. 1874, 798; Meldola a. Tommasi, C. J. 1874, 404; Cloez, A. Ch. [6] 9,

219). Nearly insol, hot water.

Bromo-acetyl derivative

NH₂CO.NH.CO.CH₂Br. Needles (from dilute alcohol) (Baeyer, A. 130, 156). Pure HNO₂ gives off CO₂ (1 vol.) and N₂O (2 vols.) (Franchiment B T C 6 218) mont, P. T. C. 6, 218).

Tri-bromo-acetyl derivative NH. CO.NH. CO.CBr. [148°]. Formed by the action of Br on an aqueous solution of di-bromobarbituric acid (Baeyer). Crystals, v. sol. hot alcohol. Vields crystalline B'Ba(OH), xaq, v. e. sol. water.

Cyano-acetyl derivative NH, CO.NH.CO.CH, Cy. [200°-210°] (Mulder, B. 12, 466). Crystalline.

Butyryl derivative NH2.CO.NHC,H,O. [176°]. Formed from urea and butyryl chloride (Moldenhauer, A. 94, 101). Plates.

Isovaleryl C₆H₁₂N₂O₂. derivative [191°]. Minute prisms, v. sl. sol. water.

Carbonyl derivative CO(NH.CO.NH.)₂. Formed by heating urea with COCl₂ at 100° (E. Schmidt, J. pr. [2] 5, 39; Emich, M. 10, 347). Powder composed of minute crystals, v. sl. sol. cold water and alcohol. Decomposed by heat into NH, and cyanuric acid. Boiling KOH also produces cyanuric acid.—B'HgO: crystalline powder, insol. water.

Succinyl derivative C₄H₄O₂(NH.CO.NH₂)₂. Formed by heating urea (2 mols.) with succinyl chloride (1 mol.) at 65° (Conrad, J. pr. [2] 9, 301). Colourless powder, v. sl. sol. hot water.

Methyl-malonyl derivative CHMe:C.O.:N.H.CO. [192°]. Formed by heating methyl-malonic acid with urea and POCl. (Franchimont a. Klobbie, R. T. C. 7, 22). Pointed plates, v. sol. water and alcohol.

Bensoyl derivative NH2.00.NHBz. 200°]. S. (alcohol) 1 in the cold; 4 at 78°. Formed by heating urea with BzCl or Bz₂O at 150° (Zinin, 4. 92, 404; Geuther, Z. [2] 4, 299). Four-sided plates (from alcohol), insol. ether).

CO(NHBz), derivative Di-bensoyl [210°] (S.); [197°] (H.). A product of the action of COCl. on benzamide at 165° (E. Schmidt, J. pr. [2] 5, 58). Formed also by heating guanidine carbonate with Bz₂O at 100° (Creath, B. 7, 1789), and by treating mercuric fulminate with BzCl (Hollemann, R. T. C. 10, 72; B. 23, 2998, 8742). Needles (from alcohol), sl. sol. water. Aniline at 180° gives NH₂, benzamide,

benzanlide, and s-di-phenyl-urea.

**u-Di-bensoyl derivative NH, CO.NBz.*

[c. 197°]. Formed by heating sodium benzoylcyanamide NaBzN.ON with BzCl, followed by boiling alcohol (Buddéus, J. pr. [2] 42, 97). White needles.

NH2.CO.NH.CO.C.H4.NO2 [1:3]. Plates (Griess, B. 8, 222).

m-Nitro-bensoyl derivative

m-Amido-bensoylderivative C8H2N3O2. Formed by boiling the preceding body with aqueous ammonium sulphide (G.). Needles (from water), decomposing at 2005. — B'HCl aq: needles. -B'2H, PtCl.

(β)-Naphthoyl derivative NH. CO.NH.CO.C, H,. [215°]. Formed from naphthoyl chloride and urea (Vieth, A. 180, 319). Minute needles, m. sol. alcohol, v. sl. sol. benzene.

Di-(\beta)-naphthoyl urea [286°] (Ekstrand, B. 20, 1353). Needles.

Lactyl derivative v. LACTYL-UREA.

Glycollyl derivative v. Hydantoïn. Reference.—OXY-UREA.

UREA CARBOXYLIC ACID v. ALLOPHANIC

UREA CHLORIDES. A name sometimes alkyl-carbamic chlorides used to denote X.NH.CO.Cl.

URECHITIN C₂₈H₄₂O₈. Occurs, together with urechitoxin, in the leaves of *Urechites* suberecta, growing in Jamaica (Bowrey, C. J. 33, 252). Crystallises from alcohol in needles (containing xaq), tastes bitter. It is very poisonous (Pr. 27, 309). Nearly insol. water, m. sol. ether. Appears to be a glucoside. Conc. H₂SO, forms a yellow liquid, turning red, and finally purple, especially in presence of an oxidising agent.

Urechitoxin C18H20O5. Bitter poison, split up by acids into sugar and urechitoxetin.

UREIDES. Compounds obtained by elimination of water between urea and an acid or an aldehyde. Many of them may be represented as amides in which NH2 is replaced by NH.CO.NH2.

BIURET v. vol. i. p. 517. URETHANE v. CARBAMIC ETHER. ISURETINE v. FORMAMIDOXIM. URIC ACID C, H, N,O, i.e.

CO NH.CO.C.NH CO (Medicus, A. 175, 243; Fischer, B. 17, 329, 1776). Mol. w. 168. S.G. 1.87. H.C.p. 461,400 (Berthelot, C. R. 110, 887; Matignon, C. R. 110, 1267); 462,500 (Stohmann, J. pr. [2] 44, 390). H.C.v. 462,700 (M.). H.F. 148,100 (M.). S. 002 at 0°; 006 at 20°; 0625 at 100° (Blarez a. Denigès, C. R. 104, 1847; cf. Garrod, Pr. 85, 63).

Occurrence.-In urine and in urinary calculi (Scheele, A.D. 1776; Liebig a. Wöhler, A. 26, 241; Horbaczewski, M. 12, 221). About 5 g. is daily excreted in human urine. Uric acid occurs in urine of carnivora, herbivora (Mittelbach, H. 12, 463), and of birds. Serpent's urine is chiefly composed of acid ammonium urate. A considerable quantity occurs in the blood of gouty subjects, and sodium urate is deposited as 'chalk stones.' Very small quantities are normally present in the liver, lungs, brain, and blood. It occurs in the green gland of the fresh-water crayfish (Griffiths, Pr. 88, 187). Urea, glycocoll, leucine, and aspartic acid given to fowls appear in the urine as uric acid (Jaffé a. Meyer, B. 10, 1930; Knierim, B. 10, 1930).

Synthesis.—1. By rapidly heating glycocoll (1 pt.) with urea (10 pts.) to 280° (Horbaczewski, B. 15, 2678; M. 8,796; 6,856).—2. By heating

Vor. IV.

URIC ACID.

urea with tri-chloro-lactic acid or its amide (Horbaczewski, M. 8, 201, 584).—3. By preparing methyl-uracil from acetoacetic ether and urea; converting methyl-uracil into nitro-uracil carboxylic acid by treatment with fuming HNO,; boiling the carboxylic acid with water, and reducing the resulting nitro-uracil to amido-uracil, and finally to isobarbituric acid by means of tin and HClAq; oxidising the isobarbituric acid to isodialuric acid; and heating the isodialuric acid (1 pt.) with urea (1 pt.) and H₂SO₄ (6 pts.) (Behrend a. Roosen, A. 251, 235).—4. By fusing urea (4 g.) with cyano-acetic acid (1 g.) (Formanek, B. 24, 3419).

Preparation.—1. Serpent's urine is dissolved in boiling KOHAq, filtered, and the uric acid ppd. by HCl or dilute H,SO. If the uric acid is much coloured a little KMnO, or K, Cr,O, (Gibbs, Am. S. [2] 48, 215) should be added to the boiling alkaline solution before ppn.-2. Guano is boiled with borax (1 pt.) and water (120 pts.), and the solution of sodium urate thus obtained ppd. by HCl.-3. Guano is extracted with dilute HClAq, and the residue dissolved in conc. H2SO4

and ppd. by water.

Properties.—White anhydrous scales (by a.). When slowly deposited from dilute soluppn.). tions (e.g. urine) it separates as large crystals (containing 2aq). Nearly insol. water, insol. alcohol and ether. Sol. conc. H₂SO₄ and reppd. by water. Its solution in hot H2SO4 deposits on cooling a deliquescent compound of uric acid with H2SO4. Uric acid is m. sol. glycerin, hot NaOAc, and sodium phosphate. Lithium carbonate (1 pt.) dissolved in water (90 pts.) can dissolve uric acid (4 pts.). Aqueous

K₂CO₂ also dissolves uric acid.

Reactions.—1. Decomposed by heat, without fusion, yielding HCy and a sublimate of cyanuric acid, ammonium cyanate, urea, and ammonium carbonate, and leaving a carbonaceous residue. 2. Water at 180° forms mycomelic acid (Hlasiwetz, A. 103, 211).-3. Potash-fusion yields NH_s and potassium oxalate, carbonate, and cyanate. - 4. Chlorine and bromine in presence of water yield alloxan, urea, and parabanic acid (Hardy, A. Ch. [4] 2, 372). Alloxan is also produced by iodine, by HNO, and by MnO, and H₂SO₄.—5. Boiling with water and PbO₂ yields allantoin. Allantoin is also produced by ozone (Gorup-Besanez, A. 110, 94), alkaline K, FeCy, KMnO₄ (Claus, B. 7, 226), and MnO₂ and water. 6. Pure HNO, decomposes uric acid in the cold, with evolution of CO₂ (14 vols.), N₂O (24 vols.), and nitrogen (4 vols.) (Franchimont, R. T. C. 6, 222). 7. KNO2 and HOAc yield urinilic acid (Sokoloff, Z. [2] 5, 78) and stryphnic acid (Gibbs, Am. S. [2] 48, 215).—8. HIAq at 165° forms glycocoll, NH₄I, and CO₂ (Strecker, Z. [2] 4, 215).—9. Exposed to the air in alkaline solution it yields exonic and uroxanic acids.—10. A dilute olution of sodic urate absorbs oxygen from the air at 35° according to the equation $C_sH_sN_sO_s + O + 2H_sO$ = $C_sH_sN_sO_s$ (uroxanic acid). Uric acid in dilute NaOH solution is completely decomposed by 85 days' digestion at 85° (Nencki a. Sieber, J. pr. [2] 26, 18). 5 g. uric acid dissolved in 200 c.c. water and 20 g. potash disappeared in 5 days. However, if no oxygen be present the uric acid is not affected.-11. In fermenting urine uric acid is completely split up into CO2 and NHa (Sestini, G. 20, 133).

Detection.-Uric acid may be recognised by its insolubility in water and HClAq, by its crystalline character under the microscope, and especially by the murexid reaction. evaporated with HNO, on a water-bath it leaves an orange residue, which is coloured violet-red by ammonia and violet-blue by potash. A solution of uric acid in aqueous Na2CO, produces a dark-brown spot of reduced silver when added to paper moistened with AgNO₃ (Schiff, A. 109, 65). Uric acid (1 mol.) reduces boiling Fehling's solution, forming a pp. of Cu₂O (1 mol.) (W. Müller, J. Th. 1881, 73). In presence of KOH uric acid dissolves CuO, forming a blue solution, which slowly deposits white cuprous urate. The compound C,H,N,O,Cu,O is ppd. when an alkaline solution of uric acid is treated with Fehling's solution and hydrox amine hydrochloride (Balke, J. pr. [2] 47,546). Uric acid evaporated with a little dilute HNO, gives a blue colour on shaking with H₂SO₄ and benzene containing thiophene (Deniges, J. Ph. [5] 18, 161).

Estimation in urine.—1. Urine (500 c.c.) is concentrated, HCl is added, and after t./enty-four hours the pp. is collected and weighed. Uric acid may be separated from xanthine by ppn. with conc. H₂SO₄ (Horbaczewski, H. 18, 341; cf. Wulff, H. 17, 634).—2. Urine (200 c.c.) is mixed with conc. Na₂CO₂Aq (10 c.c.), and after an hour conc. NH₄ClAq (20 c.c.) added. After forty-eight hours the pp. is collected, washed with HOl (1 pt. of S.G. 1-123), diluted with water (10 pts.), the washings allowed six hours to deposit any dissolved uric acid, the combined pps. washed with alcohol and dried at 110° (Ŝalkowski, Fr. 16, 373). Uric acid may also be ppd. as ammonium urate by saturating urine with solid NH₄Cl (30 g. to 100 c.c.) (Hopkins, Pr. 52, 93).—3. Fresh urine is mixed with very dilute (15 p.c.) NaOHAq, filtered from phosphates, and diluted to S.G. 1.010. To 300 c.c. of this solution is added 50 c.c. of magnesia mixture [MgSO, 7aq (1 pt.), NH,Cl (2 pts.), NH,Aq (4 pts. of S.G. 924), and water (8 pts.)]. The pp. is filtered off, and 175 c.c. of the filtrate mixed with 5 g. CaCO, and 5 c.c. of a 3 p.c. solution of AgNO₃. The pp. is washed and dried, and a nitrogen estimation made by Kjeldahl's method (Camerer, Z. B. 26, 84; cf. Salkowski, Pf. 5, 319).—4. Urine (25 c.c.) is mixed with NaHCO₂ (1g.), and NH₂Aq (2 to 3 c.c.), and ammoniacal AgNO₃ (1 to 2 c.c. of a solution of 5 g. AgNO₃ in 100 c.c. water, to which NH₃Aq is added till the ppd. Ag₂O is just re-dissolved). The pp. is collected, washed, dissolved in HNO., and the silver determined volumetrically by the sulphoyanide method (Haycraft, Er. 25, 167; 80, 648). This method is liable, according to Salkowski (H. 14, 31) and Gossage (Pr. 44, 284), to an error which may amount to 60 p.c. The error is partly due to variations in composition of the silver urate, and partly to presence of bodies of the xanthine group (Deroide, Bl. [8] 7, 863). V. also Herrmann, H. 12, 496; Czapek, H. 12, 502).—5. In a solution containing not more than 0125 p.c. of uric acid 1 c.c. of decinormal permanganate is reduced by 0074 g. uric acid (Blarez a. Deniges, C. R. 104, 789).—6. By treatURSONE. 883

ment with a standard iodine solution in presence

of alkali (Kreidl, M. 14, 109).
Salts.—(NH₄)HA". S. 06 at 15°. Slender needles.—(NH₄)₃HA"₂.—(NH₄)₄H₂A"₃ (Maly, J. 1863, 621).—K₂A". S. 3 at 16°. Small needles.— KHA". S. 13 at 20°. Amorphous.—Na,A" aq. S. 1.5. Nodules.—NaHA" aq (dried at 100°). Crystalline powder. S. .09 at 15°; ·8 at 100°. Occurs as an amorphous urinary deposit.— LiHA". S. 27 at 20; 9 at 40°; 2.5 at 100° (Schilling, A. 122, 241).—BaA"aq (dried at 100°). Granular pp. S. 013 in the cold.—BaH_A"_2 2aq (dried at 100°). Amorphous, insoluble powder (Behrend a. Boosen, A. 251, 250).—CaA". S. 066 in the cold; °C7 at 100°.—CaH_A"_2 2aq. S. 15 in the cold; 37 at 100°. More sol. KClAq.—SrA" 2aq. Minute stellate groups of needles. S. $\cdot 023$ in the cold; $\cdot 055$ at 100° .—SrH₂A"₂2aq.—MgH₂A"₂6aq. S. $\cdot 03$ in the cold; $\cdot 6$ at 100° .—PbH₂A"₂ 'dried at 100°). Insol. water.—PbA" (dried at 100°).—Cu₂A" aq.— Cu₃A"₂(OH)₂ 4aq': green pp.

**Reference.—METHYL-URIC ACID.

Iso-uric acid C,H,N,Os. Formed by boiling cyanamide (1 pt.) with an aqueous solution of alloxantin (2 pts.) (Mulder, B. 6, 1236; 7, 1633). Powder, almost insol. water. Oxidised by boiling with I and water. Its solution in K2COs reduces AgNO₈.

Pseudo-uric acid C,H,N,O, i.e.

CO NH.CO CH.NH.CO.NH2. Formed by the action of KCyO on uramil (Baeyer, A. 127, 3), and by heating uramil with urea at 180° (Grimaux, Bl. [2] 31, 535). White crystalline powder composed of small prisms, v. sl. sol. water. HNO₃ yields alloxan. Boiled with water and PbO₂ it yields oxalic acid, but no allantoin. H₂SO₄ at 150° forms xanthinine.— (NH₄)HA" aq.—KHA" aq: scales.—NaHA" 2aq: groups of prisms, v. sol. hot water.—BaA" 5aq: groups of slender needles.

URINILIC ACID C.H.N.O. Formed by the action of KNO, and HOAc on uric acid (Sokoloff, Z. 1869, 78). Prisms (from water).—K₂HA'''. Prisms, v. e. sol. water. Ca, A'''₂. Crystalline pp. insol. water and HOAc.—Sr, A'''₂.—Ba, A'''₂.— CdHA''' Saq —CuHA''' 4aq. Slender red needles. -Ag,HA". Pp.-Ag,A". Gelatinous pp. quickly

blackening in light.

UROBILIN. Occurrence.—Often in urine (Jaffé; Disqué, H. 2, 271; C. C. 1878, 711; Eikholz, J. Physiol. 14, 326). In bile, especially of the mouse. Preparation.—Urine containing urobilin is

ppd. by lead acetate and sub-acetate until the absorption band at F is removed. The pp. is extracted with alcohol, acidified with HCl or H2SO4, filtered, diluted with water, and extracted with chloroform.

Properties .- A red amorphous, shiny substance. Prepared as above it contains HCl or H₂SO₄, as the case may be. Its spectrum exhibits a dark band at F, destroyed by NH₂, but on adding NaOH another band near the red is formed (MacMunn, Pr. 30, 250; 31, 26, 206; p. also Hydrobinibush, this vol. p. 278.

Detection and estimation in urine: Jolles, H. 18, 545; Studensky, C. C. 1893, ii. 668. UROCANINIC ACID C₁₂H₁₂N₄O₄. [218°]. Ocsurs in dog's urine (Jaffé, B. 7, 1669; 8, 811).

Prisms (containing 4aq), v. sol. hot water, insol. alcohol and ether. Decomposed by fusion into CO₂ and urocanine C₁₁H₁₀N₄O, an amorphous base yielding B'H₂PtCl₃.—HA'2HCl: needles, v. e. sol. water.—HA'2HNOs: crystalline pp. insol. dilute HNO, v. sol. water.—HA'H2SO. Minute needles and plates.

UROCHLORALIC ACID C.H.;Cl.O, (?) [142°]. S. (ether) .4. Found in the urine after administration of chloral (Von Mering a. Musculus, B. 8, 662; H. 6, 483; Külz, Č. C. 1881, 486). Silky needles, v. e. sol. water and alcohol. Decomposed by boiling dilute H2SO, into tri-chloroethyl alcohol and glycuronic acid $C_0H_{10}O_7$.—NaA'.—KA'.—BaA'₂: satiny plates (from etheralcohol).

UROFUSCOHÆMATIN v. this vol. p. 274.

UROLEUCIC ACID C.H. O. . [133.5°]. An acid occurring in human urine which darkens on addition of alkalis (Kirk, Brit. Med. Journ. [1888] 2, 232). Crystals. Its solution is ppd. by Pb(OAc), and gives a transient green colour with very dilute FeCl. FeCl. colours the crystals red.

UROMELANIN $C_{18}H_{48}N_7O_{10}$. A black substance which separates on treating urine with H2SO4 or HClAq in presence in air (Thudichum, J. pr. 104, 257; C. R. 106, 1803; Plosz, H. 8, 89; Udranski, H. 11, 537; 12, 32; this vol. p. 274). It yields C₁₈H₄₀AgN₇O₉ (Thudichum).

URONITROTOLUOLIC ACID C12H15NOp. Occurs in urine of dogs after a dose of o-nitrotoluene (Jaffé, H. 2, 47). Very deliquescent. Asbestos-like mass, v. e. sol. water and alcohol. Decomposed by dilute H₂SO₄ into o-nitro-benzyl alcohol and a syrupy acid. Salts. - BaA'2: crystalline powder, v. e. sol. water, insol. alcohol. $-CO(NH_2)_2HA'2_2^1$ aq. [149°]. Needles, v.e. sol. water, sl. sol. cold alcohol.

UROROSEIN v. this vol. p. 274.

UROSULPHINIC ACID C.H.N.SO2. Formed by heating thio-uramido-barbituric acid (1 pt.) with H₂SO₄ (2 pts.) at 160° (Nencki, B. 4, 724; 5, 45). Nodules (from hot HClAq). Its K salt

orystallises in needles, m. sol. cold water.

UROXANIC ACID C.H.N.O. Formed by leaving a solution of uric acid (5 g.) in water (200 c.c.) containing KOH (20 g.) to stand for 5 days at 85°, neutralising by HOAc, filtering, and evaporating (Nencki a. Sieber, J. pr. [2] 24, 504; cf. Städeler, A. 78, 286; 80, 119; Strecker, A. 155, 177; Mulder, B. 8, 1291). Tetrahedra, sl. sol. cold water, insol. alcohol. Decomposed by boiling water into CO2, urea, and allanturio acid.—K,A" Saq. Four-sided pearly plates, v. sol. hot water.—CaA" 4aq.—BaA" 5aq.— PbA" aq. Insol. water.—Ag,A": flocculent pp., turned red by light.

URSONE C, H48O, 2aq. [265°] (Gintl, M. 14. 255). Ocsurs in the leaves of the red bear-berry (Arctostaphylos Uva-ursi) (Trommsdorff, Ar. Ph. [2] 80, 273; Hlasiwetz, J. pr. 64, 123). Needles, insol. water, acids, and alkalis, sl. sol. alcohol and ether. Conc. H₂SO₄ added to its solution in Ac₂O gives a red colour, changing through blue to green. Does not react with hydroxylamine or phenyl-hydrazine. Reduced by HI and P at

860° to O. H., (256°-267°).

Acetyl derivative [264°] Bensoyl derivative [214°].

8 H 2

URUSHIC ACID C14H18O2. S.G. 25 .985. Occurs in urushi or Japanese lacquer (Yoshida, C. J. 43, 472). Dark, pasty mass, v. sol. benzene and ether, insol. water. Br in CS₂ forms $C_{14}H_{12}Br_{8}O_{2}$. HNO₃ forms $C_{14}H_{16}(NO_{2})_{2}O_{2}$, which forms FeA', . Chromic acid mixture forms oxyurushic acid C₁₄H₁₈O₃.—PbA'₂ (dried at 100°). [110°-115°]. Flocculent pp.—FeH_pA'₁₂2aq.— FeH_pA'₈. Deep-black pp.

USNETIC ACID C.H.,O. [172°]. Occurs in small quantity, together with usnic acid, in the lichen Usnea barbata (Hesse, B. 10, 1326). Flat prisms (from alcohol), v. sol. ether. FeCl. colours its alcoholic solution bluish-violet. Bleaching-powder does not colour its alkaline solution. Is perhaps identical with decarb-

usnin.

USNIC ACID C18H16O7 (Salkowski; Paterno) or C₁₉H₁₆O₈ (Hesse), or C₁₈H₁₈O₇ (Stenhouse a. Groves, C. J. 39, 234). Carbusnic acid. [195·4° cor.]. S. (ether) 3 at 20°. Occurs in the lichens Usnea barbata, Usnea florida, Zeora sordida (Knop, A. 49, 103; Rochleder a. Heldt, A. 48, 12; Thomson, A. 53, 252; Stenhouse, Pr. 18, 222; A. 68, 97, 114; 155, 51; Hesse, A. 117, 343; Paterno, G. 8, 225; Salkowski, B. 8, 1459). Sulphur-yellow monoclinic crystals, sl. sol. alcohol, v. sol. hot ether and benzene. Does not yield betorein when heated. Conc. H2SO4 forms usnolic acid C₂,H_{2i}O₁₀ [213°] crystallising in small yellowish prisms (Stenhouse a. Groves, C. J. 39, 236). When it is heated with an alcoholic solution of aniline it yields the anilide C₁₈H₁₅O₆(NHPh), crystallising from alcohol in pale-yellow plates [171°], and forming in KOHAq a yellow solution from which HCl ppts. C22H22NO

Salts.—NaC₁₆H₁₇O₈ 2aq (Spica, G. 12, 432). Pale-yellow, silky needles. — $KC_{19}H_{18}O_8$ 3aq or $KC_{18}H_{17}O_8$ 3aq. Pale-yellow plates (from 50 p.c. alcohol) (Hesse, B. 10, 1325).— $KC_{19}H_{18}O_8$ aq or $KC_{18}H_{17}O_8$ aq. Yellow prisms (from 93 p.c. alcohol)

hol).—CuA'2: green pp.

Decarbusnein $C_{i,H_{i0}}O_{i}$. [175°]. Formed by heating usnic acid (1 pt.) with alcohol ($3\frac{1}{4}$ pts.) at 150° (Paterno, G. 12, 234). Yellow, silky Yellow, silky needles, sol. hot alcohol, sl. sol. water and ether. Not coloured by FeCl. Not attacked by AcCl. Reduces warm ammoniacal AgNO, Boiling KOHAq in absence of air splits it up into HOAc and decarbusnic acid C15H16O5, which crystallises from alcohol in lemon-yellow prisms [199°] and yields, when boiled with Ac.O, two acetyl derivatives C₁₅H₁₅AcO₅ [148°] (isomeric

with decarbusnein and C,H,Ac,O, [1317].

Pyrousnic scid C,H,O, (P.); C,H,O, (Salkowski, B. 8, 1461). [197°]. Formed, together with pyrousnetic acid, by boiling usnic acid (2 pts.) with KOH (5 pts.) and water (5 pts.) in an atmosphere of hydrogen (Paterno, G. 6, 113; 12,

242). Shining scales (from ether), v. sol. alcohol, sl. sol, ether. Its alkaline solution absorbs oxygen from the air, turning green and ultimately brown. Reduces ammoniacal AgNO, readily. Ac,O yields an acetyl derivative [205°]. Alcohol and HCl form an ether [147°]. Pyrousnic acid is split up on ary distillation in a current of H into CO2 and usneole C11H12O3, which crystallises from alcohol in yellow prismatic tablets [176°] and is converted by AcCl into a di-acetyl derivative [142°].

Pyrousnetic soid C₁₄H₁,O₂ i.e. C₁₈H₁₂O₄(CO₂H). [186°]. Formed by boiling usnic soid (10 pts.) with KOH (25 pts.) and water (25 pts.) for 15 minutes (Paterno, G. 12, 238). Colouriess plates or needles (from alcohol). Ac O yields C₁,H₁,AcO₆ [168°]. When heated in a current of H, pyrousnetic acid is split up into CO₂ and us netole C₁,H₁,O₄, which crystallises from dilute alcohol in yellowish

needles [179°].

(β)-Usnic acid v. Cladonic acid. UVIC ACID v. Pyrotritaric acid. UVINONE C14H12O4 i.e.

 $O<_{\text{CMe:C.CO.C:CMe}}^{\text{CMe:C.CO.C:CMe}}>0$ (?) [247°]. to the extent of 2 p.c., by distilling pyrotritaric acid (Dietrich a. Paal, B. 20, 1086). Yellow needles (by sublimation), sol. HOAc, v. sl. sol. alcohol. Conc. H.SO, forms a solution with green fluorescence. Br at 100° forms crystal-

line C, H, Br, O,

UVITIC AICD C,H,O, i.e. C.H.Me(CO.H)₂[1:3:5]. *Mesidic acid.* Mol. w. 180. [288°]. H.C.v. and p. 928,900. H.F. 193,100 (Stohmann, *J. pr.* [2] 40, 128). Formed, together with uvitonic acid, by boiling pyruvic acid with baryta-water (Finck, A. 122, 184). Formed also by oxidation of mesitylene or mesitylenic acid (Fittig a. Von Furtenbach, Z. [2] 4, 1; A. 147, 295), by oxidation of di-methylethyl-benzene (Wroblewsky, A. 192, 217), and by heating C_eH₂Me(SO₂NH₂)(CO₂H)₂ with conc. HClAq at 230° (Hall a. Remsen, Am. 2, 136). Slender needles (from water), v. sl. sol. hot water, m. sol. alcohol and ether. Yields toluene on heating with soda-lime (Baeyer, Z. [2] 4, 119). The Ca salt heated with lime yields mtoluic acid (Böttinger a. Ramsay, A. 168, 255). Chromic acid mixture oxidises it to trimesic

Salts.—K₂A": plates (from alcohol), v. sol. water.—BaA" aq.—CaA" aq.—Ag.A". Nearly insol. cold water.

Ethyl ether Et₂A". [35°]. Crystalline. References .- AMIDO-, BROMO-, NITRO-, and OXY-UVITIO ACID.

Isouvitic acid v. CARBOXY-PHENYL-ACETIC ACID. Uvitonic acid v. METHYL-PYRIDINE-DICARB-OXYLIC ACID.

VALERAL v. VALERIC ALDEHYDE.

VALERAL-DI-ACETONAMINE v. Pentylidene di-Acetonamine.

[41°]. Formed by the action of H₂S on isovaleric aldehyde ammonia suspended in water (Beissenhirtz, A. 90, 109; Parkinson, A. 90, 119). Formed also from thio-isovaleric aldehyde and VALERALDINE C., Hans. Thiovaleraldine. dry NH, (Schröder, B. 4, 468). Plates (from ether), with powerful smell, sol. alcohol, insol. water.-B'HCl. Needles, sol. water.

VALERCREATININ v. METHYL-GUANIDO-VALERIC ACID.

VALERIAN OIL. An essential oil obtained from the root of V-leriana officinalis. It contains a terpene $C_{10}H_{18}$ (156°) [$a]_D = -21^\circ$, a camphene, borneol $C_{10}H_{18}$ 0, di-bornyl oxide ($C_{10}H_{13}$)20 (285°-290°), formyl-, acetyl-, and valeryl-borneol, and a little valeric acid (Germann Colors) hardt, A. Ch. [3] 7, 275; Pierlot, A. Ch. [3] 56, 291; Bruylants, J. Ph. [4] 27, 349; B. 11, 452; Haller, C. R. 103, 151; Oliviero, C. R. 117, 1096).

n.VALERIC ACID C.H. $_{10}O_{2}$ i.e. CH. CH. CH. CH. CO. Mol. w. 102. (186° cor.). S.G. $_{0}^{\circ}$ 9562 (Zander, A. 224, 65); $_{15}^{15}$ 9446; ½ 9375 (Perkin); ½ 9298 (Brühl). C.E. (0°-10°) 00098 /Z.). S. 3·5 at 16°. S.V. 130.0 (Z.). M.M. 5.51. at 13.6°. $\mu_{\rm g}=1.4093$. R_{∞} = 43.16 (Brühl). H.F. 137,800 (Stohmann, *J. pr.* [2] 49, 99). Occurs in crude wood vinegar (Grodzki a. Krämer, B. 11, 1358).

Formation.—1. By saponifying its nitrile (nbutyl cyanide) (Lieben a. Rossi, A. 159, 58; G. 1, 239).—z. By oxidation of n-amyl alcohol (L. a. R.). - 3. By the action of metallic silver on a mixture of EtI and β -iodopropionic acid (W. von Schneider, Z. [2] 5,343).—4. By oxidising a-oxyn-caproic acid (Erlenmeyer, B. 9, 1840).—5. By heating \$\beta\$-acetyl-propionic acid with HI and P at 200° (Kehrer a. Tollens, A. 206, 233).—6. By reducing β-acetyl-propionic acid with sodiumamalgam (Wolff, A. 208, 110).—7. By heating n-propyl-malonic acid at 180° (Juslin, B. 17, 2504; Furth, M. 9, 308).—8. By fermentation of calcium lactate (Fitz, B. 13, 1309; 14, 1084). 9. By heating the lactone of γ-oxy-n-valeric acid with HIAq and P at 240° (Fittig, A. 226, 346).—10. A product of oxidation of castor oil by dilute HNO₃ (Wahlforss, B. 22 Ref., 438).

Properties.—Liquid, smelling like butyric acid.

Salts.-KA'. Plates (from alcohol).-CaA'2 aq. S. 10.267 at 0°; 8.144 at 60°. Least soluble at 60°-70° (F.);—CaA'₂1½aq (Schorlemmer, A. 161, 270).—BaA'₂. S. 21.693 at 0°; 23.076 at 80°.—MnA'2 aq.—CuA'2: minute green needles. More sol. cold than hot water. - ZnA'2. S. 2.6 at 25°.—AgA'. S. 229 at 3°; 641 at 70.5°.

Methyl ether MeA'. (127.3°). S.G. $\frac{0}{0}$.9097 (G.); $\frac{20}{0}$.8795. C.E. (0°-10°) .00106. S.V. 149·1 (Gartenmeister, A. 233, 273). $\mu_{\beta} = 1.3997$. $\mathbf{R}_{\infty} = 50.7.$

Ethyl ether EtA'. Mol. w. 130. cor.). S.G. § ·8939 (G.); 20 ·8765 (Lieben a. Rossi, A. 165, 117); 20 ·8661 (Brühl). S.V. 174-3. C.E. (0°-10°) ·00111. $\mu_{\theta} = 1.402$. R_{∞} 58.08 (B.).

n-Propyl ether PrA'. (167.5°). S.G. 8 *8888. C.E. (0°-10°) *00106 (G.). S.V. 197.8. n-Butyl ether C,H,A'. (185.8°). S.G.

n-Butyl ether C.H.A'. (185.8°). S.G. § .8847. C.E. (0°-10°) .00101. S.V. 222·1. n-Amyl ether C.H.A'. Mol. w. 172. (208.7°). S.G. § .8812 (G.).; § .8568 (Brühl). C.E. (0°-10°) .00097. S.V. 245·8. μ_β 1·417. R_∞ 80.48.

n-Hexyl ethor C.H.A. (223.8°). 8.G. g •8797. C.E. (0°-10°) ·00096. S.V. 272·0.

n-Heptyl ether C,H18A'. (248.6°). S.G. \$.8786. C.E. (0°-10°) .00090. S.V. 297·4.

n-Octyl ether C₂H₁,A'. (260·2°). S.G. § :8784. C.E. (0°-10°) :00088. S.V. 322·6.

Amide C₄H₂,CO.NH₂. Mol. w. 101. [116°].

Pearly plates (Weidel a. Ciamician, B. 13, 69).

V. sol. water, alcohol, and ether.

Nitrile C4H9.CN. n-Butyl cyanide. (140°). S.G. ² ·816 (Lieben a. Rossi, A. 158, 171). Formed, together with amylamine, by allowing a mixture of hexoic amide (1 mol.) and bromine 2 mols.) to run into a 10 p.c. solution of NaOH (Hofmann, B. 17, 1410).

Isovaleric acid (CH₃)₂CH.CH₂.CO₂H. Isopropyl-acetic acid. (176° cor.). S.G. $\frac{15}{16}$ 9336; $\frac{25}{25}$ 9261 (Perkin); $\frac{20}{10}$ 931 (E. a. H.). M.M. 5 635

at 16°. S. 4.24 at 20°.

Vapour tension: Richardson, C. J. 49, 767. Heat of neutralisation: Gal a. Werner, Bl.

[2] 46, 801.

Occurs in the fat of Delphinum Phocana (Chevreul [1817], Corps gras, pp. 99, 209), in valerian root (Grote, B. J. 11, 225; Trommsdorff a. Ettling, A. 4, 229; 6, 176), in the root of Angelica Archangelica (Meyer a. Zenner, A. 55, 328), in the root of Athamanta Oreoselinum, in the bark of the alder tree, in the berries and bark of the Guelder rose (Viburnum Opulus) (Moro, A. 55, 330), in the root of Viburnum prunifolium (Allen, Ph. [3] 11, 413), and in oil of geranium (O. Jacobsen, A. 157, 232). A valeric acid occurs in rosin oil (Lwoff, B. 20, 1017) and in yolk of wool (suint) (Buisine, Bl. [2] 48, 639).

Formation. -1. By oxidation of inactive fermentation amyl alcohol (Dumas a. Stas, A. 33, 156; 35, 143; Erlenmeyer a. Hell, A. 160, 257; Pedler, C. J. 21, 74).—2. By boiling its nitrile with alcoholic potash (E. a. H.; Schmidt a. Sachtleben, A. 193, 87).—3. Together with the active acid by oxidation of leucine (amido-hexoic acid) (E. a. H., cf. Neubauer, A. 106, 59).-4. By heating isopropyl-malonic acid at 180° (B. 11, 596).—5. By oxidising di-methyl-allyl-earbinol, treating the resulting CMe₂(OH).CH₂:CO₂H with HI, and reducing the β -iodo-valeric acid so formed by means of 3 p.c. sodium-amalgam applied in an acid solution (Schirokoff, J. pr. [2] 23, 286).—6. By the action of KOH or HClAq on Athamantin (Schnedermann a. Winckler, A. 51, 324).-7. By putrefactive fermentation of proteïds (Iljenko, A. 63, 269).—8. By the action of CrO_s on gelatin and albumen (Schlieper, A. 59, 7; Guckelberger, A. 64, 71).—9. From isopropylaceto-acetic ether (Frankland a. Duppa, A. 145, 84).-10. By distilling isopropyl-malonic acid (Conrad a. Bischoff, A. 204, 151).-11. A product of distillation of colophony (Renard, A. Ch. [6] 1, 253).

Preparation.-1. Isoamyl alcohol is oxidised by less than the calculated quantity of KMnO4 in the cold. Some acetic acid is formed, but this can be got rid of by distilling a dilute solution of the acids, for valeric acid passes over in the first quarter of the distillate, the acetic acid remaining behind (Duclaux, C. R. 105, 171).—2. By oxidising isoamyl alcohol (1,000 c.c.) with K2Cr2O, (1000 g.) dissolved in water (3,500 c.c.) by adding in the cold H_2SO_4 (1,400 c.c.)g.) mixed with water (800 g.). The resulting isoamyl isovalerate is saponified by potsch (Pierre a. Puchot, A. Ch. [4] 29, 229).—8. By

distilling valerian root with dilute phosphoric acid, neutralising the distillate with Na₂CO₃, and decomposing the resulting Na salt by H₂SO₄. In purifying valeric acid use may be made of the fact that the acid forms crystalline acid salts with K and Na (Lescour, Bl. [2] 27,

Properties.-Liquid, inactive to light. Its smell is powerful and unpleasant. Sl. sol. water. Separated from aqueous solution by CaCl₂. Volatile with steam. Mixes with alcohol and ether. It is set free from its salts by mineral acids, by HOAc, by oxalic, tartaric, citric, and malic acids, but not by butyric acid. When conc. valeric acid is added to a solution of cupric acetate, anhydrous cupric valerate separates in oily drops, which soon change to a greenish-blue crystalline powder of the hydrated salt; butyric acid would at once give a crystalline pp. (Larocque a. Huraut, J. Ph. [3] 9, 430).

Reactions.-1. Oxidised by chromic acid mixture at 20° to acetic acid and CO2.-2. Boiling dilute HNO, forms nitro-valeric acid, methyl-malic acid, and a little di-nitro-propane (when prepared from valerian) or di-nitrobutane (when prepared from isoamyl alcohol) (Bredt, B. 14, 1782; 15, 2319; cf. Dessaignes, A. 79, 374).—3. Dilute alkaline KMnO, forms CMe₂(OH).CH₂.CO₂H.—4. Electrolysis of the K salt in conc. solution yields octane C₂H₄Pr₂ (Kolbe, A. 69, 259).—5. Passed through a redhot tube it yields ethylene, propylene, butylene, and other hydrocarbons (Hofmann, C. J. 3, 121). 6. The Ca salt yields di-isobutyl ketone on distillation by itself, valeric aldehyde on distillation with calcium formate, and methyl isobutyl ketone on distillation with calcium acetate.

Salts.—NH,A'. V. sol. water and alcohol.
—(NH₄)H₂A'₃.—KA'. Deliquescent crystalline
mass. S. (alcohol) 26 at 20°. [140°].—
LiA' ½aq. Nodules.—TlA'. V. sol. water.—
CaA'₂ 3aq: needles (Barone, A. 165, 120; Schmidt a. Sachtleben, A. 193, 87; Schirokoff, J. pr. [2] 23, 286). S. 18.4 at 0° (Sedlitzky, M. 8, 563).—CaA'₂ 4aq.—CaA'₂ 5aq.—BaA'₂. Triclinic plates. S. 94 at 18°. S. (alcohol) 3°3 at 20°.—BaA'₂aq.—BaA'₂ 2aq.—SrA'₂.—MnA'₂ 2aq.—ZnA'₂ 2aq. Crystalline.—ZnA'₂. S. 1 in the cold. S. (80 p.c. alcohol) 1°7 in the cold; S. (ether) '2 in the cold; '5 at 35° (Wittstein, J. 1847, 557). — ZnA'₂12aq. — ZnA'₂3aq. — ZnA'₂2NH₃ (Lutschak, B. 5, 30).—CuA'₂. Green crystalline pp. — CuA'₂aq. — Bi₈A₁'(OH)₈O₃ (Schucht, Ar. Ph. [3] 2, 97).—FeA'₈.—FeA'₈(OH). -Fe,A',(OH)2.—PbA'2: easily fusible lamine. -Pb.A'202. Groups of needles, sl. sol. water.-AgA'. S. 177 at 0° (Sedlitzky); 185 at 20° (Erlenmeyer).

Methyl ether MeA'. (116°). S.G. 2 9007. C.E. (0°-10°) ·001174. S.V. 149·6 (Elsässer, A. 218, 315); 148·3 (R. Schiff, 4. 220, 334). S.H. (21°-45°) 491 (Kopp). Smells like

bananas.

Ethyl ether EtA'. (134°). (E.); (185° cor.) (Perkin). S.G. 2 ·8851 (E.); ‡ ·8714; 25 ·8632 (P.). C.E. (0°-10°) ·001034. S.V. 178·4 (E.); 178·0 (S.). M.M. 7·616 at 18°. Oil, sol. alcohol. Smells like rennet apples and water-mint (Pierre a. Puchot, A. Ch. [4] 20, 284). Converted by sodium into oxy-decoic acid, ethyl ethoxy-decoate (Hantzsch, A. 249, 64), an acid C₂₀H₂₄O₂ [c. 127°] (295°), and other bodies (Greiner, Z. [2] 2, 460; Wanklyn, C. J. 17, 871; Geuther a. Greiner, J. 1865, 819).

n-Propyl ether PrA'. (156°). S.G. 2. 8809. C.E. (0°-10°) 000997. S.V. 197.5 (Elsässer); 196.8 (Schiff).

Isopropyl ether PrA'. (142°). S.G. 2 '870; 11 '854 (Silva, 1. 153, 186).

Di-chloro-propyl ether C.H.Cl.A'. (245°) at 737 mm. S.G. 11 1149. Formed from epichlorhydrin and isovaleryl chloride CaHaClaA'.

(Truchot, A. 138, 298).

Isobutyl ether C,H,A'. (169°) (Elsässer, A. 218, 328); (173°) (P. a. P.). S.G. 2 8736 (E.). 2 8884 (P.a. P.). C.E. (0°-10°) 001027. S.V. 223·4 (E.).

Isoamyl ether C.H.A. (188°) (Kopp; R. Schiff, A. 234, 344); (196°) (Balard); (190°) (Balbiano, J. 1876, 348); (194°) (Kahlbaum). S.G. 2 870 (Balbiano, S.V. 244.5. V.D. 6.1. Smells, when dissolved in alcohol (7 pts.), like apples.

Octyl ether C₈H₁₇A'. (250°). S.G. 15

'862 (Zincke, A. 152, 6).

Cety l ether C₁₆H₃₃A'. [25°]. (280°-290°)
at 202 mm. S.G. 22 852 (Dollfus A. 131,

Allyl ether C₃H₅A'. (154°). Oil (Cahours

a. Hofmann, A. 102, 296).

Glyceryl derivative v. GLYCERIN. Chloride Pr.CH₂.COCl. (114°) at 726 mm. S.G. ²⁰ 9887. μ_θ 1.4213. R_∞ 49.14. Liquid, easily decomposed by water (Bechamp, C. R. 42, 224; Brühl, A. 203, 24).

Bromide C. H., OBr. (143°). Liquid.

Iodide C. H., OI. (168°) (Cahours, C. R.

44, 1252).

Anhydride (C,H,O),O. Mol. w. 186. (215°) (Chiozza, A. 84, 106). Converted by hydrated BaO2 into the oily peroxide (C5H2O)2O2 (Brodie, Pr. 12, 655).

Aceto-valeric anhydride (C,H,O).OAc.

(147°-160°) (Autenrieth, B. 20, 3187).

Amide C,H,CONH, [128°] (Letts, B. 5, 669; Hofmann, B. 15, 982); [135°] (Schmidt a. Sachtleben, A. 193, 102). (231°). Silky plates, v. sol. water and alcohol.

Anilide C,H,CO.NHPh. [115°] (Chiozza, A. Ch. [3] 39, 201; Kelbe, B. 16, 1200; cf. Dumas, C. R. 25, 475, 658; Dessaignes, A. 68, 333). Sl. sol. hot water, v. sol. alcohol and ether.

Pr.CH. CN. Isobutyl cyanide. (129°) (R. Schiff, B. 19, 567). V.D. 2.89. Formed by heating Pr.CH.CN. Nitrile Mol. w. 83. S.G. 2 ·823. ammonium valerate or valeramide with P2Os (Dumas, Malaguti a. Leblanc, C. R. 25, 658). Occurs among the products of oxidation of gelatin and casein by chromic acid mixture (Schlieper, A. 59, 15; Guckelberger, A. 64, 72), of the action of chlorine on leucine (Schwanert, A. 102, 228), and of the action of conc. HNO, on castor oil (Hell a. Kitrosky, B. 24, 980). Formed also, together with valeramide, by heating valeric acid with potassium sulphocyanide (Letts, B. 5, 669). It is a so got by the action of cold Ac.O on the oxim of valeric aldehyde (Dollfus, B. 25, 1915). Prepared by digesting isobutyl iodide with KCy and alcohol (Erlenmeyer a. Hell, 4. 160, 266). Liquid, smelling like almonds dissolving in about four times its volume of water.

Sodium converts it, on heating, into cyanbutine C13H27N2, crystallising in stellate groups of needles, which yield B'HCl and B', H, PtCl, and is converted by nitrous acid into C₁₅H₂₅(OH)N₂

[89°] (E. von Meyer, J. pr. [2] 37, 407).

Active valeric acid CHMeEt.CO., H. Methylethyl-acetic acid. Hydrotiglic acid. (177° i.V.). S.G. $\frac{24}{17\cdot5}$ ·938 (Saur). S.V. 129·1 (Lossen, A. 254, 60). Occurs in the essential oil from the fruit of Angelica Archangelica (R. Müller, B. 14, 2476).

Formation.—1. By oxidation of isoamyl alcohol .- 2. By reducing tiglic acid with HI and P (Schmidt a. Berendes, A. 191, 117).—3. By heating methyl-eth/l-malonic acid (Bischoff a. Conrad, A. 204, 151).—4. From methyl-ethylacetoacetic ether (Saur, A. 188, 257).—5. By reducing angelic acid (Schmidt, A. 208, 261).— 6. By reducing bromo-hydro-tiglic acid (Pagenstecher, A. 195, 171).—7. By the action of NaOEt and EtI on propionyl-propionic acid (Israel, A. 231, 219).—8. By oxidation of the corresponding aldehyde (Lieben a. Zeisel, M. 7, 56).

Properties.—The acid obtained by synthetic methods is inactive, but when obtained by oxidation of active amyl alcohol it is dextrorotatory, but is mixed with inactive isovaleric acid. By oxidation of an amyl alcohol $[a]_D = -4.4^{\circ}$ Guye and Chavanne (C. R. 116, 1454) obtained a valeric acid (174°), $[a]_D = +13.6^{\circ}$, S.G. $\frac{22}{29}$ 938. By oxidation of an amyl alcohol $[a]_D = -5 \cdot 2^\circ$ at 22° Rogers (C. J. 63, 1130) got a valerio acid (175°), S.G. $\frac{2^2}{2^2}$ 936, $[\alpha]_D = +13.9^\circ$ at 22°. Methyl-ethyl-acetic acid can be separated from isopropyl-acetic acid through the

greater solubility of its silver salt. Dilute KMnO, oxidises methyl-ethyl-acetic acid to CMeEt(OH).CO2H.

Salts.—CaA'₂5aq: needles. S. (of CaA'₂) 29.—*BaA'₂: gummy mass.—ZnA'₂. Needles, more sol. cold than hot water.—CuA'₂: bluishgreen, crystalline pp.—AgA': groups of needles. S. 1.13 at 20° (C. a. B.); 1.11 at 1° (Sedlitzky, M. 8, 568).

(133.5° i.V.). Ethyl ether EtA'. S.G.

Nitrile C, H,1N. (125°). S.G. º .8061. Formed by adding Na and EtI to a solution of acetonitrile in benzene (Hanriot a. Bouveault,

Bl. [2] 51, 173). Liquid.

Valeric acid CMe, CO,H. Tri-methyl-acetic acid. [35°]. (163·5° i.V.). S.G. 59·905. S. 2·2 at 20°. Heat of neutralisation: Gal a. Werner, Bl. [2] 46, 801. Formed by heating its nitrile with alcoholic potash or conc. HClAq at 100° (Butleroff, A. 165, 322; 170, 151; 173, 355; B. 5, 478). Formed also by action of CrO, on pinacolin (Friedel a. Silva, B. 6, 146, 826) and on tri-methyl-pyruvic acid (Glücksmann, M. 10, 777). Monometric crystals. Does not yield a bremo- derivative on heating with Br, either alone or in presence of P (Reformatzky, B. 23,

Salts.-NaA'2aq. Prisms, v. sol. water.-KH.A's. Needles, m. sol. water. - CaA', 5aq. CaA', 4aq. S. 7 at 1°.—SrA', 5aq.—BaA', 5aq. S. 34 at 2° (Landau, M. 14, 707).—MgA', 8aq. —ZnA', 2aq. S. (of ZnA',) 1.7 at 20°. The cold with a solid salt, which dissolves up again on cooling.—PbA'₂.—PbHA'₃: needles.—CuA'aq: greenish pp.—AgA'. S. 1·1 at 1°; 1·27 at 27° (Stiassny, M. 12, 599).

(101°) (Butleroff). (118.5° i.V.). S Methylether MeA'. Ethyl ether EtA'. 9 ⋅875.

Tert-butylether C, H, A'. (135° i.V.). Amyl ether CMe, CH, A'. (165°) (Tissier, Bl. [2] 24, 558).

Chloride CMe, COCI. (106°). Anhydride (CMe, CO), O. (190°).

Amide CMeg.CO.NH2. [1549]. (212°). Formed by heating the ammonium salt in sealed tubes at 230° (Franchimont a. Klobbie, R. T. C. 6, 238). Long needles. Decomposed by pure

HNO₃ (S.G. 1.58) with evolution of N₂O.

Methylamide CMe₃.CO.NHMe. [91°].

(204°).

Dimethylamide CMe3.CO.NMe2. (186°). Ethylamide CMe, CO.NHEt. [49°]. (204°). Diethylamide CMeg.CO.NEt2. S.G. 15 ·891.

Nitrile CMe, CN. Tert-butyl cyanide. [16°]. (106°). Formed, together with a polymeride (160°), by the action of tert-butyl iodide on dry potassio-mercuric cyanide below 5° (Butleroff, A. 170, 151; Freund a. Lenze, B. 23, 2866; 24, 2161). Crystalline mass with pungent smell. On heating with aniline hydrochloride it yields phenyl-valeramidine, which forms an oxalate [192°]. Hydroxylamine forms the amidoxim CMe₃.C(NH₂):NOH [116°].

References.—Amido-, Bromo-, Chloro-, Iodo-,

Nitro-, Nitroso-, and Oxy- valeric acids. n-VALERIC ALDEHYDE

CH₂·CH₂·CH₂·CH₂·CHO. (103°). S.G. 11 ·819. Formed by distilling calcium n-valerate with calcium formate (Lieben a. Rossi, A. 159, 70; Zander, A. 224, 81).

Isovaleric aldehyde (CH₃)₂CH.CH₂.CHO. Valeral. Mol. w. 86. (92·5°). S.G. ½5 ·8041; ½½ ·7961 (Perkin, C. J. 45, 477). μ_β 1·3934. R. 40·66 (Brühl). V.D. 43·06 (calc. 43) (Schröder, B. 4, 400, 468). S.V. 118.5. Formed by oxidation of isoamyl alcohol (Dumas a. Stas, A. Ch. [2] 73, 145; Parkinson, A. 90, 114; Kolbe a. Guthrie, A. 109, 296). Formed also by distilling calcium iso-valerate with calcium formate (Limpricht, A. 97, 370), by the distilla-tion of isovalerates (Chancel, A. 60, 318; Ebersbach, A. 106, 262; Schmidt, B. 5, 600), by oxidation of gluten (Keller, A. 72, 34) and castor oil (Arzbächer, A. 73, 202), and by the action of SO₃ on leucine (Schwanert, A. 102, 226).

Properties .- Neutral oil, with pungent fruity odour. Polymerises on keeping. Mixes with alcohol and ether. On heating with ammonium sulphocyanide it yields a dark-red liquid, sol. alcohol, ether, and HOAc (Brodsky, M. 8, 37). Alkaline sodium nitroprusside gives a violet-red colour, destroyed by HOAc (Von Bitto, A. 267, 876). Combines with sodium bisulphite, forming (C₅H₁₀O)NaHSO₃.

Reactions .- 1. Readily oxidised to valerie acid .- 2. Chlorine forms mono- and di- chlorovaleric aldehydes, and, at 140°, oily C₁₀H₁₂Cl₂O (204°), which is converted by alcoholic soda into C1. H1. Cl.O (209°).—8. PCl. forms C. H1. Cl. —ZnA', aq. S. (of ZnA',) 1.7 at 20°. The cold 4. Potash-fusion gives valeric acid.—5. Heat-saturated solution becomes filled on warming ing with lime forms isoamyl alcohol, calcium

valerate and other bodies (Fittig, A. 117, 68).— 6. Sodium-amalgam and water slowly form isoamyl alcohol (Wurtz, A. 134, 201).-7. Sodium forms isoamyl alcohol, sodium valerate, decyl alcohol $C_{10}H_{20}O$ (203°), an oil $(C_{10}H_{16}O)_{20}$ (250°–290°), the compound $C_{20}H_{28}O_{3}$, and the acid $C_{10}H_{18}O_{2}$ (Borodin, Z. 1864, 353; B. 5, 480).— 8. KOH at 0° produces an oily polymeride, lighter than water, not capable of combining with NaHSO₃, and yielding isovaleric aldehyde and the compounds C₁₀H₁₈O and C₂₀H₃₈O₃ on distillation (Borodin, B. 6, 982). This polymeride, left in contact with Na₂CO₃Aq, often forms needles of $C_{20}H_{42}O_{5}$ [70°].—9. Heating with KOH forms the addenged $C_{10}H_{10}O$ and a compound C₂₀H₃₀O₃ (260°-290°). S.G. '90.—
10. Heating with *sinc* at 180° forms decenoic aldehyde C₁₀H₁₀O (190°), S.G. ² ·862 (Riban, C. R. 75, 96; Kekulé, A. 162, 77) and other products.—11. ZnEt, produces the oil (C10H18O), in large quantity (Beilstein a. Rieth, A. 126, 242). ZnMe2, followed by water, forms methyl-isobutylcarbinol (Kuvsinoff, J. R. 1887, 204).-12. Dry K₂CO₂ at 50° forms an oily polymeride, which is reconverted at 180° into the aldehyde. Boiling with K₂CO₂ yields C_{1c}H_{1s}O₂ (v. supra), C_{2s}H_{2s}O₂ (265°-270°), and C_{1.1}H_{2s}O₂ (235°-240°) (Gäss a. Hell, B. 8, 369).—13. Dry K₂CO₂ added to moist isovaleric aldehyde forms, after some days, crystals of a polymeride [84°], which is reconverted at 108° into the original aldehyde (Bruylants, B. 8, 414).—14. PH,I forms a compound (O₄H₁₀O),PH,I, orystallising from ether in plates [119°], converted by cold potash into (C₅H₁₀O)₅PH,OH, crystallising in prisms [125°], insol. water (De Girard, A. Ch. [6] 2, 33; C. R. 94, 215).—15. On heating with hypophosphorous acid in an atmosphere of hydrogen the compound (C₂H₁₀O)₂H₃PO₂ [160°] is formed. It yields BaA'₂aq (Ville, C. R. 109, 71).—16. AcCl at 100° combines, forming chloro amyl acetate C4H2.CHCl.OAc (Maxwell Simpson, Pr. 27, 120). 17. m-Amido-benzoic acid forms C12H15NO2 [c. 130°].—18. Dry HCl forms (C,H,CHCl)2O (180°) (Bruylants, B. 8, 414).—19. Conc. NH, Aq forms isovaleric aldehyde ammonia C_4H_6 .CH(NH₂)OH 7aq [58°] (Erdmann, A. 130, 211; Petersen, A. 132, 158; Ljubavin, B. 6, 1460), which yields, on allowing its alcoholic solution to stand, or on boiling with potash, the oily 'trioxyamylamine' C15H28NO3, forming B'HCl [113°] and B'₂H₂PtCl₂. AgNO₃ reacts with isovaleric aldehyde-ammonia, forming (C₃H₁₁N)₂AgNO₃ and (C₃H₁₁N)₂AgNO₃ (Goldschmidt, B. 11, 1200; Mixter, J. 1878, 438).— 20. Alcoholic NH, at 150° forms valeritrine, Valerihydrovaleritrine, and other bodies. trine $C_{1,}H_{2,1}N$ is a very unstable liquid (250°–260°), yielding B'HHgCl₂ [88°], B',H₂PtCl₃, and B'C₂H₂N₃O, [130°]. Hydrovaleritrine $C_{1,}H_{2,}N$ or $C_{1,}H_{3,}N$ is a liquid, yielding crystalling. line B'HOl. S. 2.87 at 22°, not melted at 180° (Ljubavin, B. 6, 565).—21. H₂S passed into an aqueous solution forms thiovaleric a'dehyde C.H., S [69°]. An isomeride (115°) is got, together with a polymeride [94.5°], by heating isovaleric aldehyde with sulphur at 250° (Barbaglia, B. 13, 1574; 17, 2664; G. 11, 95; 16, 426).—22. CS₂ and NH₂Aq form carbovaleraldine C₁₁H₂₂N₂S₂ (Schröder, B. 4, 469; Mulder, A. 168, 237), which melts at 109°

(Guareschi, A. 222, 811).-28. NH2.CS.OEt and HCl form $C_1H_{10}(NH.OS.OEt)$, [108°] (Bischoff, B. 7, 1078).—24. H.Se passed into an aqueous solution of valeric aldehyde forms $C_5H_{10}Se$ [56.5°].—25. Acetone and dilute NaOH yield Pr.CH2.CH:CH.CO.CH, (180°) (Barbier a. Bouveault, C. R. 118, 198)

Oxim C,H,.CH:NOH. Mol. w. 101. (163°). S.G. 30 ·8934 (in vacuo). µD 1·437 (Petraczek, B. 16, 829; Trapesonzjanz, B. 26, 1428). It is a syn- oxim, because it yields valeronitrile when mixed with Ac.O and ether (Dollfus, B. 25, 1915).

Isovaleric orthaldehyde. Amylidene glycol. Acetyl derivative C.H. CH(OAc)₂. (195°). Formed by heating isovaleric aldehyde with Ac.O at 200°.

Bensoyl derivative C₄H_e.CH(OBz)₂. [111°]. (264°). Formed by heating the aldehyde with Bz2O at 260° (Guthrie a. Ko.be, A. 109, 298).

Di-methyl ether C. 1. CH(OMe). (124°). S.G. 10 852. Formed from isovaleric aldehyde (2 vols.), MeOH (5 vols.), and HOAc (1 vol.) (Alsberg, J. 1864, 486).

Di-ethyl ether C₄H₂.CH(OEt)₂. S.G. ¹²·835. Formed in like manner. (168°).

Di · isoamyl ether $C_4H_9.OH(\cup C_8H_{11})_2$ (c. 248°). S.G. $\stackrel{?}{\cancel{2}}$ *849.

Ethyl isoamyl ether C₄H_p.CH(OEt)(OC₅H₁₁). (c. 205°). S.G. 13 ·875. Formed by the action of Na on an ethereal solution of isovaleric ether (Greiner, Z. 1866, 465).

Valeric aldehyde CHMeEt.CHO. Formed by reducing tiglic aldehyde with iron and HOAc (Herzig, M. 3, 123; Lieben a. Zeisel, M. 7, 56), and by warming CH₂:CH.CHEt.OH with dilute (1 p.c.) H₂SO₄ (Kondakoff, J. R. 20,

Valeric aldehyde CMe_s.CHO. S.G. 18 .7927 (Tissier, Bl. [2] 24, 558).

Di - isovaleric aldehyde v. DECENOIC ALDE-

Reference. - DI-BROMO-, CHLORO-, IODO-, and OXY- VALERIC ALDEHYDE.

VALEROGUANAMINE C,H,,N, i.e.

ProH₂.C:N.C(NH) NH.C(NH)>NH. Butylguanamine. [173°]. Formed by heating guanidine isovalerate Bandrowski, B. 9, 240; Haaf, J. pr. [2] 43, 76). Flat plates (from water). Weak base.—B'HCl: needles, v. sol. water.—B'H₂SO₄.—B'AgNO₅.

VALEROLACTIDE v. OXY-VALERIC ACID.

VALEROLACTONE v. OXY-VALERIO ACID. VALEBONE v. DI-ISOBUTYL KETONE. VALERONITRILE v. Nitrile of VALERIC

ACID VALEROVALERIC ACID VALERYL.

DIVALERYL C10H14O2 i.s. C4H2.CO.CO.C4H2. (270°-280°). Formed by the action of Na on valeryl chloride (Brühl, B. 12, 815). Oil.

VALERYL - ACETOPHENONE v. PHENEL BUTYL METHYLENE KETONE.

VALERYL CHLORIDE v. Chloride of VALERIC ACID.

VALERYL-CYANAMIDE v. CYANIC ACIDS. VALERYLENE v. PENTINENES and also Bromo- and DI-CHLORO-VALEBYLENE.

VALERYLENE TETRABBOMIDE v. Tetra-BROMO-PENTANE.

VALERYLENE HYDRATE v. PENTENYL |

VALERYLENE HYDROCHLORIDE v. CHLORO-AMYLENE.

VALERYL-VALERIC ACID. Ethyl ether $C_{12}H_{22}O_3$ i.e. CH_2 Pr.CO.CHPr.CO₂Et. (206°) at 722 mm. Formed by the action of sodium on isovaleric ether (Greiner, Z. 1866, 461; Wohlbruck, B. 20, 2335). Oil, sol. alcohol and ether.

VALYLENE C₅H₆. Mol. w. 66. (50°). A product of the action of alcoholic potash on valerylene dibromide (Reboul, A. 185, 872). Liquid, with odour of garlic and prussic acid. Ammoniacal Cu₂Cl₂ ppts. yellow C₅H₅Cu, which yields the pure hydrocarbon on treatment with dilute HClA₆. Bromine at 0° forms crystalline C₅H₆Br_o. Ammoniacal AgNO₅ yields a white pp. of C₅H₅Ag.

VANADATES v. "ANADIUM OXYACIDS, AND SALTS AND DERIVATIVES THEREOF, p. 851.

VANADIC ACIDS v. VANADIUM OXYACIDS, p. 851.

VANADIUM. V. At. w. 51.2. Mol. w. not known. Melts at a very high temperature; probably higher than m.p. of Mo, which does not melt at 1700°-1800° (Moissan, C. R. 116, 1225). S.G. 5.5 at 15° (Roscoe, T. 1869. 679). For emission-spectrum, obtained by using electric

sparks, v. Thalén (A. Ch. [4] 18, 243).

Historical.-In 1801 Del Rio announced the discovery of a new metal in a lead ore from Zimapan in Mexico (G. A. 71, 7); to this metal he gave the name erythronium, but at a later time he thought the metal was only impure chromium. In 1830 Sefström found a new element in the bar-iron and refinery slags where iron-ore from Taberg in Sweden was smelted (P. 21, 43); this element he called vanadium (from a Scandinavian deity Vanadis). In the same year (P. 21, 49) Wöhler found that the lead ore examined in 1801 by Del Rio contained lead vanadate; and he showed that the metal which Del Rio had called erythronium, and had then supposed to be impure chromium, was really vanadium. In 1831 Berzelius supposed he had prepared vanadium by heating an oxychloride in NH, and also by heating the highest oxide with K (P. 22, 1); but in 1867 Roscoe (T. 1868. 1; 1869. 679; v. also 1870. 317) showed that one of the substances which until then had been taken to be the element V was a nitride, and that another substance, taken to be V, was an oxide of this element. Roscoe obtained pure V by heating VCl₂ to redness for many hours in dry H. Roscoe was led to doubt the accuracy of the statement of the properties and at. w. of V that had been accepted on the authority of Berzelius, because he found that several compounds of V and Pb were isomorphous, but that it was impossible to assign comparable formulæ to these compounds if the at. w. given by Berzelius for V were accepted (v. T. 1868. 1 et seg.). Berzelius gave the value 68.5 to the at. w. of V; he assigned the formulæ VO, VO2, and VO. (0=8) to the three oxides, and classed V with Or. Mo. and W. Roscoe's researches showed that the at. vs. of the metal is 51-2, that the oxides are best represented as V_2O_3 , V_2O_4 , and V_2O_{34} that the substance thought to be V was really chiefly V_2O_3 , that the compound supposed by Berzelius to be VCl_2 was $VOCl_3$, and that Vmust be classed with P and As.

Occurrence.—Small quantities of compounds of V are found fairly widely distributed; the element does not occur uncombined. The chief minerals that contain V compounds are mottramite (Cu-Pb vanadate), descloisite (Pb-Zn vanadate), dechenite (Pb-Zn-Cu vanadate), roscoeite (Al vanadate with K silicate), vanadinite (vanadate of lead with lead chloride), and some other minerals, which generally contain vanadates of Cu. Pb, Zn, or Ca. Small quantities of compounds of V are found in all pisolitic limonites (Böttger, C. C. 1873. 514), in some pitchblendes, clays, and basalts, and in a few other rocks. Compounds of V have also been found, in small quantities, in the slag from copper extraction works (Karsten, P. 52, 629; Witz a. Osmond; Bl. [2] 88, 49); in many specimens of pig-iron (v. Riley, C. J. 17, 21; cf. Hodges, C. N. 26, 238; Walz, Am. Ch. 6, 453); in some meteorites (v. Apjohn, C. J. 27, 104); in caustic soda and sodium phosphate (Schöne a. Rammelsberg, B. B. 1864. 681; Donath, D. P. J. 240, 318); and, according to Lockyer (Pr. 27, 279), probably in the sun.

Preparation.—1. From the refinery slag of the Taberg iron-ore. The finely-pow-dered slag is heated with KNO, and Na, CO, the fused substance is digested with boiling water, and solution of a salt of Ba or Pb is added to the filtered liquid; the ppd. Ba or Pb vanadate is decomposed by boiling with a slight excess of H₂SO₄Aq, and the solution is filtered from BaSO₄ or PbSO.; the filtrate is neutralised by NH, Aq and concentrated, and pieces of NH₄Cl are placed in the liquid until some NH,Cl remains undissolved; the NH4VO, which is thus ppd. as a white crystalline powder is washed with conc. NH, ClAq, and then with alcohol, and is then decomposed by heating strongly in an open vessel, whereby V₂O₅ is produced (Berzelius, P. 22, 1). Wöhler (A. 78, 125) employed a very similar process for preparing V₂O₅ from limonite.—2. From vanadinite. L'Hôte (C. R. 101, 1151) recommends to mix finely-powdered vanadinite (approximately 3Pb, V,O, PbCl₂) with four times its weight of lampblack and a little oil, to heat strongly in a closed vessel, and then to heat to 300° in a stream of dry Cl, whereby VOCl, is formed, and distilled over into a cooled U-tube; other compounds of V are readily prepared from VOCl. -3. From the cobalt-ore of Mottram in Cheshire. This source of V compounds was discovered by Roscoe (v. C. J. [2] 6, 326). The Cheshire Keuper sandstone contains carbonates of Cu, carbonate of Pb and galena, black cobalt-ochre, iron oxides, and salts of As, Ag, Mn, and Ba. The compound of V was probably vanadinite. The sandstone was crushed, and the metallic compounds were dissolved by HClAq; bleaching-powder and milk of lime were added till the reaction was alkaline; and it was from the pp. thus obtained that Roscoe prepared V and its compounds. The lime pp. contained chiefly As, Cu, Fe, Pb, V, and Ca in combination with SO₄ and PO₄; it was heated in a furnace with ground coal to remove As, then roasted with a quarter its weight of Na,CO,

and lixiviated with water; the solution was saturated with H₂S and filtered, and H₂S was boiled off; crude oxide of V was then ppd. by NH₂Aq; the oxide was dried and heated with conc. HNO₂Aq to oxidise it to vanadic acid, which was then boiled with saturated (NH₄)₂CO₂Aq, and the somewhat soluble NH₄ vanadate was crystallised from water, and then strongly heated in an open vessel; the V₂O₃ thus obtained was suspended in water into which NH₄ was passed; the solution of NH₄ vanadate was filtered from silics, phosphates, &c., evaporated to dryness, and heated in an open vessel

until V2O, remained. The metal vanadium is prepared by reducing VCl₂ in H (for preparation of VCl₂ v. Vanadium DICHLORIDE, p. 844). The process is conducted in perfectly pure and dry H; every trace of moisture and air must be excluded. The pure and dry H is passed for 12 hours through a porcelain tube, connected (by specially arranged paraffin-joints) by a glass tube to the H apparatus; the VCl2 is then placed in the porcelain tube (a special arrangement for doing this is described by Roscoe), and H is passed through the apparatus for 6 hours. The porcelain tube is then gradually heated to full redness, and the passage of H is continued so long as HCl is given off, after which the tube is allowed to cool for several hours in the stream of H. The process occupies from 40 to 80 hours according as from 1 to 4 g. of VCl₂ are used. The V thus obtained contains traces of H and O. Reduction proceeds more quickly at a white heat, but the product is not so pure; it generally contains c. 95 to 96 p.c. V. (For details, and figures of the apparatus, v. Roscoe, T. 1869. 679; 1870. 317; or C. J. [2] 8, 344; 9, 23.)
Moissan (C. R. 116, 1225) obtained V con-

Moissan (C. R. 116, 1225) obtained V containing from 17.5 to 25.7 p.c. C, by heating a mixture of C and one of the oxides of V in an electric furnace, using a current of 70 volts and 1,000 ampères; with a current of 70 volts and 350 ampères reduction proceeded very slowly.

350 ampères reduction proceeded very slowly. Vanadium, mixed with V_2O_3 , was obtained by Roscoe (*l.c.*) by strongly heating a mixture of one of the chlorides of V with Nain an atmosphere of H, and lixiviating with water; the admixed black, powdery V_2O_3 can be removed from the heavier, grey, lustrous, metallic powder, by repeated washings with water, and a substance containing c. 91 p.c. V can be obtained.

The substance supposed by Berzelius (P. 22, 1) to be V, and obtained by heating V₂O₅ with K, was shown by Roscoe to consist chiefly of V₂O₂. The black powder which Uhrlaub (P. 103, 134) prepared by heating V oxychloride in NH₅, and which he supposed to be V, was found by Schafarik (W. A. B. 33, 5) to be VN₂ (cf. Roscoe, l.c.).

Properties.—A light-grey, lustrous powder; under the microscope it is seen to be composed of crystalline, silver-white particles. Does not become coherent and dense when strongly compressed. S.G. 5.5 at 15°. Not magnetic. Is not oxidised in air at the ordinary temperature, nor by moistening and then drying in vacuo. Has not been melted, except perhaps when reduced from the oxides by C in an electric furnace (v. Moissan, C. R. 116, 1225). When powdered V is thrown into a Bunsen flame it burns brilliantly; when strongly heated in O it forms

V₂O₅; and when slowly heated in air it perhaps forms V₂O, then V₂O₃, V₂O₄, and finally V₂O₃. Burns in Cl, forming VCl₄; combines with N to form VN; also combines with S. V is insoluble in HClAq; it dissolves in hot conc. H₂SO₄ and in HNO₃Aq. V dissolves in molten NaOH, giving off H, and for ling a vanadate. V acts on glass vessels in which it is heated, forming a compound with Si; it also alloys with

The at. w. of V has been determined (1) by reducing V₂O₂ to V₂O₃ by heating in H (Berzelius, P. 22, 15 [1831]; Roscoe, T. 1868. 8); (2) by oxidising V₂O₃ to V₂O₃ by HNO₄ (B., Lc., [1831]); (3) by analysing (V₂O₂)(SO₄). 4H₂O (B., Lc., p. 18 [1831]); (4) by ppg. VOCl₃ by AgNO₃Aq, removing excess of Ag from the filtrate, evaporating to dryness, heating, and weighing the V₂O₃ produced (B., Lc. [1831]); (5) by determining the weight of Ag required to ppt. the Cl from VOCl₃; also by weighing the AgCl ppd.; also by filtering from AgCl, evaporating, and weighing the V₂O₃ obtained (Roscoe, T. 1868. 23); (6) by determining V.D.s of, and analysing, VCl₄ and VOCl₃ (g. v.). The results obtained by Berzelius (with the necessary corrections on account of the wrong formulæ used by him; v. Roscoe, Lc.) gave values for at. w. of V varying from 49·3 to 52·5; Roscoe's values from the reduction of V₂O₅ to V₂O₅ varied from 51·13 to 51·35, and from the analyses of VOCl₃ from 50·32 to 51·877. The S.H. of V has not been determined.

Vanadium is both metallic and non-metallic in its reactions. V_2O_4 interacts with strong acids to form divanadyl salts, $V_2O_2(SO_4)_2$, &c.; hypovanadates $V_4O_2(OM)_2$ are also derived from a hydrate of V_2O_4 ; V_2O_5 interacts with conc. H_2SO_4 to form $V_2O_5.3SO_8$ and $V_2O_5.2SO_3.xH_2O$; V_2O_5 also forms $V_2O_5.H_2O$ (= HVO_3) and $V_2O_5.2H_2O$ (= $H_4V_2O_7$), which react as acids, and salts of the acid H_3VO_4 are also known; V_2O_5 also combines with various acidic oxides (P_2O_5 , MoO_8 , &c.) and basic oxides, to form complex, salt-like compounds. The sulphides of V_2O_5 dissolve in alkali sulphide solutions; several thiovanadates have been isolated, e.g. (NH_4), VS_4 ; and also thio-oxyvanadates, e.g. Na_3VS_2O . The haloid compounds of V_2O_3 are generally decomposed

by water to oxyhaloid compounds.

Vanadium is placed in Group V., with N, P, As, Nb, Sb, Di, Er, Ta and Bi. V is the second member of the even series family of this group, the family consisting of N, V, Nb, Ta and an unknown element with an at. w. between those of Th (= 232) and U (= 239). Group V does not show a marked division into families; the gradation of properties from N to Bi is fairly regular, nevertheless the families (1) N and P, (2) V, Nb and Ta, (3) As, Sb and Bi are distinctly indicated. No hydrides of members of the vanadium family have been isolated with certainty. The highest oxides, M₂O₈, are saltforming; all of them interact with alkalis to form salts containing the elements M in the acidic radicles; V₂O₃ also forms basic salts (VO)₂R_p, with strong acids; normal salts have not been obtained by the reactions of M₂O₈ with acids. Oxyacids of V, viz. HVO₃ and H₄V₂O₇, have been isolated, but definite cxyacids of Nb and Ta form chlorides, corresponding with their highest

oxides M_2O_a ; but the highest haloid compound of V is VX₄, corresponding with V_2O_4 ; oxyhaloid compounds of V of the form of VX₅ exist (VOCl₅, VOBr₅) (v. Nitrogen group of elec-

MENTS, vol. iii. p. 571).

In considering the relations between vanadium and the elements of Group V., the position of V in series 4 must be taken into account: as compared with the elements that form series 3, in which series P is placed, the elements of the vanadium series are metallic (V is preceded by Ti, Sc, Ca and K, and is followed by Cr, Mn, and the iron elements); the elements of the vanadium series are nearly as metallic, and also as non-metallic, as Rb, Sr, Y, Nb, Mo, -, and the Ru metals, which elements form series 6, i.e. the niobium series. Hence V would be expected to be distinctly more like a metal, in its chemical relations, than P, and about equal to Nb in its metallic and no. metallic tendencies. is the second member of the even series family of Group V., it might be expected to resemble N, which is the first of the even series elements of this group; but it is to be remembered that the differences between the first and second even series members of the groups (i.e. the differences between Li and K, Be and Ca, B and Sc, C and Ti, N and V, O and Cr, F and Mn), become more marked in passing from Group I. to Group VII., and it is also to be remembered that the resemblances between the first even series members and the succeeding odd series members, become more marked as the atomic weights increase; F is more like Cl and Br than O is like S, Se, and Te; but O resembles S, Se, and Te more than N resembles P, As, Sb, &c.; and C is more like Si, Ge, Sn, &c., than B is like Al, Ga, In, &c.; but C less resembles Si, &c., than N resembles P. &c.

Reactions and Combinations.—(Roscoe, T. 1869. 679; 1870. 317.) 1. Heated in air, V burns to V₂O₅; lower oxides are first formed if the temperature is low.—2. Does not decompose water at 100°.—8. Heated to redness in chlorine burns to VCl₄; heated in bromine produces VBr₅; does not combine with iodine.—4. Heated with excess of sulphur, V forms V₂S₅ (v. Kay, C. J. 37, 728).—5. Heating in a stream of pure nitrogen produces VN.—6. V combines directly with silicon and also with platinum.—7. Molten potash or soda dissolves V, forming an alkali vanadate and giving off H.—8. V dissolves in conc. sulphuric acid (? forming a vanadyl salt); also in nitric acid, probably forming a basic nitrate; also, slowly, in hydrofluoric acid

(? forming VF_s).

Detection and Estimation.—Small quantities of compounds of V give a colourless borax or microcosmic salt bead in the outer flame, becoming green in the reducing flame; if much V compound is present the bead is yellow in the outer flame. Solutions of V₂O₅ in acids are not ppd. by H₂S, but are reduced to solutions of V₂O₂ which are lavender-blue; if reduction is effected by Zn, Cd, or Na-amalgam the liquid becomes green, then blue, and finally lavender-blue; after reduction the solution bleaches indigo very rapidly, and soon absorbs O. Solutions of vanadates, either neutral or acidified by accetic acid, give finely-divided black pps. with tincture of galls, and the liquids appear like ink.

When an acidified solution of a vanadate is shaken with an ethereal solution of H_2O_2 , the latter solution becomes dark red (Werther, J.pr. 88, 195).

Variadium is generally estimated by converting it into an ammonium salt, strongly heating this in air, and either weighing the V_2O_3 thus produced or heating in H and weighing the V_2O_3 that remains. For a volumetric method of estimating V v. Gerlach $(B.\ 10,\ 1216)$.

Vanadium, acids of, v. Vanadium oxyacids, and

SALTS AND DERIVATIVES THEREOF, p. 851.

Vanadium, bromides of. Only one bromide

of V has been obtained.

Vanadium tribromide VBr_s. Mol. w. not known; from analogy of PBr3, AsBr3, &c., the formula VBr, is probably molecular. Formed by heating V or VN to redness in vapour of Br. Also by passing Br vapour over a mixture of V₂O₃ and C kept at a red heat; VOBr₃ is formed, then VOBr2, and then VBr3 sublimes (Roscoe, T. 1870. 317). Roscoe (l.c.) recommends to prepare VBr_s by placing pure VN in a porcelain boat in a tube of hard glass, driving out all air by dry CO₂, then passing vapour of Br through the tube while that part where the boat is placed is heated to redness, when brown vapours are given off and condense on the cooler parts of the tube, and finally driving out all traces of Br by a stream of dry CO₂. VBr₃ is a greyish black, opaque, amorphous, very deliquescent solid; loses Br even in a sealed tube; heated gently in air gives V₂O₃, and at a higher temperature V₂O₃. Dissolves in water, without giving off any Br, forming a brown liquid which goes green on addition of a few drops of HClAq; this reaction is said by R. to be characteristic of a salt of V₂O₃.

Vanadium, chlorides of. Three chlorides have been isolated: VCl₂, VCl₃, and VCl₄. V burns when heated in Cl, forming VCl₄, and the other chlorides are formed by reducing VCl₄ by H. The V.D. of VCl₄ has been determined, and the formula VCl₄ is molecular; the formula VCl₅ is probably molecular, from the analogy of AsCl₅, PCl₃, BiCl₃, &c.; VCl₂, AsI₂, PI₂ and BiCl₅ are the only haloid compounds of the form MX that have been isolated in Group V.; the mol. w.s of these compounds have not been deter-

mined.

Vanadium tetrachloride VCl₄. Mol. w. 192.68. Prepared by Roscoe (T. 1869. 691). Formation.—1. V or VN is heated in Cl.—

Formation.—1. V or VN is heated in Ol.—2. The vapour of VOCl₃, mixed with excess of Cl, is passed slowly over a long layer of sugar-

charcoal heated to redness.

Preparation.—VN is heated to redness in a tube of hard glass, in a current of pure, dry Cl, which must be free from air. The first few drops of distillate are rejected, as they contain traces of VOCl, formed from traces of oxide in the nitride used. The dark-red liquid that is formed is saturated with dry Cl, then heated for some hours in a current of dry CO₂ in a vessel connected with an upright condenser, and then fractionated.

Properties and Reactions.—A thick, dark, brownish-red liquid; boils at 154° under 760 mm. pressure. S.G. 1.8584 at 0°, 1.8363 at 8°, 1.8159 at 30°. V.D. 98.1 at c. 200°. Does not solidify at any temperature above —18°. Slowly decomposes to VCl₂ and Cl at ordinary tempera-

tures, more quickly when exposed to light, and | rapidly when heated. Gives off white fumes in the air. Dissolves in water, forming a blue solution, which does not bleach; this solution is oxidised by KMnO₄Aq to HClAq and V₂O₅Aq, 2VCl, taking up O. Reacts violently with alcohol and ether, forming deep-coloured liquids. When heated with Br in a sealed tube, VCl, is formed. VCl, does not combine with Cl to form a higher chloride. H2S reacts with VCl, to form

VCl₃, HCl, and S (Kay, C. J. 38, 736).

VANADIUM TRICHLORIDE VCl₃. M Mol. w. not determined, but formula is probably molecular from analogy of PCl2, AsCl2, &c. Prepared by heating VCl, in a retort as long as Cl is given off, and heating the residual solid to 160° in a stream of dry CO_2 (Roscoe, T. 1869. 691). Also by heating pure V_2S_3 in a stream of dry, air-free Cl, separating S_2Ol_2 by repeatedly distilling the liquid product, and heating the residue at 140° in dry CO, to remove the last traces of S₂Cl₂ (Halberstadt, B. 15, 1619). Lustrous, peachblossom coloured tablets, resembling CrCl,; S.G. 3.0 at 18°. Deliquesces very rapidly in air to a brown liquid, which goes green on addition of a drop or two of HClAq; this solution is oxidised by KMnO₄Aq, 2VCl₂ taking up 2O, to HClAq and V₂O₂Aq. Soluble in alcohol and in ether. Heated in air forms V₂O₅. Heated in H is reduced to VCl2 and then, gradually, to V.

Vanadium dichloride VCl₂. Mol. w. not known. Formula possibly V₂Cl₂. Prepared by passing vapour of VCl₄, mixed with a large excess of pure, dry H, through a glass tube heated to dull redness. The distillation of VCl₄ should be slow, and the mixture of vapour of VCl, and H should be brought at once into the heated part of the tube. Forms apple-green, micaceous, hexagonal plates; S.G. 3.23 at 18°; very hygroscopic. Dissolves in water, forming a violet solution, which bleaches indigo and litmus, and reacts as a solution of a salt of V2O2. Dissolves in alcohol and in ether. The solution in water is oxidised by KMnO₄Aq to HClAq and V₂O₅Aq, 2VCl₂ taking up 3O. Heated in H, is slowly reduced to V; heating to whiteness in NH, produces VN (Roscoe, l.c.).

Vanadium, ferrocyanide of, v. vol. ii. p. 337.

Vanadium, fluorides of. Only one fluoride, VF, has been isolated; there are indications of the existence of a higher fluoride in solutions in conc. HFAq.

VANADIUM TRIFLUORIDE VF .. 3H2O. not known; formula probably molecular, from analogy of trihalides of other elements of Group V. Prepared by evaporating a solution of V_2O_2 in HFAq at 100° , and crystallising from water containing a little HFAq. A white crystalline crust, consisting of rhombohedra. Easily soluble in water; solution reacts acid. Insoluble in conc. alcohol. Effloresces in air; loses some water at 100°, and all at 130°, with absorption of O; V2Oa is formed on heating to redness in air. An aqueous solution reduces Ag salts to Ag, Hg and Cu salts to mercurous and cuprous salts; alkalis or alkali carbonates ppt. V₂O₂.xH₂O from VF₂Aq (E. Petersen, J. pr. [2] 40, 44).

Vanadofluorides. (Fluovanadites.) A number of compounds of the forms VF₂.xM¹F

and VF.xMnF. were obtained by Petersen (l.c.) by adding various fluorides to solutions of V.O.

These salts are in HFAq, and crystallising. generally greenish coloured, and crystalline; most of them are soluble in water, but not in KFAq; they lose water of crystallisation from 100° to 200°. The vanadofluorides may be regarded as salts of hypothetical acids HVF, H2VFs, and H2VFs; they may also be formulated as double salts (v. supra). The following salts are described by Petersen (l.c.):

Ammonium vanadofluorides NH,VF. 2aq,

 $(NH_1)_2 \nabla F_5$, aq, and $(NH_1)_2 \nabla F_6$.

vanadofluoride Cadmium CdVF, 7aq (Piccini a. Giorgis, G. 22 [1] 55).

Cobalt vanadofluoride CoVF, 7aq. Nickel vanadoftuoride NiVF₅.7aq. Potassium vanadoftuoride K₂VF₅.aq Sodium vanadofluoride Na, V.F.11. aq

= 2VF₃.5NaF. aq (? Na₂VF₃. aq).

Zinc vanadofiuoride ZnVF₃. 7aq (P. a. G.,

l.c.).
Vanadoxyfluorides. (Fluoxyvanadates.) These salts are generally produced by dissolving V_2O_5 in HFAq, adding metallic fluorides, and crystallising; also by dissolving V_2O_5 in solutions of metallic fluorides, and crystallising; some of them are obtained by treating solutions of V2Os in metallic fluoride solutions with reducing agents, and then crystallising. The vanadoxyfluorides have been examined by Baker (C. J. 33, 388 [1878]), Ditte (C. R. 105, 1067; 106, 270 [1888]), Piccini a. Giorgis (G. 18, 186 [1889]; G. 22 [1] 55 [1892]), and Petersen (B. 21, 3257 [1888]; more fully in J. pr. [2] 40, 193, 271 [1890]). There are still considerable discrepancies between the results obtained by the different observers. For a criticism of Baker's results v. Petersen (J. pr. [2] 40, 289-92). The vanadoxyfluorides belong to two main classes: (1) those obtained by dissolving V2Os in HFAq, adding a fluoride (or carbonate), and crystallising: many of these salts are also formed by dissolving V₂O₃ in solutions of fluorides, and crystallising; (2) those obtained by partially reducing solutions of V2O, in HFAq, then adding fluorides, and crystallising. The first class of salts may be divided into (i.) those wherein one atom of O in V2O, is substituted by 2F, and which, therefore, are represented as xMF.yVO₂F $(V_2O_5$ gives $V_2O_4F_2 = 2VO_2F)$; and (ii.) those wherein 3O in V_2O_5 is substituted by 6F, and which, therefore, are represented as xMF.yVOF, $(\nabla_2 O_5$ gives $\nabla_2 O_2 F_6 = 2 \text{VOF}_5$). Salts of the second class are regarded as compounds of MF with VOF_2 , and VOF_2 is looked on as obtained by substituting 20 in V_2O_4 by 4F (V_2O_4 gives $V_2O_2F_4 = 2VOF_2$). The salts derived from V_2O_4 . i.e. xMF_3VOF_4 and xMF_3VOF_5 , are often called fluoxyvanadates, and those derived from V₂O₄, i.e. xMF.yVOF₂, are often called fluoxy-hypovanadates; the former class of salts will here be called vanadoxyfluorides, and the latter hypovanadoxyfluorides.

Ammonium vanadoxyfluorides. Three salts are described by Baker (C. J. 33, 388 [1878]), and also by Piccini a. Giorgis (G. 22 [1] 55, [1892]; the formulæ given by the different experimenters do not agree.

Lamellar ammonium vanadoxyfluoride.
This salt was obtained by Baker by dissolving
V.O. in NH,HF,Aq, and then adding more
NH.HF. it servents in the control of the control o NH.HF.; it separates in small yellow, pearly,

hexagonal plates. P. a. G. obtained the salt by dissolving V₂O₅ in a slight excess of HFAq, adding NH₄F in the ratio 8NH₄F:2V₂O₅, and evaporating. B. gives the formula $6NH_*F.V_*O_*P.VOF_*$. 2aq; P.a.G. give the formula $3NH_*F.2VO_*F$. If both formulæ are expressed as $3NH_*F.2VO_*T$, then $x=1\frac{n}{2}$ in Baker's formula and =2 in the formula of P. a. G. A salt obtained in the formula of P. a. Q. 40, 103, 271 tained by Petersen (J. pr. [2] 40, 193, 271 [1890]), by a similar process to those used by B. and by P. a. G., was formulated by him as 7NH,F.4VO,F.HF.

Pyramidal ammonium vanadoxyfluoride. Obtained by Baker by adding NH, HF, to a solution of the lamellar salt; P. a. G. prepared the salt by dissolving V2O, in excess of HFAq, and exactly neutralising the hot solution by NH3Aq. This salt crystallises in yellow, simple pyramids The formula assigned by B. is 12NH, F.V2O5.2VC. while that given by P. a. G. is 3NH,F.VO,F; Petersen gives the same formula as P. a. G. If Baker's formula is expressed as $3NH_4F.VO_xF^y$, x must be put = $1\frac{3}{4}$, and $y=1\frac{1}{2}$.

Acicularammonium vanadoxyfluoride. Baker obtained this salt by dissolving the pyramidal salt in warm HFAq and allowing to cool; P. a. G. prepared it by a similar reaction. This salt crystallises in pale-yellow acicular prisms; when dry it has the odour of HF; it acts on glass. The formula given by P. a. G. is 3NH,F.2VOF, aq; Baker assigned to this salt the formula 3NH,HF,2VOF, but his analyses were, admittedly, not very trustworthy.

Adopting the formulæ given by P. a. G. for

the three ammonium vanadoxyfluorides, these

salts are represented as

(1) 8NH,F.2VO,F = (NH₄)₃V₂O₄F₃ (2) 8NH,F.VO₂F = (NH₄)₃VO₂F₄ (3) 3NH₄F.2VOF₃ = (NH₄)₃V₂O₂F₉.

A few of the other vanadoxyfluorides seem to belong to one or other of these types; P.a.G. and also Petersen, describe a potassium vanadoxy-fluoride $3KF.2VO_2F = K_3V_2O_4F_3$, belonging to the same form as the lamellar ammonium salt. But most of the vanadoxyfluorides described belong to forms which are not represented by the ammonium salts. The following are the principal salts :-

Ammonium salts. 9NH, F.5VOF, 3HF; 7NH, F.4VOF, HF (Petersen).

Potassium salts 2Kf.VO.F; 2KF.VOF.; 3KF.2VOF₈.HF; 4KF.VF₈.VOF₈ (P.)

Sodium salt 3NaF.VOF. VO2F. aq a. G.).

Zinc salts. ZnF₂.VO₂F. 7aq (P. a. G.); ZnF₂.ZnO.2VOF₂. 14aq (B.).

In none of these compounds does the atomic ratio of V to O exceed 1:2. According to Ditte (C. R. 105, 1067; 106, 270 [1888]), many compounds of alkali fluorides with V2O, are formed by heating the constituents together, digesting with water, and crystallising; the atomic ratio of V to O in these compounds, as given by Ditte, is 2:5. The formulæ assigned to the compounds may be generalised as xV_2O_xyMF ; M=K or Na, x=1, 8, and 4, and y=1, 2, 4, and 8. Ditte (i.e.) says that crystals of V_2O_x 4NH, F.4aq are obtained by dissolving V_2O_x in hot conc. NH, FAq, and that this compound is formed, along with V_2O_x 4NH, F.4aq when V_2O_x 4 is dissolved to add V.O. 8NH, F. 4aq, when V2Os is dissolved in cold conc. NH,FAq; but the experiments of Baker, Piccini a. Giorgis, and Petersen, all agree in assigning to the compounds formed in these reactions formulæ wherein the atomic ratio of

 ∇ to O does not exceed 1:2.

Hypovanadoxyfluorides. (Fluoxyhypovanadates.) These compounds are formed by partially reducing mixtures of V2Os and fluorides, or carbonates, dissolved in HFAq. If sufficient F is assigned to the metal present to form the normal fluoride, then the atomic ratio of V to the remaining F in these compounds is 1:2, and that of V to O is 1:1; the formulæ are written xMF.yVOF_s. All the salts described by Baker and Piccini a. Giorgis, except one, belong to the form $2M^1F.VOF_2$ or $M^1F_2.VOF_2$; these formulæ may evidently be written $M^1_*(\text{or}M^{11})VOF_4$.

Ammonium hypovanadoxyfluoride $2NH_4F.VOF_2$. $aq(=(NH_4)_2VOF_4$. aq) was prepared by Baker (C. J. 33, 395) by passing H₂S into a solution of V_2O_3 in HFAq, concentrating, filtering, and adding NH₄F. The same salt was prepared by Piccini a. Giorgis (G. 22 [1] 55) by electrolytically reducing a solution of NH VO in HFAq, after adding NH,F; also by reducing NH, VO, in HFAq by SO₂Aq, neutralising by NH₃Aq, and adding NH,F; and in each case dissolving the crystals that were formed (said by P. a. G. to be SNH₄F.VOF₂) in hot HFAq, and crystallising. Petersen (J. pr. [2] 40, 196) also obtained this salt. The salt crystallises in clear blue monoclinic forms; a:b=.9653:1 (Baker, l.c.).

Petersen (J. pr. [2] 40, 197) described two other ammonium hypovanadoxyfluorides,

7NH,F.4VOF, 5aq and 3NH,F.VOF,

The other hypovanadoxyfluorides described by P. a. G. (l.c.) are 2KF.VOF₂ and MF₂.VOF₂.2aq where M = Cd, Co, Ni and Zn. The K salt was prepared by reducing V2Os in HFAq by SO2Aq and adding KF; the other salts by dissolving MO, or MCO_s, along with V₂O_s in HFAq, and reducing electrolytically. According to Petersen (l.c. pp. 199, 200), the salts 7KF.3VOF, and 8NaF.3VOF, 2aq also exist.

P. s. G. (l.c.) remark that the tendency to

form highly fluorinated compounds increases in the family V, Nb, Ta as the atomic weight increases; thus, VO₂F is not affected by a little HFAq, while Nb₂OF forms NbOF₃, and TaOF₄ is converted into TaF₅. They also point out that the three compounds, ZnF₂,VO₂F. 7aq, ZnF₂,VOF₂. 7aq, and ZnF₂,VF₃. 7aq, all crystallise with the same number of molecules of water, all contain the same number of atoms, have the same crystalline form, and are geometrically isomorphous.

Vanadium pentafluoride VF₅. According to Petersen (J. pr. [2] 40, 271), this compound exists in a solution of V2O5 in conc. HFAq. No compound was isolated, nor is any trustworthy evidence given of the existence of VF, in solu-

tion.

Vanadium, haloid compounds of. When V is heared in excess of Cl the tetrachloride VCl, is produced; heating in Br produces the tribromide VBr. V does not combine with I. The haloid compounds of V belong to the forms ∇X_1 , ∇X_2 , and ∇X_2 ; X = Cl in each case, X = Br or F in the one case of ∇X_2 . The formula ∇X_4 is molecular; the other formulæ are probably molecular. VOl. BiCl., AsI., and PI. are the

only dihalides known of members of Group V. Attempts to prepare halides higher than VX. have failed. Solutions in water of the haloid compounds of V behave like solutions of the corresponding exides in haloid acids; VCl,Aq like a solution of V₂O₄ in HClAq, VX₂Aq like a solution of V₂O₅ in HXAq, and VCl₂Aq like a solution of V₂O₅ in HClAq. VCl, is decomposed by heat to VCl₃ and Cl, and heating VCl₂ with H produces VCl2. VF, combines with metallic fluorides to form vanadofluorides, many of which belong to the form M¹₂VF₅ or M¹¹VF₅ (v. p. 844). Several oxychlorides and oxybromides of V have been isolated: VOX₅ and VOX₂, where X = Cl or Br; also VOCl, V₂O₂Cl, and VO₂Cl; the existence of other oxybromides also is probable. No oxyfluoride has been isolated with certainty. But many compounds exist which may be regarded as containing the oxyfluorides VO₂F, VOF₃, and VOF₂ respectively; the compounds VO₂F.xMF and VOF₃.xMF are obtained by dissolving V2O, and fluorides in HFAq and crystallising; they are described as vanadoxyfluorides (p. 844); the compounds VOF .xMF are obtained by partially reducing solutions of V2O5 and fluorides in HFAq, they are described as hypovanadoxyfluorides (p. 845).

Vanadium, iodides of. No iodide of V has been isolated. No reaction occurs when vapour of I is passed over VN heated to redness, nor do I and $\nabla_2 O_2$ react at any temperature (Roscoe, C. J. [2] 9, 28). I does not react with $\nabla_2 O_4$ when digested therewith for a long time in presence of water or alcohol (Guyard, Bl. [2] 25,

Vanadium, nitrides of. V and N combine

directly. Two nitrides are known.

VANADIUM MONONITRIDE VN. Mol. w. not known. Obtained by heating V in a stream of pure N; also by saturating VOCl, with dry NH, and heating to full whiteness in a stream of NH₂. Better prepared by heating NH₄VO₂ in the air, and then heating the residue to a full white heat in a stream of dry NH₃ (Roscoe, C. J. [2] 8, 844). VN may also be prepared by heating V₂O₂ to whiteness in NH₃ (R., *l.c.*). VN is a greyish-brown, metal-like powder; heated in air it is oxidised to V2O, and then to V2O,;

heated with soda-lime it gives off NH.

VANADIUM DINITRIDE VN. A black powder;
exposed to air gives off NH, and is oxidised.
Prepared by saturating VOCI, with dry NH, heating the solid so obtained in a glass tube as long as NH₄Cl sublimes, washing the residue with water containing a little NH₃, and drying in vacuo over H.SO. (Roscoe, C. J. [2] 6, 849); examined by Uhrlaub, who did not succeed in

determining its composition (P. 108, 184; cf. Schafarik, W. A. B. 33, 5).

Vanadium, oxides and hydrated oxides of.
By gently heating V in air the oxide V.O is possibly formed, and, on continued heating, exidation proceeds to V_2O_2 , V_2O_3 , V_2O_4 , and finally V_2O_4 . The exides V_2O_4 , V_2O_5 , and V_2O_2 are also formed by reducing V_2O_4 by heating with Hor K. Several oxides intermediate between V_2O_4 and V_2O_5 seem also to exist. The oxides V_2O_3 , V_2O_4 , V_2O_4 , and V_2O_5 are feebly basic; V_2O_4 and V_2O_5 are also acidic. The oxides of V_2O_4 oxides of V_2O_5 are also acidic. the oxides of N; corresponding oxides of Bi

are known. The names generally given to the oxides of V do not, unfortunately, tally with the names given to the corresponding oxides of N; thus, V₂O₂ is generally called hypovanadous oxide, while NO is called nitric oxide; V₂O₃ is called vanadic oxide, while the name nitrogen peroxide is often quite wrongly applied to NO₂.

The mol. w. of none of the oxides of V is known.
? VANADIUM MONOXIDE (? V₂O). 'When the powdered metal is . . . slowly Leated in a current of air it glows brightly, with absorption of oxygen, forming in the first place a brown oxide (V2O or V2O2); and on further heating this oxide again glows, and passes through the black trioxide and blue tetroxide to pentoxide ' (Roscoe, C. J. [2] 8, 357). This statement seems to contain all that is known of an oxide lower than V₂O₂.

VANADIUM DIOXIDE V2O ... (Hypovanadous oxide.) This substance was supposed by Berzelius (P. 22, 1) to be V; Roscoe (T. 1868. 1) proved it to be an oxide with the atomic ratio V:O=1:1. The mol. w. is not known; the formula is generally written $\nabla_2 O_2$ rather than VO, as the double formula is, on the whole, more in keeping with the reactions of the com-

pound.

Formation.—1. By heating one of the higher oxides with K (Berzelius, l.c.).—2. By passing vapour of VOCl3, mixed with much dry H, over strongly heating V_2O_3 , and V_2O_5 , mixed in the ratio V_2O_3 : V_2O_5 , out of contact with air (Petersen, J. pr. [2] 40, 193).

Preparation.—A mixture of much dry H and vapour of VOCl₃ (q. v. p. 854) is passed through a hard glass tube containing powdered charcoal and kept red hot. Some solid oxychlorides are deposited on the cooler parts of the tube, while the V2O2 remains mixed with the C. The contents of the tube are then st.ongly heated in a current of dry H to remove traces of Cl compounds; and the grey metal-like V₂O₂ is separated from the C (Roscoe, C. J. [2] 6, 334). When Zn, Cd, or Na-amalgam is added to a solution of V_2O_5 in H_2SO_4Aq (prepared by dissolving the oxide in hot cone. acid and diluting with 50 pts. water), the liquid becomes blue. green, and finally lavender. The lavender solu-tion absorbs O very rapidly; exposure to the air for a few seconds causes the colour to change to deep chocolate-brown. By determining the quantity of standardised KMnO₄Aq decolourised by the lavender solution, Boscoe (C. J. [2] 6, 334) proved that this solution contained V_2O_2 , probably combined with SO. After passing air into the lavender solution until it becomes permanently blue, the solution contains V_2O_4 ; if the free acid in the lavender solution is neutralised by zinc, and the liquid is then exposed to the air until permanently brown, and a little acid is then added, a green solution is formed which contains V₂O₃ (R., l.c.).

Properties and Reactions.—A light-grey, lustrous powder; or a lustrous, metal-like, crystalline crust, S. 7. 8-64; very brittle; fuses at a very high temperature. Heated to redness in sir, $\nabla_2 O_3$ burns to $\nabla_2 O_4$; heated in O, burns to $\nabla_2 O_4$. Insoluble in water; dissolves in dilute acids, giving a lavender solution, which bleaches strongly; from these solutions KOHAq or

NH, Aq ppts. a brown hydrated oxide, which | rapidly oxidises (Brierley, C. J. 49, 824). According to Petersen (J. pr. [2] 40, 194), V₂O₂ dissolves in HFAq, and on evaporation, solution of the residue in water, and evaporation over H₂SO₄, blue, microscopic crystals are obtained, which are, perhaps, VOF2. xaq; no analyses were made. Addition of alkali fluorides to V2O2 in HFAq produces various compounds of the form $x \nabla \tilde{\text{OF}}_{2}.y \text{MF}$ (v. Hypovanadoxyfluorides, p.

HYDRATE OF VANADIUM DIOXIDE. By adding NH₃Aq to a lavender-coloured solution of V₂O₃ (? in H₂SO₄Aq), Brierley (C. J. 49, 824) obtained a deep-purple gelatinous pp. which he said was hydrated V2O2; the pp. rapidly oxidised in the

Vanadium trioxide $\nabla_2 O_3$. (Vanadous oxide. Vanadium sesquioxide.) Mol. w. not known. This compound was represented by Berzelius as VO $(\nabla = 68.5, O = 8)$.

Formation.—1. By heating $\nabla_2 O_5$ to redness in H (Schäfarik, A. 109, 85). The presence of traces of $P_2 O_5$ prevents reduction (Roscoe, C. J. [2] 6, 331).—2. By heating $\nabla_2 O_5$ in a carbon crucible (Berzelius, P. 22, 1).

Preparation.—Pure V2Os is heated in perfectly pure and dry H until the substance ceases to lose weight; the tube is allowed to cool completely in H, and then a stream of dry air is passed through it (Roscoe, C. J. [2] 6, 331). When a solution of V_2O_3 in conc. H_2SO_4 is much diluted, and then treated with Mg until the liquid is green, this solution contains V₂O₃, combined with SO₃ (v. Vanadous salts, infra). A similar solution is obtained from the layender solution of V2O2 in H2SO4Aq (prepared from V_2O_5 in H_2SO_4 , v. supra) by neutralising free acid by zinc, exposing to the air, and adding a little

acid (Roscoe, C. J. [2] 6, 337).

Properties and Reactions.—V₂O₃ is a black powder, resembling graphite; or, formed by reducing VOCl, by H, it is a black, lustrous, crystalline crust. S.G. 4.72 at 16°. Does not fuse in the blowpipe flame. When warm V_2O_3 is exposed to the air it glows and oxidises to V_2O_5 ; and even at the ordinary temperature it takes up O, being changed after some months to V₂O₄ (R., Lc.). Heated in Cl gives V_2O_3 and $VOCl_3$ (R., Lc.). Heated in NO_2 to $300^\circ-400^\circ$ V_2O_3 is oxidised to V_2O_3 (Sabatier a. Senderens, C. R. 115, 236). Heating to redness in H₂S produces V₂S₂ (Kay, C. J. 37, 728). V₂O₃ dissolves in hot conc. H₂SO₄, forming V₂O₃.xSO₂ (v. Vanadous salts. infra). V₂O₃ dissolves in HFAq, and VF₃.3H₂O has been isolated from this solution (v. Vanadium TRIFLUORIDE, p. 844).

HYDRATE OF VANADIUM TRIOXIDE. By adding KOHAq, or NH, Aq, to an aqueous solution of $V_{2}O_{3}.48O_{2}.9H_{2}O$ (v. infra) Brierley (C. J. 49, 824), obtained a dirty green, gelatinous pp., which oxidised rapidly in air. According to B., this

pp. was hydrated V_2O_3 . VANADOUS SALTS V_2O_3 .xX (X = acidic oxide) or V_2R_2 ^{II} (R^{II} = acidic radicle). Only one salt derived from an oxyacid has been certainly isolated; it has the composition V2O, 4SO, xH2O, or the formula may be written V2H2(SO4)4.yH2O.

Vanadous tetrasulphate, or acid vanadous sulphate, V.O. 486. xH.O or V.H. (80.). yH.O. A hydrate with x=3 or y=2 was obtained by Gerland (B. 10, 2111 [1877]) by heating a solution of V₂O₄ in a large excess of conc. H₂SO, for a long time at 120°, and treating the blue crystalline crust thus formed (consisting chiefly of V2O4.3SO2. xaq, v. HYPOVANADIO SALTS, p. 848) with boiling water, when V₂O₂.4SO₂.3H₂O remained. G. supposed that reduction of V2O4

was effected by dust in the air.

Brierley (C. J. 49, 822 [1886]) prepared $V_2O_s.4SO_s.9H_sO(=V_2H_s(SO_4)_s.8H_sO)$ by electrolysing a solution of V_2O_t in H_sSO_t . The solution was prepared by reducing V_2O_s in conc. H_sSO_t , after dilution, by SO_s . The deep-blue solution was placed in a Pt dish which formed the negative electrode, and a porous pot containing dilute H.SO.Aq was arranged so that it just touched the surface of the liquid in the Pt dish; a plate of Pt immersed in the H₂SO₄Aq formed the positive electrode. The Pt dish was warmed during the process, so that electrolysis and evaporation proceeded together. Electrolysis was continued until the liquid was deep green, and a drop placed in a little water showed no trace of blue colour. Conc. H₂SO₄, equal to twice the bulk of the green liquid, was added; after 24 hours the green, sandy pp. was collected, washed with conc. alcohol, and dried over H_2SO_4 in coal-gas. The salt dissolved in water, and alkali ppd. from this solution a dirtygreen gelatinous substance, said by B. to be V₂O₂.xH₂O. This pp. dissolved in an aqueous solution of V2O3.4SO3.9H2O, forming a chocolatebrown solution, giving a lustrous green solid on evaporation, which dissolved in water to form a brown solution. B. concluded that the brown solution contained a compound of V₂O₂ and SO₂

more basic than $\nabla_2 O_3 \cdot 4SO_3$. Gerland (B. 11, 106 [1878]) mentions the normal vanadous sulphate $\nabla_2 (SO_4)_3 (= \nabla_2 O_3 \cdot SSO_3)$

as if he had prepared it; no details are given.

VANADIUM TETROXIDE V₂O₄. (Hypovanadic oxide.) Mol. w. not known. Represented by

Berzelius as VO₂(V = 68.5, O = 8).

Formation.—1. By allowing V₂O₃ to absorb O from the air (cf. Hypovanado-vanado oxides, p. 851).—2. By strongly heating a mixture of 5 pts. V₂O₃ and 6 pts. V₂O₅, in absence of air.— 3. By the electrolysis of molten V_2O_5 (Buff a. Wöhler, A. 110, 277).—4. By heating V_2O_4 Cl₂. 5aq

in a stream of CO₂ (Crow, C. J. [2] 15, 453).

Preparation.—A solution of V₂O₄ in H₂SO₄, obtained by reducing V₂O₅ in conc. H₂SO₄ by SO2, is ppd. by Na2CO2Aq, and the pp. is washed and dried by heating in vacuo (Berzelius, P. 22, 1).-2. V_2O_3 is exposed to the air for some months, until the colour has changed to blue (Roscoe, C. J. [2] 6, 838).—3. V₂O₅ is dissolved in hot conc. HClAq; when Cl ceases to be given off the liquid is saturated with H2S, filtered from S, evaporated to dryness at 100° and the V2O4Cl2/xaq thus formed is heated to redness in a stream of dry CO₂ (Crow, l.c. pp. 457, 458).

Properties and Reactions.—A blue, lustrous,

crystalline powder; prepared by heating V2O4Cl2 in CO₂ it forms a dark-green, amorphous powder (Crow, l.c.). Infusible at a high temperature. Insoluble in water. When kept in a loosely-stoppered bottle V₂O₄ gradually absorbs O, forming V₂O₅ (Crow, l.c.). Dissolves easing acids, forming bright-blue liquids; the same solutions are obtained by the action of moderate

reducers, such as SO₂Aq, H₂SAq, or H₂C₂O₄Aq, on solutions of V_2O_5 in acids, also by passing air into V_2O_2 in H₂SO₄Aq until the liquid becomes permanently blue (Roscoe, l.c.; v. infra, Hypo-VANADIC SALTS). Petersen (J. pr. [2] 40, 194) supposed that the blue microscopic crystals he obtained by dissolving $\nabla_2 O_4$ in excess of HFAq and evaporating were VOF_2 , xaq, but he did not obtain enough to make an analysis. V_2O_4 dissolves readily in alkali solutions, forming hypovanadates, $M_2V_4O_9$ (q. v. under Vanadium oxy-ACIDS, AND SALTS AND DERIVATIVES THEREOF, D.

HYDRATES OF HYPOVANADIC OXIDE. By adding Na₂CO₂Aq, drop by drop, to a solution of V₂O in HClAq or H.SO.Aq, till the supernatant liquid was colourless, filtering rapidly in CO2, and drying on a porous tile over H₂SO, in vacuo, Crow (l.c.) obtained the black, amorphous heptahydrate V₂O₄. 7aq; and on heating this in CO₂ for some hours at 100° he obtained the trihydrate

V₂O₄. 3aq

HYPOVANADIC (or vanadyl) SALTS V_2O_4 .xX (X = acidic oxide). Only a few compounds of V2O4 with acidic oxides have been isolated. The sulphates have been studied more than any other salts; they are most simply represented as $V_2O_4.xSO_3.yH_2O$. The compound $V_2O_4.2SO_3$ may be called the normal sulphate; the formula may be expressed as V_2O_2 (SO₄)₂ or with acidic oxides are written V2O4.X the compounds are generally called hypovanadic salts; when the formulæ are written $\nabla_2 O_{xx} R$ (R = acidic radicle) the name divanadyl salts is usually given to the compounds; and when these compounds are regarded as VO.xR, it is customary to call them vanadyl salts. Hypovanadic sulphates are obtained (1) by dissolv- | vanadic sulphates are as follows:—

H₂SO₄ and heating to the boiling-point of the liquid for a long time. The salt separated as a green-blue powder; insoluble in water, cold or hot; insoluble HClAq and H2SO4. (G. does not say how the salt was obtained pure for analysis; no analyses are given of the salt prepared exactly as described above.) Various hydrates are described by G. obtained by different processes. Crow (C. J. [2] 15, 456 [1876]), obtained the heptahydrate by repeatedly treating hexa-hydrated trisulphate (v. infra) with absolute

Hypovanadic trisulphate, or acid divanadyl

sulphate, $V_2O_4.3SO_3.xH_2O$ or $(V_2O_2)H_2(SO_4)_3.yH_2O$ [y=x-1]. (A sulphate $(VO.SO_4)_2.H_2SO_4.yH_2O$.) (Acid vanadyl Hydrates with x=4 and 3 (or y=3 and 2) were prepared by Gerland (B. 10, 2, 109); and a hydrate with x = 6 (or y = 5) by Crow (C. J. [2] 15.

By dissolving $\nabla_2 O_4$ in a large excess of conc. H₂SO₄, heating for a long time at c. 120°, and drying the small, blue, transparent or stals that formed on a porous tile over H2SO4, Gerland obtained a substance which gave analytical results approaching those required by the formula V₂O₄.3SO₂.4H₂O. The salt was not obtained quite free from admixed V₂O₂.4SO₂.xH₂O. This tetrahydrate was partially decomposed by cold water, giving lustrous scales which, after washing with alcohol and ether and drying over H₂SO₄, had the composition of a trihydrate, V2O4.3SO8.3H2O.

Crow obtained a hexahydrate $V_2O_4.3SO_9.6H_2O$, by reducing by SO_2 a solution of V_2O_5 in conc. H_2SO_4 , heating at 100° for some time, drying the light-blue crystals that separated on a tile, removing acid by washing with ether, and drying by pressure between paper. The crystals deliquesced in air to a blue syrup; they were insoluble in ther, and scarcely

soluble in absolute alcohol.

The formulæ of the various hydrated hypo-

Hypovanadic sulphates;	or	Divanaayi suiphates;	or	vanadyi suipnates.
$\nabla_2 O_4.2SO_4$		$\nabla_2 O_2 \cdot (SO_4)_2$		VO.SO, (Gerland).
V,O,.2SO,.3H,O		V ₂ O ₂ .(SO ₄) ₂ .3H ₂ O		2(VO.SO).3H2O (G.)
$\nabla_{2}^{2}O_{4}^{2}.2SO_{3}^{2}.4H_{2}^{2}O$		$\nabla_2^2 O_2^2 (SO_4^2)_2 \cdot 4H_2^2 O$		VO.SO.2H ₂ O (Crow, confirming Berzelius).
V ₂ O ₄ .2SO ₄ .7H ₂ O		$\nabla_2 O_2 \cdot (SO_4)_2 \cdot 7H_2O$		2(VO.SO ₄).7H ₂ O (G., also C.)
V.O.2SO.10H,O		V₂O₂.(SO),.10H₂O		VO.SO, 5H,O (G.)
V.O.280.13H.0		V ₂ O ₂ (SO ₄) ₂ 18H ₂ O		2(VO.SO ₄).18H, O (G.)
V ₂ O ₄ .3SO ₄ .3H ₂ O		$(\nabla_2 O_2) H_2 (SO_4)_3.2 H_2 O$		2(VO.SO,).H ₂ SO,2H ₂ O (G.)
$\nabla_{2}O_{4}.38O_{4}.4H_{2}O$		(V,O,)H,(SO,),3H,O		2(VO.SO ₄).H ₂ SO ₄ .3H ₂ O (G.)
$\nabla_2 O_4.38O_8.6H_2O$		$(\nabla_2 O_2) H_2 (SO_4)_3.5 H_2 O$		2(VO.SO ₄).H ₂ SO ₄ .5H ₂ O (C.)

ing V_2O_4 in hot cone. H_2SO_4 and boiling; (2) by dissolving V_2O_4 in conc. H_2SO_4 , reducing by SO_2 or $H_2C_2O_4$, and evaporating; (3) by dissolving V_2O_4 in conc. H_2SO_4 and boiling in the air for some time. In reaction (3) V₂O₃.xSO₄ seems to be formed and then partially reduced, by dust or products of combustion of coal-gas, to $V_2O_4xSO_2$; the products are mixtures of $V_2O_4xSO_3$ and $V_2O_4xSO_3$ (Gerland, B. 11, 98).

Hypovanadic disulphate, or normal divanadyl sulphate, V.O.,280, or V.O.,(80.).
"Vanadyl sulphate VO.SO.") Prepared by Gerland (B. 10, 2109 [1877]; supplementing B. 9, 869) by dissolving V.O. in a large excess of conc.

Bevan (C. N. 38, 294) obtained divanadyl dithionate by adding BaS, O, Aq to V,O, (SO,),Aq, filtering, and evaporating in vacuo: the salt was not obtained pure. By dissolving V₂O₄ in HNO₂Aq Berzelius (P. 22, 1) obtained a blue liquid which probably contained hypovanadic nitrate; but no solid could be isolated, as on evaporation to dryness decomposition occurred, with formation of V_2O_3 . V_2O_4 also dissolves in H_2PO_4Aq ; on evaporation at 50° blue crystals are obtained, which deliquesce to a blue syrup; these crystals are probably hypovanadic phosphate, but no analyses and given (B. Lc.). Berzelius obtained what were probably compounds of ∇_2O_4 with As_2O_5 , B_2O_5 , CrO_5 , MoO_5 , and a few other acidic oxides (v. Lehrbuch [5th ed.] 3, 1053).

Vanadium pentoxide V_2O_3 . (Vanadic oxide. Vanadic anhydride.) Mol. w. not known. Represented by Berzelius as VO_3 (V=68.5, O=8).

Formation.—1. By strongly heating any of the lower oxides in air or O.—2. By heating NH,.VO₂ in air to dull redness.—3. By decomposing VOCl₂ by water.—4. By heating V₂O₂,4SO₂, 9aq to bright redness (Brierley, C. J. 49, 824).

Preparation.—Pure VOCl₂ is decomposed by water, and the ppd V₂O₂ is washed till the washings are free from acid (HClAq is produced) and heated until dry (Schafarik, J. pr. 76, 142). For the preparation of V₂O₃ from vanadinite and other sources of V compounds, v. Vanadium, Pre-

paration of (p. 841).

Properties.—A nale-yellow powder, with a reddish tinge. When heated to full redness out of contact with any reducing agents, V2O5 melts, and solidifies on cooling to a mass of yellow-red, crystalline needles, appearing ruby-red by transmitted light; should the V₂O₅ used contain V₂O₄, which is often the case when the $\nabla_2 O_5$ is prepared by heating NH4. VO3, the solid obtained by fusion and cooling appears violet to nearly black, and does not seem to be crystalline (Berzelius, P. 22, 1). As molten $\nabla_2 O_5$ cools nearly to its solidifying-point much heat is given out, and the mass glows until solidification is complete (B., l.c.). V.O. is said not to be changed at a white heat, provided no trace of a reducing agent is present (B., l.c.); but according to Read (C. J. 65, 313 [1894]) molten V_2O_5 gives off O, producing V_2O_8 , which on cooling in air takes up O, forming steel-blue crystals of V2O4. The crystals of V2O8 are rhombic (v. Nordenskjold, P. 112, 160). S.G. (crystalline) 3.5 at 20° (Schafarik, J. pr. 76, 142). V₂O₅ is odourless and tasteless; it reddens moist litmus paper; 1,000 pts. boiling water dissolve scarcely 1 pt. V₂O₅ (B., l.c.). With molten alkalis or alkali carbonates $\nabla_2 O_5$ produces vanadates (q.v. V₂O₅ dissolves in conc. H₂SO₄, HClAq, . 851).

HNO, Aq, or HFAq (cf. Vanadic sales, p. 850).

According to Ditte (C. R. 101, 698), V₂O₈ exists in three forms: (1) red, obtained by heating NH₄.VO₂ in a closed crucible, treating with HNO₃ and repeatedly heating to redness; (2) yellow, obtained by heating NH₄.VO₂ to 440° in a stream of air; and (3) crystalline, obtained by treating the yellow form with HNO₃, and then fusing. D. says that the red form is somewhat soluble in water, 1,000 c.c. dissolving 8 g. at 20°; that the yellow form is very slightly soluble, and the crystalline form is insoluble, in water. He also says that the red form absorbs water from the air, producing hydrates with 2H₂O and 5H₂O; and that addition of NHCl, KCl (and other salts) to an aqueous solution of the red variety causes ppn. of a hydrate of the yellow form of V₂O₃, which hydrate, when dry, is a dark-red powder having the composition V₂O₂.2H₂O (=H₄V₂O₇) and is almost insoluble in water. (No analyses

are given by D.)

Reactions.—1. When heated and kept molter, V_2O_2 is formed, and this absorbs O as it cools in air and forms V_2O_4 (Read, C. J. 65, 313 [1894]). 2. Heated to refiness in hydrogen till it ceases to lose weight, V_2O_2 is reduced to V_3O_3 (Schafarik, Vol. 17.

A. 109, 85); traces of P₂O₅ prevent the reducwith potassium produces V_1O_2 (Berzelius, P. 22, 1).—4. V_2O_3 is produced by heating V_2O_4 in a carbon crucible (B., l.c.). For action of in a carbon crucible (B., l.c.). For action of Mg, Zn, &c. on solutions of V₂O₄, v. Vanadic sales, (p. 850).—5. When mixed with carbon And heated in chlorine, V₂O₅ yields VOCl₃ (Roscoe, C. J. [2] 6, 342).—6. Electrolysis of molten V₂O₅ produces V₂O₄ (Buff a. Wöhler, A. 110, 277).—7. According to Ditte (C. R. 103, 55), fusion with potassium cyanide produces V₂O₄ (Buff a. Wöhler, A. 100, 277).—7. According to Ditte (C. R. 103, 55), fusion with potassium cyanide produces to the control of the con V_2O_3 (N, K_2O , and CO_2 being also formed). Carnot (C. R. 104, 1803, 1850; 105, 119) says that a boiling solution of potassium cyanide produces KVO₃ (CO₂ and N being given off); and that by dropping V2O5 into excess of boiling KCNAq there is formed K₂V₄O₉ (with evolution of N and CO₂). No analyses are given either by D. or C.—8. Ditte (C. R. 103, 55) asserts that the products of the interaction of V₂O₅ and potassium iodide or bromide are K₂V₂O₅, KVO₃, and I or Br. The reactions are said to occur between $\nabla_2 O_n$ and molten KI or KBr; and KIAq is said to react in the same way as the molten salt.—9. When V2O3 is heated with molten boron trioxide, a pale-yellow glass is formed, which is said by Guyard (Bl. [2] 40, 354) to be a vanadic borate; no analyses, composition, or methods of purification are given.—10. V_2O_3 dissolves in hot conc. sulphuric acid, forming $V_2O_3.xSO_3$, generally along with more or less $V_2O_4.xSO_3$ (v. Vana-DIG SALTS, p. 850).—11. V₂O₅ dissolves in conc. hydrochloric acid; Cl is given off on heating; after the passage of H2S followed by evaporation at 100° the oxychloride VO2Cl. xaq is obtained (Crow, C. J. 30, 457; cf. Vanadium dioxymono-chloride, p. 854).—12. By dissolving V₂O₅ in conc. hydrochloric acid in presence of platinic chloride, Brauner (M. 3, 58) obtained large tablets having the composition 2(VOCl₂.PtCl₄). 21aq.-13. An oxyiodide 2VOI₂. 9aq is obtained, according to Ditte (C. R. 102, 1310), by adding excess of hydriodic acid to a warm aqueous solution of red V_2O_5 , removing free I by shaking with silver powder, filtering, and evaporating in vacuo. (No analyses are given.)—14. V₂O₃ dissolves in hydrofluoric acid, forming a pale-yellow solution, the colour of which becomes deeper on dilution (Piccini a. Giorgis, G. 22 [1] 55). Petersen (J. pr. [2] 40, 271) supposed that a solution in conc. HFAq contained VF.,; but he did not separate this compound, nor does he give any direct evidence in favour of its existtence beyond the preparation from the solution of V₂O₅ in HFAq of a compound which may be represented as 4KF.VF₅.VOF₃, and other compounds of the forms xMF.yVOF₃ and $xMF.yVO_2F$. By adding metallic fluorides to solutions of V_2O_5 in HFAq, and crystallising, vanadoxyfluorides are formed; these compounds are of the forms xMF.yVOF, and xMF.yVO.F (v. VanadoxyrLuorides, p. 844). — 15. Vanadoxyr fluorides are also formed by dissolving V2Os in alkali fluoride solutions, and crystallising (cf. infra, Combinations, No. 5) .- 16. If a solution of $\nabla_2 O_b$ in hydrofluoric acid is partially reduced, and then evaporated with metallic fluorides, hypovanadcryfluorides (q. v. p. 845) are formed; these salts are of the form xMF.yVOF2.-17. 1.1 boiling V₂O₃ with hydrogen peroxide solution a

black liquid was obtained that deposited a blackbrown crust of indefinite composition (Cammerer,

Chem. Zeit. 19, 957).—18. By fusion with alkali or alkali carbonates vanadates are formed (v. Vanadates, p. 851).

Combinations.—1. With water, but not directly, to form V₂O₂,H₂O and V₂O₃,2H₂O (v. Metavanadic acid and Pyrovanadic acid, under VANADIUM OXYACIDS, AND SALTS AND DERIVATIVES THEREOF, p. 851).—2. With various acidic oxides: (1) With SO₂ to form V₂O₃.xSO₃, formed by dissolving V₂O₅ in hot cone. H₂SO₄; (2) with As₂O₅ to form V₂O₅.xAs₂O₅; (3) with P₂O₅ to form V₂O₅.xP₂O₅; (4) with I₂O₅ to form V₂O₅.xMoO₅; (6) with WO₅ to form V₂O₅.xWO₆; (7) with P₂O₅ and SiO₂ to form V₂O₅.xP₂O₅.ySiO₂. Most of these compounds combine with the stronger VANADIUM OXYACIDS, AND SALTS AND DERIVATIVES of these compounds combine with the stronger bases M_2O and MO to form salt-like compounds. The compounds of V_2O_8 with SO_8 are described BE vanadic sulphates under VANADIC BALTE (infra); the compounds with other acidic oxides are described respectively as arseno-vanadic acids and salts, iodo-vanadic acids, molybdo-vanadic acids and salts, phospho-vanadic acids and salts, phospho-silico-vanadic acid, and tungsto-vanadic acids and salts, under VANADIUM OXYACIDS, AND SALTS AND DERIVATIVES THEREOF (v. pp. 853-4).-8. With vanadium tetroxide to form oxides intermediate between V₂O₄ and V₂O₅ (v. Hyro-Vanado-Vanadic oxides, p. 851).—4. With vana-dium trioxide to form V₂O₄; produced by heat-ing equivalent weights of the two oxides out of contact with air (Berzelius, P. 22, 1; Petersen, J. pr. [2] 40, 198).—5. With alkali fluorides. According to Ditte (C. R. 105, 1067; 106, 270), several compounds of the form $xV_2O_4.yMF$ are formed by fusing V2O, with alkali fluorides, extracting with water, and crystallising; D. gives formulæ where x=1, 3 and 4, y=1, 2, 4 and 8, and M=K and Na; but he does not give analyses of the salts (v. Vanadoxyfluorides, p. 844).

HYDRATES OF VANADIC OXIDE. The hydrates $\nabla_2 O_3 \cdot H_2 O = H V O_3$ and $\nabla_2 O_3 \cdot 2 H_2 O = H_4 \nabla_2 O_7$ have been isolated; these compounds are acids (v. Vanadic acids, p. 851). Ditte (C. R. 101, 698) asserts, without giving analyses, that a pentahydrate V₂O₃.5H₂O is formed by absorption of water from the air by the red form of V2Os (v.

VANADIUM PENTOXIDE, Properties, p. 849).

VANADIC SALTS V₂O₂...X (X = acidic oxide). A few compounds of this form have been isolated. The sulphates have been better studied than the other salts. Two sulphates, $V_2O_4.2SO_2.xH_2O$ and $V_2O_4.8SO_2.xH_2O$, have been isolated. If x is taken to be 2 in the first salt and 3 in the second, then the formulæ may be

written as (VO₂)₂SO₄.H₂SO₄.H₂O and (VO₂)₂SO₄.2H₂SO₄.H₂O respectively:

 $[(VO_4)_2SO_4.H_2SO_4.H_2O = V_2O_4.2SO_4.2H_2O;$ $(VO_4)_2SO_4.2H_2SO_4.H_2O = V_2O_4.3SO_2.3H_2O].$ If the salts are anhydrous, as seems most probable from the analyses, then the formulæ may be written $(VO_2)_2SO_4.SO_3$ and $(VO_2)_2SO_4.2SO_3$ respectively. Neither of the salts can be regarded as the normal salt. The sulphates are obtained by dissolving V.O. in hot conc. H2SO.

98), and Münzig (C. C. 1889 (ii.) 908). The salt seems to be without water of crystallisation; F. (l.c.) says it crystallises with H₂O. V₂O₅ is dissolved in hot conc. H2SO4, and the solution is boiled for a considerable time, when small crystals separate, some of which are brown and some ruby-red, lustrous, and transparent. The crystals are heated in a bath of molten lead until SO, ceases to be given off (G., l.c.). According to Münzig (l.c.), the orange-coloured crystalline powder that separates when V_2O_3 is added to boiling cone. H₂SO₄ until the acid is saturated is V_2O_3 .2SO₃. By keeping a solution of V_2O_4 in a large excess of conc. H₂SO₄ at 130°-150° for a long time, Gerland (l.c.) obtained a hard, opaque, sealing-wax red, crystalline crust composed of V_2O_s and SO_s in approximately the ratio V_2O_s :2SO_s. V_2O_s :2SO_s [= $(VO_2)_s$ SO_s.SO_s] is described by

G. (l.c.) as a clear red, crysta'line, deliquescent solid; addition of a drop or two of water ppts.

 $\nabla_2 O_3 x H_2 O_4$.

Vanadic trisulphate $\nabla_2 O_3 \cdot 3SO_8 (? x H_2 O)_4$. This salt seems to have been obtained by Berzelius (l.c.). Ditte (C. R. 102, 757) gave x the value 3; Gerland's analyses (l.c.) were inconclusive. the percentage of water found varying from 8 to 7.3. G. (l.c.) prepared this salt by long-continued boiling of a solution of V₂O₅ in a large excess of conc. H₂SO₄; small crystals separated, some of them brown and some ruby-red and transparent (probably regular octahedra). When the boiling was not continued for so long a time, G. noticed that fine, golden-yellow, lustrous needles separated. The composition of the crystals in each case was very nearly that required by the formula V2O3.3SO3. More or less V2O4.2SO4 was always formed; G. found that addition of a little HClO, prevented the formation of much V₂O₄.2SO₂.

V₂O₅.3SO₅ is very deliquescent; soluble in cold water or alcohol; a slight rise of temperature suffices to ppt. V2O3.xH2O. By adding to a solution of this salt in cold water the proper quantity of K₂SO₄, G. (l.c.) obtained small, amber-yellow crystals, which, after a few washings with cold water, pressure between paper, and drying over H2SO4, had the composition of the double salt V2O3.2SO3.K2O. 6aq. The tion of the account sair v₂O₄.25O₃.n.₂O₄.caq. and same crystals were obtained by using the disulphate V₂O₃.2SO₂. Replacement of K₂SO₄ by (NH₄)₂SO₄ produced the double salt V₂O₃.2SO₂.(NH₄)₂O₄Aq. No double salts were obtained by using Na₂SO₄ or LigSO₄, but the contained by using Na₂SO₄ or LigSO₄, but the contained to CSO₄ acceptablished unchanged. trisulphate (V.O. 8SO.) crystallised unchanged.

Gerland (B. 11, 102) found that when a solution of $\nabla_2 O_1$ in hot. conc. $H_2 SO_4$ was largely diluted and dialysed, $H_2 SO_4 \Delta q$ passed rapidly through the dialyser, along with small quantities of V compounds, until the liquid in the dialyser contained V2Os and SOs in the ratio V2O3.3SO3, after which SO3Aq passed very slowly through, and the composition of the substance in the dialyser gradually approached $V_2O_3.2SO_3$. By dialysing a diluted solution of V₂O₃ in cold conc. H2SO4, the conditions being the same as in the experimen, with the hot soid, C. found that SO, Aq passed through the dialyser regularly, and that, after a time, the ratio of V,O, to SO, in the dialyser was 1:1.82. G. supposed these Vanadic distiphate $V_2O_1.2FO_2(? xH_2O)$. and that, after a time, the ratio of V_2O_3 to SO_3 Prepared by Berzelius (P. 22, 39); examined also by Fritsche (J. pr. 58, 98), Gerland (B. 11, 20), results to mean that a solution of V_2O_3 in cold

H₂SO₄ contains loose, easily decomposed compounds of V_2O_5 and H_2SO_4 , but that a solution of V_2O_5 in hot H_2SO_4 contains V_2O_5 .3SO,, which gradually loses SO_3 until $V_2O_5.2SO_5$ is formed.

Solutions of V_2O_5 in conc. H_2SO_4 may be reduced to sulphates of lower oxides (Roscoe, C. J.

[2] 6, 334; Rammelsberg, B. B. 1880. 787):—Zn, Cd, or Na-amalgam causes reduction to lavender solutions of V₂O₂.xSO₃; Mg effects reduction to green solutions of V₂O₃,xSO₃; and H₂S, SO₃, or exalic acid reduces only to blue solutions of V2O4.xSO2.

By fusing $\nabla_2 O_5$ with $B_2 O_4$ (Guyard, Bl. [2] 25, 354) obtained a green glass, which he took to be a vanadic borate; no analyses or descrip-

tions are given.

The compounds of V₂O₅ with As₂O₅, I₂O₅, P₂O₅, MoO₂, and WO₂ respectively—or some of these compounds—may perhaps be regarded as vanadic arsenates, iodates, &c.; v. Arseno-vanadio acids and salts, Iodo-vanad acids, &c., p. 853.

HYPOVANADO-VANADIC OXIDES. Several oxides seem to exist intermediate between V2O4 and V_2O_3 . Hoscoe noticed that V_2O_3 gradually absorbed O and H_2O from the air, forming a greenish solid; Brierley (C. J. 49, 30) found that a rule grass-green substance was formed after some months, having the composition $V_2O_4.2V_2O_5.8aq = V_3O_7.4aq$. By dissolving this solid in dilute H_2SO_4Aq , and neutralising by KHOAq, B. obtained $2V_2O_4.4V_2O_5.5K_2O.aq$. By gently heating the compound

2V₂O₄.3V₂O₅.3(NH₄)₂O.6aq (v. Hypovanadovanadates, p. 853), B. (l.c.) obtained a darkgreen solid that absorbed water from the air, and then had the composition 3(V₂O₄.V₂O₅). 8aq 2V₂O₄.3V₂O₅.3(NH₄)₂O₂.6aq $=3V_4O_9$. 8aq (cf. Ditte, C. R. 101, 151; also Manasse, A. 240, 23).

Compounds of SO_s with oxides intermediate between V_2O_4 and V_2O_5 are formed by the reducing action of zinc on solutions of V_2O_5 in H₂SO₄ (v. Rammelsberg, B. B. 1880. 787).

Vanadium oxyacids, and salts and derivatives thereof. Two oxyacids have been isolated, HVO, and $H_4V_2O_7$, corresponding with HPO, and $H_4P_2O_7$; salts of these, also salts of the tribasic acid H₂VO₄, and several more complex salts, are known. Salts derived from the hypothetical hypovanadic acid $V_4O_7(OH)_2$ (= $2\overline{V}_2O_4.H_2O$) have also been prepared; and several compounds ere known which are most simply regarded as $xV_2O_4.yV_2O_5.zMO$, and which may be called hypovanado-vanadates. V₂O₅ also combines with several acidic and basic oxides simultaneously to form compounds of the general form xV_{*}O_{**}yX:xMO; these compounds are generally described as arseno-vanadates, phospho-vanadates, &c.

VANADIC ACIDS. Two vanadic acids, HVO, and $H_4V_2O_7$, have been isolated; salts of the third acid, H_5VO_4 , are known, but the acid

itself has not certainly been prepared.

seid HVO. (Monobasic Monohydrate of vanadic oxide Metavanadic vanadic acid. V₂O₃.H₂O_.) Gerland (B. 9,872 [1876]) prepared this acid by adding an excess of NH₄ClAq to cold CuSO,Aq, then adding NF,VO,Aq till 1 permanent pp. formed, heating to 75°, washing the lustrous yellow scales that slowly separated with dilute P\$SO,Aq and SO,Aq (to remove V₂O_a), then washing with water, and drying at

100° until the smell of NH2 ceased. Manasse (A. 240, 52) obtained HVO, by decomposing conc. solutions of the Ca, Mg, or Sr salt by a little HNO, Aq, and washing the pp. with dilute HNO, Aq, and then with hot water. Metavanadic acid forms golden-yellow crystalline scales; it is unchanged in air; dissolves very slightly in water; insoluble in absolute alcohol; soluble in NH,Aq, KOHAq, or NaOHAq. By adding NH,VO,Aq to CuSO,Aq, filtering from the pp. that formed, and dialysing for several days, Gerland (l.c. p. 874) obtained a solution in the dialyser of vanadic acid, which remained clear when heated, and on continued evaporation gave a pp. of red amorphous V₂O₂. Metavanadic acid forms a series of salts (v. Metavanadates, infra).

This compound is used as a colour in place of gold bronze, under the name of vanadium bronze. According to Guyard (Bl. [2] 25, 856), the substance commonly known as vanadium bronze is really an ammonium vanadate.

Pyrovanadic acid H, V2O7. (Tetrabasic vanadic acid. Dihydrate of vanadic oxide V₂O₅.2H₂O_.) This acid is obtained by decomposing a fairly conc. solution of an alkali vanadate of the form $M_2V_4O_{11}$ or $M_2V_4O_{14}$ (v. TETRAVANADATES and HEXAVANADATES, p. 852) by HNO, Aq, washing with cold water, and drying in the air (von Hauer, J. pr. 80, 324). According to Ditte (C. R. 101, 698), the red form of $\nabla_2 O_8$ (v. p. 849) absorbs H₂O from the air, forming V₂O₂.2H₂O, and also V₂O₃.5H₂O. Pyrovanadic acid is described as a brown, amorphous powder, resembling FeO,H,; dried over H2SO, it is said to lose half its water. Very slightly soluble in water; the solution is clear yellow; it reddens litmus. H₄V₂O₇ is insol. in absolute alcohol; it dissolves readily in NH, Aq. For salts of this

acid v. Pyrovanadates (p. 852).

VANADATES. The greater number of the VANADATES. The greater number of the vanadates belong to the series (1) MVO₂, (2) M₄V₂O₇, (8) M₂VO₄, (4) M₂V₄O₁₁, or (5) M₂V₅O₁₄. Series (1), (2), and (3) may be called normal salts; the formulæ may be written as (1) M₂O.V₂O₃, (2) 2M₂O.V₂O₃, and (3) 3M₂O.V₂O₄. Series (4) and (5) may be called acid salts; the formulæ may be written as (4) M₂O.V₂O₃ and (5) M₂O.V₃O₄. Acid salts (4) M₂O.2V₂O₅, and (5) M₂O.3V₂O₅. Acid salts are also known intermediate between series (1) and (4); the chief of these are (6) $3M_2O.4V_2O_3$ and (4); the chief of these are (6) $3M_2O.5V_2O_3$ = $M_2V_3O_2s$ ($M_2O:V_2O_3 = 1:1\frac{1}{2}$), (7) $3M_2O.5V_2O_3 = M_2V_1O_2s$ ($M_2O:V_2O_3 = 1:1\frac{1}{2}$), and (8) $2M_2O.3V_2O_3 = M_2V_2O_3$ ($M_2O:V_2O_3 = 1:1\frac{1}{2}$). A few salts of the form $4M_2O.V_2O_3 = M_2V_2O_3$ ($M_2O:V_2O_3 = 1:\frac{1}{2}$) may be called basic salts. Some salts, especially those of Sr and Tl belong to more complex those of Sr and Tl, belong to more complex forms.

Roscoe (C. J. [2] 9, 28) has shown that the alkali orthovanadates, M₂VO₄ or 3M₂O.V₂O₅, are easily decomposed in aqueous solution at the ordinary temperature to pyrovanadates, M₄V₂O₂, and MOHAq; but that orthovanadates are formed, rather than meta- or pyro-vanadates, by fusing V.O. with alkali carbonates; further, that aqueous solutions of alkali pyrovanadates are readily decomposed by CO, to alkali carbonates and metavanrdates.

METAVANADATES M'VO, or M.O.V.O.

Ammonium metavanadate NH, VO. Obtained by dissolving V.O. EH, O in excess of NH,Aq, warming, and allowing to evaporate **3** 1 2

Forms a white, crystalline crust. Dissolves slowly in cold water, quickly in boiling water. Decomposed by heat, giving V2O3 when air is present. According to Berzelius (P. 22, 1), when a solution of $V_2O_8.xH_2O$ in excess of NH, Aq is allowed to evaporate without having been heated, NH, VO, separates in lemon-yellow crystals (v. B., l.c.; Roscoe, C. J. [2] 6, 322; Ditte, C. R. 102, 918; Norblad, B. 8, 126; Wagner, D. P. J. 223, 633; Böttger, C. C. 1873.

Ba(VO₃)₂. aq. A Barium metavanadate yellowish-white pp.; obtained, along with red crystals, by ppg. NH₄VO₂Aq by BaCl₂Aq (Manasse, A. 240, 23). Sl. sol. in water. Loses H₂O when heated to c. 200° (Berzelius; Norblad,

Potassium metavanadate KVO. xaq. Formed by dissolving V_2O_5 in KOHAq in the ratio V_2O_5 :KOH. Various hydrates have been obtained as white crystalline solids, melting to a clear yellow liquid, which solidifies to white, lustrous KVO₂ (v. Berzelius, *l.c.*; Ditte, C. R. 104, 902, 1061, 1168).

Sodium metavanadate NaVO₂. Obtained by Roscoe (C. J. [2] 9, 31) by fusing V₂O₃ and Na₂CO₃ in the ratio V₂O₃:2Na₂CO₃, dissolving in water, crystallising Na₄V₂O₇. 18aq, and passing CO₂ into an aqueous solution of this salt, when NaVO. crystallised out and Na2COs remained in solution. Small yellowish-white crystals, probably monoclinic prisms, melting readily and solidifying to a yellow crystalline mass (cf. Norblad, l.c.). Hydrates with 2, 4, and 5 H₂O have been obtained (cf. Ditte, l.c.).

The other metavanadates that have been prepared are $Cd(VO_8)_2$ (Ditte, C. R. 104, 1705); Ca(VO₃)₂ 4aq (von Hauer, J. pr. 69, 385; 80, 324; Manasse, A. 240, 23; cf. Roscoe, C. J. [2] 9, 32); Co(VO₃)₂. Saq (D., l.c.; Radau, A. 251, 114), forms double salts with KVO₃ (R., l.c.); 114₁, forms acouse sats with KVO_s (R., L.c.);
 Pb(VO_s)₂ (R., L.c.; D., L.c.); LiVO₃. 2aq (Ditte, C. R. 104, 1168);
 Mg(VO_s)₂. 6aq (D., L.c.);
 Manasse, L.c.);
 Mn(VO_s)₂. 4aq (Radau, L.c.);
 AgVO_s.4NH_s. 2aq (D., L.c.);
 Sr(VO_s)₂. 2aq (D., L.c.).
 Orthovandates M¹₂VO_s or 3M₂O. Ohtsined

Packetium orthoverselate K VO. Ohtsined

Potassium orthovanadate $K_3 \nabla \hat{O}_4$. Obtained by fusing V_2O_5 with K_2CO_3 ; decomposed by water, giving KOHAq and $K_4V_2O_7$ (Rammels-

berg, B. B. 1883. 8).

Sodium orthovanadate Na. VO. xH.O. Roscoe (C. J. [2] 9, 29) obtained the salt with 16H₂O by fusing V₂O₅ and Na₂CO₅ in the ratio V₂O₅:8Na₂CO₅, allowing to cool, dissolving in as little cold water as possible, quickly adding excess of conc. alcohol, allowing to stand for some hours, washing the crystals that formed with small quantities of alcohol, and drying on a porous plate over H_3SO_s in vacuo (cf. Czudnowicz, P. 120, 34; Rammelsberg, B. B. 1883. 3). Baker (C. J. 47, 853) obtained $Na_sVO_sxH_sO$ with x=8(?), 10, and 12; he also obtained the double salt $Na_sVO_sNaF.19H_sO$. The crystalline forms are given by B., and the various hydrates of Na.VO, are shown to be isomorphous with corresponding arsenates and phosphates. Na, VO, Aq changes to Na, V2O, Aq and NaOHAq; the change occurs slowly at ordinary temperatures, and rapidly at higher temperatures (B., l.c.).

The other orthovanadates that have been pre-

pared are Ca₂(VO₄)₂.CaCl₂(Hautefeuille, C. R. 103, 600; 104, 501); Pb₃(VO₄)₂ and 3Pb₃(VO₄)₂.PbCl₂ (R., I.c.); Ag₃VO₄ (R., I.c.); Tl₂VO₄ (Carnelley, C. J. 26, 323).

Pyrovanadates M1, V2O, or 2M2O.V2O5.

Barium pyrovanadate Ba₂V₂O₃. Obtained, as a white amorphous powder, by adding BaCl₂Aq to freshly-prepared Na, VO, Aq, washing, and drying at 100°. The salt is slightly soluble in

water (Roscoe, C. J. [2] 9, 83).

Potassium pyrovanadate K₄V₂O₇. Hard, deliquescent monoclinic prisms, formed by adding KOHAq to KVO, Aq, rapidly evaporating to a syrup, and placing over H2SO4 (Norblad, B. 8,

126).

Na, V20, 16H2O. Sodium pyrovanadate Formed by fusing V2O, and Na2CO, in the ratio V2Oa:2Na2COs, dissolving, and crystallising; also by exposing Na, VO, Aq to air free from CC, until addition of alcohol ppts. silby scales $(Na_4V_2O_7)$ (Roscoe, *l.c.*, p. 31). White six-sided tables; e. sol. water, insol. alcohol. $Na_4V_2O_7Aq$ is decomposed by CO₂ to Na₂CO₂Aq and NaVO₂Aq

The other pyrovanadates that have been prepared are 2Ca, V2O, 5aq (Roscoe, l.c.; cf. Ditte,

Tetravanadates M21V4O11 or M2O.2V2O5 (also

called divanadates).

Ammonium tetravanadate $(NH_4)_2V_4O_{11}$ and; x=3 or 4. Formed by adding a little acetic acid to boiling NH_4VO_3Aq and evaporating in vacuo; also by saturating NH, Aq with V2Os in a closed, warmed flask, and allowing to evaporate; red prismatic crystals with a golden-yellow sheen (Berzelius, P. 22, 1; Rammelsberg, B. B. 1883. 3; von Hauer, l.c.; Ditte, C. R. 102, 918)

Potassium tetravanadate $\mathbb{K}_{2}\mathbb{V}_{4}O_{11}$. xaq; x=8, 4, 7,and 10. Obtained by saturating K₂CO₂Aq at 80° with excess of V₂O₅ and allowing to cool; also by adding acetic acid to $\nabla_2 O_n$ in KOHAq, concentrating at 80°, and letting cool (Ditte, C. R. 104, 902, 1061, 1168; cf. Norblad, B. 8, 126). Orange-coloured tablets (v. also Gibbons, C. N. 80, 267).

The other tetravanadates are CaV₄O₁₁. 9aq (Manasse, A. 240, 23); PbV₄O₁₁ (Ditte, l.c.); Li₂V₄O₁₁. 9aq (Norblad, l.c.); MgV₄O₁₁. 9aq. (B., l.c.; von Hauer, l.c.); NiV₄O₁₁. 9aq (Radau, A. 251, 114); Na₂V₄O₁₁. 9aq (Berzelius, P. 22, 1; von Hauer, l.c.; Norblad, l.c.); SrV₄O₁₁. 9aq (B., l.c.; von H₄).

von H., l.c.).

HEXAVANADATES M2 1V O16. xH2O or

M20.8V20, xH20 (also called trivanadates). These salts have been described by Norblad (B. 8, 126) and Ditte (C. R. 104, 902, 1061, 1168). The chief are those where $M_2 = (NH_4)_2$

Cd, Ca, K₂ and Na₂.

The principal vanadates belonging to other mantioned are the folseries than those already mentioned are the fol-

The state of the

1048 ; Manasse, l.c. ; Norblad, l.c.) ; (4) $4M_2O.V_2O_5 (= M_8V_2O_9)$, M = Li and Na (v. Ditte, l.c.).

Several other vanadates not belonging to any of these series have also been described; for Ca and Sr salts (and double salts), v. Manasse (l.c.);

and for Tl salts v. Carnelley (C. J. 26, 323).

HYPOVANADATES. Salts of hypothetical hypovanadic acid $V_{\bullet}O_{\bullet}(OH)_{2}$ (=2 $V_{\bullet}O_{\bullet}H_{\bullet}O)$. These salts belong to the form $M^{1}_{\bullet}V_{\bullet}O_{\bullet}xH_{\bullet}O$ (= $M_{\bullet}O.2V_{\bullet}O_{\bullet}xH_{\bullet}O$). The formation of the NH₄ and K salts was described by Berzelius; a series of salts was prepared and examined by Crow (C. J. [2] 15, 458, The alkali salts are obtained by treating fairly conc. VO2ClAq (v. HYDRATED VANADIUM DIOXYMONOCHLORIDE, p. 854) with excess of caustic alkalis, allowing the pps. to settle in closed vessels, washing with alkali solution, then with dilute alcohol containing a little acetic acid, and finally with alcohol alone, and drying between paper. The salts of Pb and Ag are ob-tained by ppn. from a solution of the K salt. The salts form brown to black crystalline powders; the alkali salts dissolve in water. Crow prepared salts with M, = (NH₄), Ba, Pb, K₂, Ag₂, and Nag

HYPOVANADO-VANADATES. Salts of the form $nV_2O_4.mV_2O_5.pM_2O.xH_2O$. Alkali salts of this series were obtained by Brierley (C. J. 49, 32) by reducing V_2O_5Aq by SO_2 , adding V_2O_5 in alkali, then a slight excess of alkali, and then acetic acid; pouring into alkali acetate solution, washing the pps. with alcohol, and drying over CaCl₂. The salts crystallise in greenish-black, or black, prisms. The following were prepared:

(1) 2V₂O₄, V₂O₃, 2Na₃O.13H₂O; (2) 2V₂O₄, V₂O₃, 2K₂O.6H₂O; (3) 2V₂O₄, 2V₂O₅, (NH₄)₂O.14H₂O; these salts are soluble in water: (4) 2V₂O₄.4V₂O₅.5K₂O.H₂O; (5) 2V₂O₄.4V₂O₅.3(NH₄)₂O.6H₂O; these are insoluble in water.

COMPLEX VANADIC ACIDS AND SALTS. Vanadic oxide V2O5 combines with several anhydrides, such as As₂O₃, I₂O₃, P₂O₃, &c., and many of the compounds thus produced combine with The compounds of V2O, with basic oxides. certain anhydrides are generally regarded as acidic; and the compounds formed of V₂O₄, anhydrides, and bases are usually classed as saltlike substances. It is quite likely that many of the compounds of V_2O_3 with anhydrides, such as As_2O_3 , P_2O_3 , &c., would be better classed as salts, just as the compounds of V_2O_3 with SO_3 are classed as salts.

ABSENO-VANADIC ACIDS AND SALTS. By boiling V₂O₅ with excess of H₂AsO₄Aq, and concentrating the solution, Fernandez (B. 17, 1632) obtained lustrous, golden-yellow crystals of V₂O₅.As₂O₅.11aq (cf. Friedheim a. Schmitz-Dumont, B. 23, 2600). Gibbs (P. Am. A. 21, 50) obtained $x\nabla_2O_s.yAs_2O_s.xaq$ (x and y probably either 5 and 8, or 7 and 6), by decomposing a mixture of Na vanadate and arsenate by HNO, Aq. For various compounds of the form $mV_2O_5.nAs_2O_5.pMO$, where m and n are 1 and 1, 2 and 3, 1 and 2, 2 and 1, &c., and p varies from 1 to 5, v. F. a. S.-D. (l.c.). IODO-VANADIO ACIDS. Ditte (C. R. 102, 757)

says that compounds of V2O, and LO, are formed by heating V.O. with conc. HIO.Aq; he

gives formulæ, but no analyses.

MOLYBDO-VANADIC ACIDS AND SALTS. For descriptions of numerous compounds of the forms $x\nabla_2O_3.yMoO_3.eaq$, $x\nabla_2O_3.yMoO_3.pMO.eaq$, and also $x \nabla_2 O_3 \cdot y \text{MoO}_3 \cdot m P_2 O_3 \cdot p \text{MO} \cdot saq$, v. Gibbs (P. Am. A. 18, 232).

PHOSPHO-VANADIC ACIDS AND SALTS. compounds of V_2O_5 , P_2O_5 , and H_2O are described by Gibbs $(P.\ Am.\ A.\ 21,\ 50)$; Ditte $(C.\ R.\ 102,$ 757) also assigned formulæ to substances obtained by heating V₂O₅ with conc. H₂PO₄. Friedheim (B. 23, 1530) obtained V₂O₅.P₂O₅.2H₂O by dissolving V2O, in syrupy H2PO4.

For descriptions of a great many compounds of the form $xV_2O_5.yP_2O_5.mMO.zaq$, v. Gibbs

(l.c.); and cf. Friedheim (l.c.).

F. divides the compounds formed by reacting with V₂O₅ on alkali phosphates into two classes, which he calls luteo and purpureo com-

pounds.

Luteo- compounds are formed by (1) the interactions of phosphoric and vanadic acids in solution; (2) the interactions of solutions of phosphates and vanadates in presence of small quantities of acids; (3) the interactions of solutions of phosphates and vanadic acid; (4) the interactions of vanadates and phosphoric acid, by heating solutions nearly to boiling and eva-porating. These compounds are yellow, granular, and indistinctly crystalline; the alkali compounds are very slightly soluble in water, solution being accompanied by decomposition. The following luteo- compounds are described by F .:

V₂O₃,P₂O₄,2H₂O₅. 9aq, V₂O₃,P₂O₄,(NH₄)₂O,H₂O₅. 2aq, 2V₂O₃,P₂O₅,(NH₄)₂O, 7aq, V₂O₅,P₂O₅,K₂O,H₂O₅. 2aq, and $2\bar{V}_2\bar{O}_3.\bar{P}_2\bar{O}_3.\bar{K}_2O.\bar{7}$ aq.

Purpureo- compounds are formed by dissolving vanadic acid in conc. solutions of alkali phosphates, and by adding phosphoric acid to conc. solutions of alkali vanadates; in each case the solutions should be strongly heated. The purpureo- compounds described by F. belong to the form 12V2Os.P2Os.7M2O. 26aq; they are dark-red crystalline substances, sol. in water.

F. regards the luteo-compound of the form V₂O₃.P₂O₃.2H₂O.9aq as acid phosphate of the radicle VO₂, thus (VO₂)H₂PO₄.9aq; he compares this with the acid orthophosphates MH₂PO₄. The compounds of the form MH_2PO_4 . The compounds of the form $\nabla_2O_3.P_2O_3.M_2O.H_2O.x$ aq are looked on as double acid phosphates of alkali metal and VO, thus NH, (VO,) HPO. The luteo-compounds 2V2O5.P2O5.M2O. xaq are formulated by F. as double compounds of alkali metavanadates and thus 2V₂O₃.P₂O₃.K₂O.2H₂O. 5aq ₂)H₂PO₄.2½aq). The purpureoto the form 12V,O,P,O,7M,O. xaq are looked on by F. as double compounds of acid vanadates and acid pho phates; this view of their constitution is expressed by the statement $12V_2O_3P_2O_3.7M_2O.xH_2O =$

 $P_2O_3.2M_2O.H_2O + 12V_2O_3.5M_2O + (x-1)H_2O.$ These views of F. are considerably modified in a later paper (Zeit. f. anorg. Chemie, 5, 437), where the purpureo- compounds are looked on as divanadates with some V₂O₅ isomorphously replaced by P.O. 'According to F., many of the substances described by Gibbs and by Ditte do not exist.

Berzelius (P. 22, 1) described a compound

of V₂O₅, P₂O₅, and SiO₂, with H₂O, which may be named phospho-silico-vanadic acid.

TUNGSTO-VANADIC ACIDS AND SALTS. The compounds of V2O, with WO, and bases that have been described are very numerous, and the formulæ given are generally extremely complex (v. Gibbs, P. Am. A. 18, 232; Rosenheim, A. 251, 197, 234; and especially Friedheim, B. 17, **1505**; 23, 1505; 24, 1173).

Vanadium, oxybromides of. Two compounds of V with O and Br have been isolated; VOBr₂ and VOBr₃, corresponding with two of the five exychlorides. According to Schafarik (W. A. B. 33, 14; 47 [11] 251), other oxybromides also

Vanadium oxydibromide VOBr₂. (Vanadyl dibromide.) A brown, deliquescent powder. Prepared by heating VOBr₃ (v. infra); decomposition to VOBr₂ and Br occurs slowly at temperatures below 180°, and suddenly at 180°. Heated in air forms V_2O_5 ; dissolves in water, forming a blue solution (Roscoe, C. J. [2] 9, 26). Ditte (C. R. 102, 1310) says that a compound VOBr₂4H₂O is formed, as dark-green, deliquescent crystals, by treating red V2O, with

HBr. D. writes the formula V₂O₃Br₃.2HBr.7H₂O.

Vanadium oxytribenomide VOBr₃. (Vanadyl tribromide.) Prepared by passing vapour of pure Br over V₂O₃ heated to redness, condensing the yellowish-white vapours that are formed, removing excess of Br from the red liquid by heating to 40°-50° at c. 100 mm. pressure in a current of perfectly dry air, and then distilling in vacuo (Roscoe, C. J. [2] 9, 24). VOBr, is a red, very hygroscopic liquid that boils at 180°-136° at 100 mm. pressure; S.G. 2.9673 at 0°, 2.9325 at 14.5°; decomposes slowly at the ordinary temperature, and suddenly at 180°, to VOBr. and Br (R., l.c.).

Vanadium, oxychlorides of. Five compounds of V with O and Cl have been isolated: VOCl,

VOCl₂, VOCl₂, V₂O₂Cl, and VO₂Cl. xaq.

VANADIUM OXYMONOCHLORIDE VOCI. (Vanadyl monochloride.) Obtained by the regulated reduction of VOCl, by H. The mixed vapours are passed through a red-hot tube; VOCl is de-posited near the end whereat the vapours enter the tube. A brown, light, flocculent powder; insol. in water; easily dissolved by HNO, Aq

(Boscoe, C. J. [2] 6, 347).

VANADIUM OXYDICHLORIDE VOCI2. (Vanadyl dichloride.) Prepared by heating a slight excess of VOCl, with zinc for some days in a sealed tube, at 400°, cutting off the part of the tube on which the sublimate of VOCl2 has formed (V2O4 and ZnCl₂ are the other products), quickly placing it in a wider tube, and removing VOCI, by heating in a stream of dry CO₂ at 130°. Also obtained by passing vapour of VOCI₂ and H through a red-hot tube; VOCl, is deposited near the end of the tube whereat the gases enter, and V.O.Cl at the further end of the tube (Roscoe, l.c., p. 348). Lustrous, grass-green, deliquescent tablets; S.G. 2.88 at 13°; slowly decomposed by water; soluble in dilute HNO,Aq. A compound with PtCl, having the composition 2(VOCl2.PtCl4). 21aq, is said by Brauner (M. 3, 58) to be formed by dissolving V₂O₄ in conc.

HClac in presence of PtCl,Aq.

Vanadium oxytrichloride VOCl. (Vanadyl

trichloride.) Mol. w. 178.27.

Formation.—1. By heating V2O2 gently in a stream of Cl (Berzelius, P. 22, 1).-2. By heating a mixture of V₂O₅ and C in Cl.

Preparation.—A mixture of V2O5 and sugarcharcoal is heated to redness in a stream of dry H, and allowed to cool in that ges; a stream of dry Cl is then passed over the mixture, which is heated to redness, and the product is condensed; the liquid thus obtained is heated for some hours in a flask with an inverted condenser, in a current of dry CO₂, and then rectified over Na (Roscoe, C. J. [2] 6, 842).

Properties and Reactions .- A clear, mobile, lemon-yellow liquid (Berzelius l.c.; Roscoe, l.c.; Schafarik, A. 109, 85). S.G. 1.836 at 17.5°, 1.828 at 24° (R., l.c.); 1.86534 at $\frac{0^{\circ}}{0^{\circ}}$, 1.63073 at b.p. (Thorpe, C. J. 37, 348). B.P. 127·19° (T., l.c.). S.V. 106·25. Does no solidify above -15° (R., l.c.). V.D. 88·3° at 186° (R., l.c.; cf. T., C. N. 24, 827; Schatarik, l.c.). Gives off thick, yellowish-red fumes in the air. Dissolves

in a little water, forming a thick, red liquid; dissolves in much water, forming a clear yellow liquid; dissolves in absolute alcohol with a red colour, but the solution soon becomes blue, owing to reduction. Heated to 60°-70° in a closed tube with ether forms needles of the composition VOCl₂.Et₂O (Bedson, C. J. 29, 309).

DIVANADIUM DIOXYMONOCHLORIDE (Divanadyl monochloride.) Yellow-bronze, microscopic crystals, resembling mosaic gold; formed by passing VOCl, and H through a redhot tube. The crystals are deposited on the part of the tube farthest from the end whereat the gases enter (Roscoe, C. J. [2] 6, 348). Insoluble in water; soluble in HNO, Aq. compound was thought by Schafarik (A. 109, 85) to be V.

HYDRATED DIOXYMONOCHLORIDE VANADIUM

VO.Cl. xaq. (Hypovanadic chloride V.O.Cl. xaq.) Formed, as a brown, deliquescent solid, by passing H_2S into a solution of V_2O_5 in hot HClAq, filtering from S, and evaporating (Crow, C. J. 30, 457). Heated in a stream of dry CO₂ gives off HCl and H₂O, and leaves V₂O₄. C. (l.c.) gives the hydration as $V_2O_4Cl_2$. 5aq.

Vanadium, exyfluorides of. No compound of V with O and F has been isolated with certainty. V2O2 dissolves in excess of HFAq; on evaporation, solution in water, and evaporation over H₂SO₄, Petersen (J. pr. [2] 40, 194) obtained blue, microscopic crystals which he supposed might be VOF₂. xaq, inasmuch as a solution of these crystals in HFAq treated with fluorides yielded salts of the form xMF.yVOF, (v. Hypo-VANADOXYFLUORIDES, p. 845). P. did not obtain enough of the blue crystals for analysis.

Vanadium, oxyiodides of. According to Ditte (C. R. 102, 1310), an oxyiodide 2VOL, 9H,O is formed, as a black, deliquescent mass, by adding excess of HIAq to a warm solution of red V₂O₂, removing free I by shaking with finely-divided Ag, filtering, and evaporating in vacuo. The compound is said to react with NH₄Aq, giving (NH₄)₂V₄O₉. D. writes the formula as V₂O₁L₂2H1.8H₂O; but no analyses are given.

No reaction occurs when I is heated with V₂O₂ (Roscoe, C. J. [2] 9, 28); nor when V₂O₃ is digested with water, or alcohol, and I (Guyard,

Bl. [2] 25, 351). V_2O_5 and I_2O_5 probably com-

bine (v. Ditte, C. R. 102, 757, 1019).

Vanadium, oxysulphides of. None of these compounds has been certainly isolated. Oxysulphides may perhaps be formed by decomposing alkali thiovanadates in solution by dilute

acids (v. Kay, C. J. 37, 728).

Vanadium, salts of. Compounds of the oxides V2O3, V2O4, and V2O3 with acidic oxides have been isolated; the formulæ of one or two of the compounds may be written as derived from oxyacids by replacing H by V, but it is simpler, and more in keeping with the facts that are known, to re; resent all the compounds by the general formula V₂O₂.yX, where X is an acidic oxide. The best-studied compounds of V₂O₄ with acidic oxides may be represented as salts of the radicle V₂O₂ (divanadyl), or of the radicle VO (vanadyl); and the compounds of V2Os with acidic oxides as acid salts of the radicle VO2. The suphates have been better studied than the other salts; the following table presents the compositions of the sul-

I. Vanadous sulphates (v. p. 847): $V_2O_3.4SO_3.xaq = V_2H_2(SO_4)_4.yaq (y=x-1)$; $??V_2O_3.3SO_3 = V_2(SO_4)_3$

II. Hypovanadic sulphates (v. p. 848):

 $\begin{array}{l} V_2O_4.2SO_4 = V_2O_2.(SO_4)_3 = (VO)SO_4; \\ V_2O_4.3SO_3.\ xaq = V_2O_2.H_2(SO_4)_3.\ yaq\ (y=x-1) \\ = (VO.SO_4)_2.H_2SO_4.\ yaq. \end{array}$

III. Vanadic sulphates (v. p. 850): $V_2O_3.2SO_3 = (VO_2)_2SO_4.SO_3$; or if the salt contains xaq, then $V_2O_3.2SO_3.xaq = (VO_2)_2SO_3.xaq = (VO_2)_2SO_3.xaq$

 $(VO_2)_2SO_4$, H_2SO_4 , yaq (y=x-1); V_2O_3 .3SO₃ = $(VO_2)_2SO_4$.2SO₃; or if the salt contains xaq, then

 $V_2O_3.3SO_3. xaq = (VO_2)_2SO_4.2H_2SO_4. yH_2O_3$

(y = x - 2).Alkalis probably ppt. V₂O₃, xaq from solutions of vanadous sulphates (v. p. 847); from solutions of hypovanadic sulphates alkalis ppt.

V₂O₄, xaq (v. p. 848). V₂O₂ dissolves in oxyacids, forming lavender solutions which probably contain V₂O₂ xX, where

X is an acidic oxide (v. p. 846). Various salts of V_2O_3 , V_2O_4 , and V_2O_5 besides the sulphates probably exist, but they have not been thoroughly examined (v. HYPOVANADIO SALTS, p. 848; VANADIC SALTS, p. 850; ARSENO-VANADIC ACIDS AND SALTS, p. 853; IODO-VANADIC ACIDS, p. 853; MOLYBDO-VANADIC ACIDS AND SALTS, p. 853; Phospho-vanadic acids and salts, p. 853; and Tungsto-Vanadic acids and salts, p. 854).

Vanadium, silicide of. Roscoe (C. J. [2] 8, 858) says that V acts on glass or porcelain vessels when strongly heated therein, forming a compound with Si; tubes in which V chlorides are reduced by heating in H get coated with a grey lustrous mirror of this compound. No details or

analyses are given.

Vanadium, silicofluoride of. By boiling V₂O₅ with H₂SiF₆Aq and alcohol, Guyard (Bl. [2] 40, 352) obtained a greyish, uncrystallisable mass which he took to be a silicofluoride of V; no analyses or details of preparation are given.

Vanadium, sulphides of. Berzelius ($P_{\perp}22$, 1) described two compounds of V and S, one obtained by ppg. a solution of an alkali thiovanadate by dilute acid, and the other by heating V₂O₂ in H₂S. B. gave the formula VS₂ and

VS, to the compounds he described. The sulphides of V were re-examined by Kay in 1880 (C. J. 37, 728), who found that the pp. obtained by adding dilute HClAq or H2SO, Aq to a solution of sodium vanadate saturated with $H_{\nu}S$, or to solution of $V_{\nu}O_{\nu}$ in an alkali hydrosulphide, contained O_{ν} but probably had not a definite composition; and that the product of heating V_2O_3 in H_2S is the trisulphide V_2S_3 ; Kay prepared three sulphides, V_2S_2 , V_2S_3 and V_2S_3 , corresponding to three of the four oxides; the sulphides dissolve in solutions of alkali sulphides.

probably forming thiovanadates (q. v. infra).

Vanadium disulphide V₂S₂. (Hypovanadous sulphide.) Prepared by heating V₂S₃ to full redness, for a long time, in H quite free from O. Forms black lustrous plates, or a brownish-black powder; S.G. 4.2 to 4.4. Absorbs O very quickly when heated in air, giving V_2O_3 , V_2O_4 , and then V_2O_5 , with evolution of SO_2 . Not acted on by boiling HClAq, dilute or conc., nor by boiling dilute H₂SO₄Aq, or cold conc. H₂SO₄. Dissolves in hot conc. H₂SO₄; dissolves in HNO₅Aq. Slowly acted on by NaOHAq or KOHAq, hot or cold; dissolves in (NH₄)₂SAq or KHSAq, forming purple to reddish-brown solutions (Kay, Lc. p. 735).

(Vanadous VANADIUM TRISULPHIDE V2S. sulphide.) Formed by strongly heating V2O, in a stream of H2S (Berzelius, l.c.; Kay, l.c. p. 736); also by heating any chloride of V, or VOCl, to redness in H2S (K., l.c.); and by strongly heating V2O, in a stream of vapour of CS, so long as any reaction occurs (K., l.c.). Forms dark, lustrous plates, or a black amorphous powder; S.G. 3.7 to 4.0. Oxidised by heating in air, finally to V₂O₂, giving off SO₂. Scarcely acted on by dilute HClAq or H₂SO₄Aq, hot or cold, nor by conc. HClAq; readily oxidised by HNO,Aq. Soluble in (NH₄)₂SAq or KHSAq, forming purplered to golden-red solutions; also somewhat soluble in KOHAq, NaOHAq or NH.OHAq (Kay, l.c.).

VANADIUM PENTASULPHIDE V₂S₃. (Vanadic sulphide.) Prepared by mixing V₂S₃ with 1 its weight of pure powdered S, heating to fusion (c. 400°) in a narrow tube quite filled with CO, and sealed, cooling, and dissolving out excess of 8 with CS₂. A black powder; S.G. 3.0. Heated in a gas that does not react with it, V2S, gives off S and leaves V₂S₃; heated in air gives off SO₂, forming V₂O₃ and finally V₂O₅. Behaves towards acids similarly to V₂S₃. Dissolves in hot NaOHAq; also in (NH₄)₂SAq or KHSAq, forming yellow to red solutions (Kay, l.c. p. 738).

Vanadium thio-acids, and salts thereof. compound of V with H and S has been isolated, but some thiovanadates, and also some oxythiovanadates, have been prepared by Krüss a.

Ohnmais (B. 23, 2547).

Ammonium thio-orthovanadate (NH4), VS4. Obtained, in purple crystals, resembling KMnO., S.G. 1.62, by passing H.S into a solu-tion of NH, VO. in NH, Aq S.G. 1898; the solution is saturated in the cold, and is kept cold while H.S is passed in. A brown pp. is formed, and this dissolves after passing in more H2S to a dark-violet liquid, from which the salt crystallises after some time.

Sodium monoxy-thio-orthovanadate Na, VOS, 5aq. Obtained by saturating 80 c.c. NaOHAq S.G. 1.122, with H2S, adding a solution of 3 g. Na, V2O, in 6 c.c. water, cooling the liquid by ice, and passing in H2S for 4 hours. The salt was also obtained without water of crystallisation by Krüss (Zeit. f. anorg. Chemie, 3, 264) by fusing a mixture of V_2O_5 , Na,CO_s, and S, until excess of S was removed. Small crystals, that melt at the ordinary temperature to a red, oily liquid (K. a. O., l.c.).

Sodium trioxy-thio-orthovanadats Na, VO, S. 10aq. Obtained by heating Na, V2O, Aq to boiling, adding freshly prepared NaSHAq, cooling by ice, and adding alcohol. A red oil, that solidifies to a crystalline mass which melts

at 18° (K. a. O., l.c.).

Ammonium monoxy-thio-pyrovanadate (NH₄)₄V₂OS₆. Formed by passing H₂S into a cooled solution of NH₄VO₃ in NH₃Aq S.G. greater than .898, and allowing the liquid to stand for some months, when crystals separate having S.G. 1.715 (K. a. O., l.c.).

Potassium monoxy-thio-pyrovanadate K₄V₂OS₄. 3aq. Obtained by passing H₂S into KVO₃ in KOHAq S.G. 1·472, air being excluded and the liquid cooled by ice. Crystals resemble KMnO4; S.G. 2.144; loses all water slowly at 150°. By evaporating the mother-liquor from this salt in vacuo, large crystals of the salt 2K₄V₂OS₆. 3aq are obtained (K. a. O., l.c.).

M. M. P. M. VANADOXYFLUORIDES and Hypovanad-

ocyfluorides; v. pp. 844-5.
VANADYL COMPOUNDS; compounds of the radicle VO: v. Vanadium oxybromides (p. 854), Vanadium oxychlorides (p. 854), and Hypo-▼ANADIC SALTS (p. 848).

VANILLIC ACID v. Methyl derivative of

PROTOCATECHUIC ACID.

VANILLIN v. Methyl derivative of Pro-TOCATECHUIC ALDEHYDE.

VANILLO-DIACETONAMINE v. ACETON-AMINE

VAPOUR DENSITIES. The term 'vapour density' is now generally employed to signify the specific gravity of a gas referred to hydrogen as unity. For descriptions of the principles of the methods used in determining vapour densities, v. Densities, relative, vol. ii. p. 374; and for an account of the application of vapour densities to finding molecular weights, v. Atomic AND MOLECULAR WEIGHTS, vol. i. p. 340.

VEGETABLE PROTEÏDS v. PROTEÏDS.

VERATRALBINE v. JERVINE.

VERATRIC ACID v. Di-methyl derivative of PROTOCATECHUIC ACID.

Homo-veratric acid v. DI-OXY-PHENYL-ACETIC ACID.

VERATROLE v. Di-methyl ether of Pyro-CATECHIN.

VERATRUM ALKALOIDS.

Veratrine C₃₂H₄₀NO₉. Cevadine. [205°]. S. 12 at 15° Occurs in the seeds of Veratrum Sabadilla (Meissner, N. J. T. 5, 3; Pelletier a. Caventou, A. Ch. [2] 14, 69; Couerbe, A. Ch. [2] 52, 352; Merck, A. 95, 200; Ar. Ph. 231, 135; Delondre, J. Ph. [3] 27, 417; Weigelin, C. C. 1872, 229; Schmidt, Ar. Ph. [8] 10, 511; B. 9, 1115; A. 185, 224). Occurs also in the root of Sarracenia purpurea (St. Martin, Z. [2] 2, 442; Hétet, C. R. 88, 185). Prepared by extracting the seeds with boiling alcohol containing a little tartaric acid, concentrating the extract, adding water, filtering from resin, adding Na,CO,, and shaking with ether. The ethereal solution is shaken with dilute tartaric acid solution, and the acid solution mixed with Na₂CO₂ and extracted with ether. The ethereal extract is mixed with ligroin and allowed to evaporate spontaneously, when a viscid mass first separates, followed by crystals which are rccrystallised from alcohol (Wright a. Luff, C. J. 83, 338). If commercial veratrine [144°] be dissolved in alcohol at 70°, water added till turbidity ensues, and the solution evaporated at 50° to 60°, crystalline veratrine separates first, then a resinous mixture of veratrine and veratridine, while the mother-liquor contains veratridine and veratroïn veratrate (Rosetti, Ar. Ph. [3] 21, 81).

Properties.—Needles (from alcohol), sol. ether, insol. water. Inactive to light. poisonous, a small quant y producing vomiting and purging. Introduced into the nose it produces sneezing. Alkaline to test papers H₂SO₄ forms a yellow colour changing to crimson, 1 pt. colouring 3,000 pts. H2SO4 (Vasmer, Ar. Ph. 2, 74). Conc. HClAq forms a violet solution on warming. HNO₃ forms a red solution, becoming yellow. Veratrine mixed with sugar (3 pts.) is coloured by H₂SO₄ dark green and finally deep blue (Weppen, Fr. 13, A solution of ammonium selenite (1 g.) in H_2SO_4 (20 c.c.) gives a yellow colour at 30°; in 3 hours a red pp. is formed, the liquid remaining yellow (Da Silva, C. R. 112, 1267). On heating with alcoholic potash or baryta it yields angelic acid and cevine (W. a. L.; Rosetti; Stransky, M. 11, 482). ICl forms a yellow flocculent pp., sol. hot HClAq (Dittmar, B. 18, 1612). Conc. HClAq yields tiglic acid. Veratrine yields tiglic acid and (3)-methyl-pyridine on distillation. On distilling veratrine with lime the products are (8)-methyl-pyridine and its hexahydride and isobutyric acid (Ahrens, B. 23, 2705). trine is not affected by boiling dilute H2SO.

Salts.—B'HCl. Amorphous.—B'2H2PtCl4: amorphous.—B'HAuCl. Yellow needles (from alcohol).—B'HAuCl. 2aq (Rosetti, J. 1883, 1351). -B'HHgCl. Crystalline pp., v. sol. alcohol.-B'2H2SO4 (dried at 100°).—B'HI3. Reddishbrown amorphous solid (Bauer, J. 1874, 861).

Benzoyl derivative C₃₂H₁₈BzNO₉. [170°-180°]. Brown crystals (containing 1 aq) (from

ether).—B'HAuCl.

Dibromide C32H49Br2NO9. Formed by allowing the tetrabromide to stand in contact with dilute KOHAq. Light-yellow amorphous solid

Tetrabromide C₃₂H₁₉Br₄NO₉. Formed by shaking veratrine with bromine-water. Yellow amorphous powder, insol. water, v. sol. alcohol

and ether (Ahrens, B. 23, 2701).

Cevine $C_{27}H_{48}NO_{9}$ (W. s. L.); $C_{27}H_{48}NO_{9}$ (Rosetti). Cevedine. [145°] (W. a. L.); [182°–185°] (R.). Formed by boiling veratrine with alcoholic NaOH (W. a. L.). Yellow resin, v. sol. alcohol, sl. sol. ether. Its aqueous solution becomes turbid on warming. Does not attack the mucous membrane, gives a crimson colour with H.SO,, and a brown colour with cane-sugar and H₂SO₄. Its salts are amorphous.—B'HHgI, (dried at 100°). Precipitate.

Veratridine C₃, H₃₅NO₁₁ (W. a. L.); H₄₆NO₉ (Rosetti). Veratrine. [1 C₁₂H₄₈NO, (Rosetti). [180° cor.] (W. a. L.); [150°-155°] (R.). S. 8 at 15°. If the viscid mass which separates before veratrine when a solution of the crude base in alcoholligroin is evaporated be shaken with ether, cevadilline remains undissolved, while veratridine dissolves in the ethor (Wright a. Luff). Amorphous resin, sl. sol. ether. Decomposed by alcoholic potash into veratric acid and verine $C_{28}H_{18}NO_8$ [c. 95°] (W. a. L.) or veratroin $C_{38}H_{22}N_1O_{16}$ [143°-148°] (R.), an amorphous base, sol. ether. Veratridine dissolves in boiling water, being converted into veratroin veratrate $\rm C_{55}H_{92}N_2O_{16}C_9H_{10}O_4$ 2aq (Rosetti), which melts, when anhydrous, at 165°-170°. $\rm H_2SO_4$ forms a yellow solution which turns crimson.—Salts. B'HAnCl, Amorphous. - B'H2SO, 10aq, crystalline.

Cevadilline C₃₄H₃₀NO₃. Obtained as above (W. a. L.). Amorphous, sl. sol. ether, m. sol. benzene. Decomposed by alcoholic potash.

Salts. - B'HAuCl₄. - B'HHgI₃. Gelatinous. Cevadilline is perhaps identical with the sabadilline $C_{41}H_{da}N_2O_{13}$ of Wiegelin (C. C. 1872, 229), to which Hesse (A. 192, 186) assigns the formula $C_{21}H_{35}NO_7$; while sabatrine $C_{51}H_{86}N_2O_{17}$ (W.) or $C_{29}H_{15}NO_{9}$ (H.) was probably a mixture of decomposition-products (Wright a. Luff).

Veratrum aloum. The alkaloids in the root of this plant, and in that of V. viride, are de-

scribed under Jervine

VERATRUMIC ACID is VERATRIC ACID.

VERNIN C₁₈H₂₀N₈O₈ 3aq. Occurs in young vetch plants (Vicia sativa), in young red clover (Trifolium pratense), in pumpkin seeds, in ergot, and in the blossom of Corylus avellana and Pinus sylvestris (E. Schulze, J. pr. [2] 32, 447; H. 10, 80, 326). Separated from asparagin by crystallisation from hot water. Minute silky prisms, v. sol. hot water, insol. alcohol. Its aqueous solution is neutral, gives no pp. with Pb(OAc)2, but is ppd. by AgNO2, by pieric acid, and by phosphotungstic acid in presence of HCl. Boiling hydrochloric acid forms guanine.-Ag₂C₁₆H₁₈N₈O₈. Gelatinous pp.

VERNONIN C₁₀H₂₄O₇. Occurs in the root of

Vernonia nigritiana, used on the west coast of Africa as a febrifuge (Heckel a. Schlagdenhauffen, C. R. 106, 1446). White powder, sl. sol. ether and chloroform, sol. alcohol. Conc. H2SO, gives a brown colour, changing to purple. Cardiac poison, 80 pts. being required to produce the effect of 1 pt. of digitalin. Decomposed by boiling dilute HCl into glucose and a resin

 $C_4H_{10}O_3$.

VICIN $C_{28}H_{51}N_{11}O_{21}$. S. 1 at 23°. Occurs in the seeds of *Vicia sativa*, *V. Faba*, and Pitthausen *J. pr.* [2] 2, 333; V. Faba minor (Ritthausen, J. pr. [2] 2, 333; 24, 202; 29, 359). Obtained by extracting the seeds with cold dilute H2SO4 (1:50), neutralising with lime, filtering from CaSO4, evaporating to dryness and crystallising from 85 p.c. alcohol. The yield is 25 p.c.

Properties.—Tufts of small needles, nearly Loses 2H₂O at 160°. Sol. insol. alcohol. alkalis and alkaline earths, reppd. on neutralisation. Not eaffected by boiling baryta-water. Sol. dilute HCl and H2SO, but after boiling the solution gives a deep-blue colour with FeCl and

and a violet pp. with baryta. Boiling KOHAq (S.G. 1.1) also forms divicin. evaporated with HNO, (S.G. 1.2) the residue is edged with deep violet. Potash-fusion forms

Salts. - B',4H,SO, - B',11HCl. Slender

needles.

Divicin C₈₁H₅₀N₈₀O₁₆. By boiling vicin with water (5 pts.) containing H₂SO₄ (1 pt.) at 0° there is formed crystalline (C₃₂H₂₈N₂₀O₉)2½SO₄, which yields divicin on treatment with exactly the calculated quantity of KOHAq. Flat prisms (from water). Reduces AgNO, at once. Not ppd. by baryta. A little FeCl, followed by NH, gives a splendid blue colour. B'8HNO₃. Whetstone-shaped crystals, got by adding HNO, to a solution of divicin.

Convicin C20H28N6O14 2aq. Obtained from powdered vetch seeds by extracting with alcohol, allowing vicin to crystallise from the extract, ppg. the mother-liquor with HgCl2 and potash, decomposing the pp. with aqueous H.S. and evaporating. The mixture of vicin and convicin so obtained is treated with dilute H2SO4, which dissolves the vicin only. Thin plates (from water), often resembling leucine. Hardly sol. cold water, sl. sol. alcohol. Not decomposed by boiling KOHAq (S.G. 1·1). Potash-fusion gives off NH, but forms no KCy. Insol. cold dilute HCl and H₂SO₄. Its aqueous solution gives a flocculent pp. with Hg(NO₃)₂.

VINACONIC ACID v. TRI-METHYLENE DI-

CARBOXYLIC ACID.

VINCETOXIN $C_{16}H_{12}O_{6}$. [59°]. [a]_D = -50°. Extracted by milk of lime from powdered asclepias root (Tanret, C. R. 100, 277; Bl. [2] 43, $6\overline{2}0$). Occurs in two forms, one soluble and one insoluble in water. Both forms are lævorotatory, amorphous, sl. alcohol, insol. ether. Boiling dilute HCl yields an amorphous, inactive sugar, which does not ferment with yeast.

VINYL ALCOHOL CH,:CH.OH. This substance might be expected to be identical with aldehyde CH₈.CHO, but Poleck a. Thümmel (B. 22, 2863) suppose it to be present in ether that has been exposed to air and sunlight, and that its presence is indicated by the formation of a pp. C₂H₂OHgOHg₂Cl₂ when a solution of mercury oxychloride in Na2CO3Aq is added to commercial ether. This pp. is white, and is converted by boiling potash into explosive greenish-black 'acetylene mercury' C2H3O4Hg4, and by adding nitric acid to its alkaline solution into C2HHg2OCl2, which is not explosive. H2S passed into water containing the compound C₂H₃O₂Hg₃Cl₂ yields (γ)-tri-thio-acetic aldehyde [76°]

VINYL-DIACETONAMINE v. ACETONAMINES. VINYL-AMINE CH2:CH.NH2. Formed by the action of moist Ag2O or of KOHAq on brom -ethylamine hydrobromide at 48° (Gabriel, B. 21, 1049, 2665). Known only in aqueous solution. Volatile with steam. Decomposes in aqueous solution even in the cold. SO₂ converts it into taurine.

Salts .- *B'HCl. Poisonous. Its aqueous solution is decomposed by heat.—B'₂H₂PtCl_e. Crystals, v. sol. water.—B'₂BiI₂. Minute scarlet hexagonal leaves.—B'HAuCl_e. Golden G∩lden crystals.—B'C.H.N.O. [142°]. Slender yellow needles.

Reference. — DI-IODOVINYLAMINE.
VINYL BROMIDE v. BROMO-ETHYLENE.
VINYL tribromide v. Tri-Bromo-ETHANE.
VINYL CHLORIDE v. CHLORO-ETHYLENE.
VINYL ETHANE v. BUTYLENE.
VINYL-ETHYL-CARBINOL v. PENTENYL

ALCOHOL. VINYL-ETHYLENE v. BUTINENE.

VINYL ETHYL OXIDE CH.:CH.OEt. (85.5° i.V.). S.G. 14.8 7625. Formed by heating chloro-acetal CH.2Cl.CH(OEt), with sodium at 140° (Wislicenus, A. 192, 106). Liquid, smelling like ether mixed with allyl compounds. Combines with Cl and Br, forming di-chloro- and dibromo-di-ethyl oxide. A small quantity (1g.) of iodine converts large quantities of the oxide (200 g.) into a viscid liquid. Dilute (1:4) H2SO4 forms aldehyde and EthSO4.

Reference.—CHLORO-VINYL-ETHYL OXIDE.

VINYL-ETHYL-PYRIDINE

C₃H₂NEt.CH:CH₂ (98°-102°at 21 mm.). Formed
by heating C₃H₃NEt.CH₂.CH₂OH with conc. HCl
at 170° (Prausnitz, B. 25, 2394). Oil, v. sol.
ether.—B'₂H₂Hg,Cl₁₂. Needles, sl. sol. hot water.
VINYL 10DIDE v. IODO-ETHYLENE.

VINYL-MALONIC ACID v. TRIMETHYLENE

DICARBOXYLIC ACID.

VINYL OXIDE (C₂H₃)₂O. (39°). Formed by the action of dry Ag₂O on vinyl sulphide (Semmler, A. 241, 90).

Reference.—HEXA-CHLORO-DI-VINYL OXIDE.

o-VINYL-PHENOL CH.: CH.C. H. OH. Methyl ether CH.: CH.C. H. OMe. o-Vinyl-anisole. (c. 198°). S.G. 15 10095, 30 10005. Formed from either of the methyl derivatives of o-oxy-phenyl-acrylic acid by successive treatment with HI and Na₂CO₂Aq (Perkin, C. J. 33, 211; 39, 429). Oil. Polymerises readily (at 150°), forming a glassy mass, which, however, on distillation, regenerates the original oil. It smells like high-boiling coal-tar naphtha. It forms a colourless compound with bromine. Forms a red solid with H₂SO₄.

p-Vinyl-phenol. Methylether CH₂:CH.C.H.O.Me. [3°]. (205°). S.G. ½5 1-0029; §3° 9956. Formed by distilling methoxyphenyl-acrylic acid (q.v.) and from the same acid by successive treatment with conc. HI (S.G. 1-94) and Na₂CO₃.

VINYL-

Reference.—Bromo- and Nitro-

PHENOL.

VINYL-PIPERIDINE $C_bH_{1a}(C_2H_3)N$ (?). (147°). Formed by dehydration of oxy-ethyl-piperidine [32°] (Ladenburg, B. 22, 2587). Liquid, smelling like tropidine and conline, v. solwater.

Reference.—Bromo-vinyl-piperidine.

p-VINYL-ISOPROPYL-BENZENE C₁₁H₁₄ i.e. C₆H₄Pr.CH:CH₂. (204°). S.G. ¹⁵ ·8902. Formed by distilling cumyl-acrylic acid at 210° or by boiling a-bromo-\$\text{B}\$-cumyl-propionic acid with Na₂CO₂Aq (Perkin, C. J. 1877, ii. 660). Oil, smelling like cuminic aldehyde. Partiall \$\text{J}\$ polymerises on boiling, and also on keeping, forming a glassy mass, reconverted into the original hydrocarbon by heat. Yields C₁₁H₁,Br₂ [71°].

a-VINYL-PYRIDINE C,H,N i.e.

NC(C₂H₂):CH>CH. (159°). S.G. 2 9985. Formed by passing a mixture of pyridine and

ethylene through a red-hot tube (Ladenburg, B, 20, 1643). Formed also by distilling oxy-ethylpyridine under high pressure or in presence of KOH (Ladenburg, B. 22, 2585), and by the action of NaOHAq on \$\beta\$-bromo-\$\beta\$-pyridyl-propionic acid (Einhorn, \$A\$. 265, 229). Liquid smelling like conyrine, m. sol. water, v. sol. alcohol. Decomposed by distillation under atmospheric pressure, but boils at \$1^\circ\$ under 29 mm. Oxidised by KMnO, to picolinic acid, and reduced in alcoholic solution by Na to ethyl-pyridine.—Salts: B'_H_PtCl_s. [174°]. Crystals, m. sol. water.—B'HAuCl_t. [144°]. Yellow needles.

Tetra-vinyl-pyridine C₅Vi(C₂H₅)₄N. (277°). S.G. 2 1-0515. Formed in the preparation of γ-ethyl-pyridine by heating pyridine ethyloidide in sealed tubes at 320° (Karan, B. 25, 2776). Sl. sol. water.—B'₂H₂PtCl₄. [175°].—B'HAuCl₄. [148°].—B'HHgCl . [146°]. Needles.

VINYL-QUINOLINE v. QUINOLYL-ETHYLENE. VINYL SULPHIDE C.H. S i.e. S(CH:CH₂)₂. (101°). S.G. 913. Constitutes the chief part of the essential oil of Allium ursinum (Semmler,

A. 241, 90). Liquid, smelling like allyl sulphide. Reactions.—1. Dry Ag₂O forms vinyl oxide.—2. Moist Ag₂O gives aldehyde.—3. Alcoholic HgCl₂ forms crystals of C₈H₁₂Cl₄Hg₂S₂, which, when heated with potassium sulphocyanide, yields vinyl sulphocyanide.—4. PtCl₄ added to its alcoholic solution ppts. (C₂H₃)₈Cl₂PtS₃, which is decomposed by ammonium sulphide into vinyl chloride and dark-brown (C₂H₃)₂PtS₃.—5. AgNO₃ forms (C₂H₃)₂SAgNO₃.—6. Br gives (C₂H₃Br₂)₂SBr₂ [195°].—7. Oxidising agents yield CO₂, oxalic acid, and H₂SO₄ only.

VINYL-TOLUIDINE so called is DI-p-TOLYL-DI-ETHYLENE-DIAMINE.

VIOLAQUERCITRIN v. this vol. p. 373. VIOLANTIN C_kH_eN_eO₉ 4aq. Formed by mixing hot conc. solutions of vitroso- and nitrobarbituric acids (violuric and dilituric acids). Formed also by warming hydurilic acid with dilute HNO₂ (Baeyer, A. 127, 223). Yellowishwhite, crystalline powder, decomposed by water into its two component acids, but may be recrystallised from HOAc or 50 p.c. alcohol. It is also split up into its components by salts of the stronger acids.

VIOLURIC ACID C.H.N.O. i.e.

C₂O₂<NH C:NOH. Nitroso-barbituric acid. Mol. w. 157.

Formation.—1. From hydurilic acid by the action of nitric acid (S.G. 1·2) or nitrous acid (Baeyer, A. 127, 200).—2. By heating dilituric (nitro-barbituric) acid with glycerin.—3. By warming ferrous diliturate with KCy.—4. By boiling an aqueous solution of alloxantin with hydroxylamine hydrochloride (Pellizzari, G. 17, 258).—5. By adding hydroxylamine to an aqueous solution of alloxan (Ceresole, B. 16, 1133).—6. By adding KNO₂ to barbituric acid (Baeyer, A. 130, 140).

Properties.—Trimetric crystals (containing aq); a:b:c=83:1:1.92. M. sol. cold water, sl. sol. alcohol. Its aqueous solution is ppd. by alcohol. FeSO, gives a deep indigo-blue colour. HNO, forms nitro-barbituric acid. Br forms di-bromo-barbituric acid and nitrous fumes. Beducing agents form uranil (àmido-barbituric

heating.

Salts .-- NH, A'. Dark-blue prisms .-- KA'2aq. Deep-blue crystals, v. sol. water, forming a blue solution; turned red by excess of KOH. solution of the K salt in conc. HClAq deposits (KA')2(HCl)3 6aq in colourless efflorescent prisms. -BaA', 4aq. Red dimetric tables, nearly insol. cold water.—MgA'2 6aq. Purple-red crystals.—PbA'2'4aq. Small red crystals.—AgA'.

Bensyl ether CO NH.CO C:NOC,H,. [226°]. Formed by the action of benzyl chloride on silver violurate (Conrad a. Guthzeit, B. 15, 2849). Silvery scales, sol. hot water and alcohol.

VIRIDIC ACID v. CAFFETANNIC ACID. **VIRIDINE** $C_{12}H_{19}N$. (251°). S.G. 1.024. A homologue of pyridine occurring in coal-tar (Thenius, C. C. 1862, 53). Yellowish oil, with slight greenish fluorescence, sl. sol. water, v. sol. alcohol and ether. — B'2H2PtCl6. Greenish brown, insol. water, alcohol, and ether. The mercuric chloride double salt melts at 85° and crystallises from water.

Isomeride C₁₂H₁₈N. (230°-235°). Obtained, with other bases, by heating methyl-ethyl-acroleïn with alcoholic ammonia (Hoppe-Seyler, M.

9, 651).—B'HAuCl. [93°].—B',H,PtCl. [135°].

VISCIN C₁₀H₂₁O₄ (?). The glutinous constituent of the stalk, leaves, and berries of the mistletoe (Viscum album). Extracted from the bark by kneading with water, washing the sticky mass with 90 p.c. alcohol, and extracting the viscin with cold ether. The residue consists of viscaoutchin and woody-fibre (Reinsch, C. C. 1861, 145). Colourless, tasteless, semi-fluid mass, S.G. 10, decomposed by distillation, yielding oily viscene (226°) S.G. 85, which forms a crystalline Na salt with conc. NaOHAq. Viscaoutchin is very glutinous. Its S.G. is

seid). Conc. HClAq forms hydroxylamine on | 978, and it is insol. alcohol and ether, sol. oil of turpentine.

VIŜCOSE is DEXTRANE (q.v.). VITELLIN v. PROTEÏDS. VITELLOSE v. PROTEÏDS.

VOLUMES, SPECIFIC: v. Specific volumes. p. 498.

VULPIC ACID C, H, O, i.e.

CPh:C(OH).C:CPh.CO,Me Methyl pulvate. CO-[148°]. Occurs in Cetraria vulpina, a lichen growing in Norway, and used there, mixed with nux vomica, as poison for wolves (Bebert, A. 2, 342; Strecker a. Möller, A. 113, 56; Spiegel, B. 13, The lichen con-1629; 14, 1686; A. 219, 15). tains 21 to 4 p.c. of the acid, which may be extracted by warm milk of lime. Vulpic acid is also formed by dissolving pulvic anhydride in a solution of KOH in MeOH. It appears to occur in the lichen Parmelia parietina (Berzelius; Stein, J. 1864, 553).

Properties.—Yellow plates or needles, sol. alcohol and ether, v. e. sol. chloroform, nearly insol. boiling water. Decomposed above 200° into MeOH and pulvic anhydride. Boiling milk of lime converts it into pulvic acid. Boiling KOHAq forms di-benzyl-glycollic acid and CO.

Salts .- NH, A'. Yellow crystals, sol. water. -BaA'₂2aq. Yellow needles (from water).-BaA'₂7aq.-KA'aq. Light-yellow needles, sl. sol. water.-AgA'. Pp. Blackens at 100°.

Acetyl derivative C10H12AcO4. [156°]. Colourless needles, insol. NaOHAq.

Methyl ether v. Di-methyl ether of Pulvic

Isovulpic acid C₁₈H₁₁MeO₅. [124°]. Formed, in small quantity, together with pulvic anhydride, by heating vulpic acid at 200° (Spiegel). Thin golden plates (from alcohol). Forms orange solutions in alkalis.

 ${f w}$

WACKENRODER'S SOLUTION. The solution obtained by passing H2S for a long time into nearly saturated SO₂Aq. The solution contains H₂S₄O₅, much H₂S₄O₆, H₂S₄O₆, and probably H₂S₆O₆, along with H₂SO₄, dissolved colloidal S, and a little S in suspension; v. THIONIC ACIDS, p. 698.

WALDIVIN C₁₈H₂₄O₁₀. [230°]. S.G. 1·46. S. 17 at 15°; 3 at 100°. S. (alcohol) 53. Extracted by dilute alcohol from the powdered fruit of Simaba waldinia (Tanret, Bl. [2] 35, 104; C. R. 91, 886). Hexagonal prisms (containing 5aq), v. sol. chloroform, insol. ether. Neutral to litmus. Inactive to light. Tastes bitter.

WATER. H₂O. (Hydrogen monoxide.) Mol. w. 17.96. (For physical data v. Properties.)

Occurrence. - Pure water is never found in nature. The properties of different specimens of naturally occurring waters depend on the impurities they contain, and these impurities are derived from the substances with which the water has come into contact; hence it is customary to classify natural waters in accordance with their origin, as rain-water, surface-water, well-water, mineral spring-water, and sea-water. The composition of the substances found in these waters, and the properties of the waters themselves-that is, of the various more or less dilute aqueous solutions - are discussed in pp. 939-960 of vol. iv. of the Dictionary of Applied Chemis-TRY. Solid water, more or less pure, is found as ice and snow. Water vapour is a constant constituent of the atmosphere. A great many minerals, and also many organic substances, contain water combined with other compounds.

Historical.—In 1781 Cavendish showed experimentally that water was the only product of burning H and O mixed in certain proportions. and that almost the whole of the H and O disappeared. The account of the experiments made by Cavendish was published in 1784 (T. 1784. 116).

When a mixture of inflammable and dephlogisticated when a mixture or inflammable and dephlogisticated air [4.e. in modern language, hydrogen and oxygen] is exploded in such proportions that the burnt air is not much phlogisticated, the condensed liquor contains a little acid, which is always of the nitrous kind...; but if it... proportions be such that the burnt air is almost entirely phlogisticated the condensed liquor to a calculate. phlogisticated, the condensed liquor is not at all acid, but

seems pure water, without any addition whatever; and as, when they are mixed in that proportion, very little air remains after the explosion, almost the whole being condensed, it follows that almost the whole of the infiammable and dephlogisticated air is converted into pure water' (Lc.

Translated into modern language, this statement would be taken as asserting that water is formed by exploding a mixture of H and O in proper proportions. It is, however, worthy of note that Cavendish did not himself interpret his experimental results as we interpret them to-day. He regarded 'dephlogisticated air' [oxygen] as 'nothing but dephlogisticated water, or water deprived of its phlogiston.' He said: 'We must allow . . . that inflammable air [hydrogen] is either pure phlogiston . . . or else water united to phlogiston' (l.c. pp. 137, 140).

The formation of water by burning H and O was thought of by Cavendish as the restoration of phlogiston to water that had been deprived of this principle. 'Water,' he said, 'consists of dephlogisticated air united to phlogiston. Cavendish evidently thought of H and O as what we might now call forms of water; one of these was water with too little phlogiston, and the other was water with too much phlogiston; the explosion restored the phlogistic balance, and the properties of water were apparent. When Lavoisier had interpreted Cavendish's results, Cavendish spoke of Lavoisier's explanation as an hypothesis: 'According to this hypothesis we must suppose that water consists of inflammable air united to dephlogisticated air' (l.c. p. 150). Cavendish established the fact that water is the product of burning a mixture of H and O in the ratio (approximately) of 2 vols. H to 1 vol. O; but he stated this fact in language that no longer carries a definite meaning with it. Lavoisier added to the experimental basis whereon the fact rested, and he expressed the fact in language that still is clear, definite, and descriptive.

Formation.-1. By the direct union of H and O by igniting a mixture of these elements. 2. By deoxidising metallic oxides, and many other compounds that contain O, by heating with H .- 3. By the decomposition of many compounds containing H and O, by heat, or by

reactions with other substances.

According to Freyer a. V. Meyer (B. 25, 622), a mixture of H and O in the ratio 2H:O does not explode when slowly passed through a glass tube at 606°, and the temperature of ignition of the wet, gaseous mixture is between 650° and 730°. Askenasy a. V. Meyer (A. 269, 49) found that when pure, dry electrolytic gas was passed at a moderate rate through a glass tube heated to 518°, only c. '7 to 1.7 mgms. of water were produced in ten hours; and that a little more water, but still only a very small quantity in proportion to the total quantity of H and O. was formed at 606°. Experiments made to determine the relation between the quantity of water formed and the time of the experiment showed that no constant relation could be arrived at, even when every precaution was taken to insure equality of conditions; the irregular action of the surfaces of the vessels was probably the cause of the irregularities in the rosults.

Davy (T. 1817) found that electrolytic gas did not explode when the pressure was so reduced

that the gas was rarefied to $\frac{1}{18}$ of its ordinary density. Thomas (C. J. 85, 215) found that the gas exploded at 168 mm. pressure. L. Meyer a. Scubert (C. J. 45, 586) found that the sparks from a Ruhmkorff coil caused the combination of c. 3 of a quantity of electrolytic gas at c. 70 mm. pressure, and thus the remainder combined when the pressure was increased until it became the same as before the first explosion; this result is in keeping with Bullsen's determinations of the quantity of oxygen needed to prevent the explosion of 2H + O (v. M. a. S., l.c. p. 588). Dixon (T. 1884. 634) noticed that elec trolytic gas did not explode at a pressure under 70 mm., but that explosion occurred under 75 mm. pressure (cf. D., l.c. p. 642).

According to the experiments of Dixon (l.c.), 'the union of oxygen and hydrogen is not affected by the presence or absence of water'; dry electrolytic gas exploded by the spark at a pressure between 70 mill. and 75 mm., and the

wet gas exploded at the same pressure.

The velocity of explosion of electrolytic gas was found by Berthelot and Vieille (C. R. 95, 151) to be 2,810 metres per second (cf. Explo-

sion, vol. ii. p. 530).

Preparation .- Stas (Chem. Proport. 110) prepared pure water as follows. When large quantities were required, spring-water was very slowly distilled through a long copper tube, bent into zigzag form, completely filled with pure copper turnings that had been oxidised by strongly heating in O, the copper tube being surrounded by alumina and sand, and heated to full redness; the distillate was then distilled in an apparatus of platinum.

The second method recommended by Stas, especially when comparatively small quantities of pure water are required, is based upon destroying the organic matter in distilled water by the action of K manganate and permanganate. The process is described by Stas as follows:-

I prepared potassium manganate by reacting on manganese oxide with caustic potash and potassium chlorate. I shook up the powdered product with water, just sufficient to dissolve the manganate that had been formed, and allowed the mixture to settle in a closed vessel. I then added 4 or 5 p.c. of the clear, dark-green solution to the spring water which was to be distilled, and allowed the components of this mixture to react for 24 hours. poured into the distillation vessel one or two litres of the cone, solution of potassium manganate that had been mixed with an equal volume of conc. caustic potash solution; this solution of potash was sufficiently conc. to make the salt so stable that its dilute solution could be heated for a salt so stable that its dilute solution could be heated for a long time without decomposition. I then filled the distillation vessel to a fa with the water which had been in contact with the potassium manganate, and distilled in the ordinary way. When boiling began I moderated the heat, in order to prevent the liquid, which frothed much for some minutes, from passing over. When the frothing has stopped, the water may be boiled rapidly without the least inconvenience. When fa of the water has distilled over, that which then distills is completely free from organic substances, and also from mineral substances, if the upper part of the distillation vessel is furnished with disphragms to hold back the extremely small drops that are always carried forward when a liquid is boiled vigorously.

Stassays that water thus prepared is perfectly free from organic matter. When he wished to obtain water absolutely free from any form of solid matter, Stas re-distilled the water that had been purified as described above, using as condenser a long tube of platinum soldered with gold. It is advisable to distil the water just before it is to be used.

On one occasion Stas used rain-water instead of well-water, and he found distinct quantities of ammonia in the distilled water thus prepared. To remove this he recommends to re-distil with

1000 part of NaHSO, or KHSO.

Composition of water.—The gravimetric composition of water was determined by Berzelius a. Dulong (A. Ch. [2] 15, 86) and by Dumas (A. Ch. [3] 8, 189) by passing pure H over a weighed quantity of red-hot CuO, and weighing the water produced and the copper which remained. The results gave the ratio H:O=1:8 (B. a. D.) and 1:7.98 (D.). An extended series of measurements by the same method, with many precautions, by Dittmar a. Henderson (C. N. 67, 127, 139, 151, 164 [1893]) gave the ratio H:O=1:7.9327.

Several measurements have been made of the proportion by volume in which H and O combine to form water. Gay-Lussac in 1805 and Humboldt in 1805 (A. Ch. 53, 239), found the volumetric ratio of H:O to be 2:1. Morley, in 1891 (Am. S. [3] 41, 220, 276), determined the ratio of H:O to be 2:00023:1 by directly measuring the volumes of the gascs. In 1892 Leduc (C. R. 116, 1248) found the ratio H:O = 2:0037:1, from determinations of the relative densities of H, O, and electrolytic gas. In 1893 Scott (T. 184, 543) completed a most carefully performed series of syntheses of water by sparking mixtures of H and O, and determined the most probable value of the volumetric ratio H:O to be 2:00245:1.

Properties. - Water is a clear, transparent, almost colourless, tasteless, odourless liquid. A column of water appears slightly blue when looked at lengthwise. Bunsen (A. 72, 44) pointed out that the slight blue colour of water may be observed by looking at a shining white object through a column of water 2 metres long, contained in a tube blackened inside. V. Meyer (B. 15, 297) recommends to join five wide, thinwalled glass tubes, c. 40 mm. internal diameter, and each c. 1\frac{1}{2} metres long, by wide caoutchouce tubing, and thus to form a tube c. 7\frac{1}{2} metres long; to lay the tube perfectly horizontal, and to close the end by smooth glass plates held in position by metallic clasps; then to cover the tube with black cloth. On looking through the tube the field of view appears quite colourless, but on now filling the tube with pure water (by means of brass tubes passing through the metallic clasps) a deep-blue colour is seen on looking through the column of water.

The boiling-point of water is 100° under the pressure of 760 mm. Zeuner (Grundsuge der mechanischen Wärmetheorie, Tab. 10 [1877]) gives the following table, showing the increase of boiling-point with increase of pressure:—

Pressure in atmos.	Boiling- point.	Pressure in atmos.	Boiling- point.
1 .	100	8	. 170.81
2.	120.6	9	. 175.77
З,	. 133.91	10	. 180.31
4,	144	11	. 184.50
5.	. 152-22	12	. 188.41
6.	159-22	13.	. 192.08
7.	165.34	14	. 195.53

An elaborate table is given by Broch (Trav. et Mém. du Burşau internat. des Poids et Mes.,

1, 46 [1881]) based on Regnault's determinations; the table gives the b.p. of water for each '1 mm. from 680 to 800 mm. pressure. (The table is given in Landolt a. Börnstein's *Physikalisch-Chemische Tabellen* [Berlin, 1883], pp. 47–49.)

The melting-point of ice is slightly lowered by pressure. J. Thomson (T. E. 16) calculated that the m.p. would be lowered by n-0075° for an increase of n atmospheres; W. Thomson (P. M. [3] 37, 123) confirmed this calculation by determining the m.p. of ice at 8·1 and 16·8 atmos. Mousson (A. Ch. [3] 56, 252) kept water liquid at -5° by greatly increasing pressure, and he found that at c. 15,000 atmos. pressure ice melted at -18° .

The specific gravity of water is greater at 4° than at any other temperature. Exner gives the temperature of maximum density as 3.945° (older determinations are tabulated by Exner, W. A. B. 68 (ii.), 463 [1873]). The following table, showing the density and volume of water from 0° to 100°, is given by Volkmann (W. 14, 260 [1881]); it is based on the determinations of Hagen, Matthiessen, Pierre, Kopp, and Jolly:

Temp. Density (in seaso) Le. wt. of 1 c.c. water in grams. Volume of 1 gram water in c.c.			
0 .999878 1.000122 1 .999933 1.000067 2 .999972 1.000028 8 .999993 1.000000 4 1.000000 1.000008 6 .999969 1.000081 7 .99933 1.000067 8 .999882 1.000118 9 .999819 1.000261 10 .999739 1.000261 11 .999650 1.000350 12 .999544 1.000466 13 .999430 1.000570 14 .999297 1.000466 13 .999154 1.000847 16 .99904 1.000947 17 .998839 1.001162 18 .998663 1.001839 19 .998475 1.001527 20 .998272 1.001731 21 .998065 1.001939 22 .997849 1.002156 23 .997623 1.002288	Temp.	Density (in vacuo) i.e. wt. of 1 c.c. water in grams.	Volume of 1 gram water in c.c.
0 .999878 1.000122 1 .999933 1.000067 2 .999972 1.000028 8 .999993 1.000000 4 1.000000 1.000008 6 .999969 1.000081 7 .99933 1.000067 8 .999882 1.000118 9 .999819 1.000261 10 .999739 1.000261 11 .999650 1.000350 12 .999544 1.000466 13 .999430 1.000570 14 .999297 1.000466 13 .999154 1.000847 16 .99904 1.000947 17 .998839 1.001162 18 .998663 1.001839 19 .998475 1.001527 20 .998272 1.001731 21 .998065 1.001939 22 .997849 1.002156 23 .997623 1.002288	_		
1		-999878	1.000122
2			
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65	55	·98584	1.01436
70	30	•98334	1.01694
75	65	·98071	1.01967
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Rossetti (P. Ergünzbd. 5, 268 [1871]) gives the densities and volumes of water for each degree from —10° to 100°, referred both to water at 0° and to water at 4° as unity. The S.G. of ice is c. '916 at 0° (water at 0°=1); according to recent determinations by Zakrzevski (W. 47, 155 [1893]) the value is '916660.

The expansion of water for various intervals of temperature has been measured by various observers; putting $V_t = V_0$ (1+at+bt²+ct³), the following values are given by Kopp (P. 72, 1 [1847]; cf. Pierre, P. 86, 451; Weidner, P. 123, 800; Matthiessen, P. M. [4] 31, 149; Rossetti, P. Ergánzbd. 5, 258; Hirn, A. Ch. [4] 10, 32):

For the expansion of water above 100° v.

Mendeléeff (A. 119, 1).

As water freezes it expands by c. $\frac{1}{11}$ of its volume; one volume of water at 0° becoming 1.09082 volumes of ice at 0°. It expands when heated at temperatures below 0°; Zakrzevski (W. 47, 155) gives the co-efficient of expansion -000077 (v. also Brunner, P. 64, 116; Struve, P. 66, 298; Marchand, J. pr. 35, 254).

The compressibility of water is small. Röntgen a. Schneider (W. 33, 644) give the absolute compressibility at 17.95° as '0000462 per atmosphere of pressure (v. also Ramsay a. Young, T. 1892; and cf. Grassi, A. Ch. [3] 31, 437; and Rankine, P. M. [4] 1, 548; also Amaury a. Descamps, C. R. 68, 1564; and Cailletet, C. R. 75, 77).

A table showing the volume of 1 kilo. of

saturated water vapour and the weight (in kilos.) of 1 c. metre of the vapour, at temperatures from 0° to 200°, is given by Zeuner (v. Landolt a. Börnstein's Physikalisch-chemische Tabellen [Berlin, 1883], p. 53; cf. Dieterici, W. 38, 1). According to Dieterici (l.c.) water vapour saturated at 0° behaves like a perfect

The vapour pressure of water varies from 1.0288 mm. at -19° to 20926.4 mm. at 230°; for complete tables calculated from Regnault's determinations v. Landolt a. Börnstein's Physikalisch-chemische Tabellen [Berlin, 1883] pp. 40-46 (the vapour pressure is given for each ·1° from -19°

 $\begin{array}{c} {}^{b} \\ {}^{c} \\ {}^{c$

to 101°, and for each 1° from 101° to 230°). Ramsay a. Young $(T.\ 1.62)$ give a table of the vapour pressures of water up to 270°. In connection with the vapour pressures of water and ice, v. R. a. V. $(T.\ 1884.\ 470)$. For an expression representing the vapour pressure of water at any temperature up to 325°, v. Antoine $(C.\ R.\ 113,\ 328)$.

The spec. heat of water increases as temperature rises; the quantity of heat required to raise 1 g. of water from t^o to t^o+1 is taken as unity in determinations of the spec. heats of other substances. The following table presents the data for S.H. of water at intervals of 10^o from 0^o to 230^o (the memoirs by the different observers are: Regnault, Acad. 21, 729 [1847]; Jamin a. Amaury, C. R. 70, 661 [1870]; Bosscha, P. Jubelbd. 549 [1874]; von Münchhausen, W. 1,

(air therm.)	Regnault	Jamin a. Amaury	Bosscha	v. Münchhausen	Henrichsen	Baumgartne
00	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000
10	1.0005	1.0111	1.0022	1.0048	1.0036	1.0031
20	1.0012	1.0225	1.0044	1.0085	1.0079	1.0061
30	1.0020	1.0341	1.0066	1.0128	1.0131	1.0092
40	1.0030	1.0459	1.0088	1.0170	1.0191	1.0123
50	1.0042	1.0580	1.0110	1.0213	1.0259	1.0154
-60	1.0056	1.0703	1.0132	1.0255	1.0335	1.0184
70	1.0072	1.0829	1.0154	1.0298	1.0419	1.0215
80	1.0090	1.0957	1.0176	1.0340	1.0511	1.0246
90	1.0109	1.1087	1.0198	1.0383	1.0612	1.0276
100	1.0130	1.1220	1.0220	1.0425	1.0720	1.0307
110	1.0158	1.1855	1.0242	1.0468	1.0837	1.0338
120	1.0177	1.1493	1.0264	1.0510	1.0961	1.0368
130	1.0204	1.1632	1.0286	1.0558	1.1094	1.0399
140	1.0232	1.1775	1.0308	1.0595	1.1235	1.0430
150	1.0262	1.1920	1.0330	1.0638	1.1384	1.0461
160	1.0294	1.2067	1.0352	1.0680	1.1540	1.0491
170	1.0328	1.2217	1.0374	1.0723	1.1706	1.0522
180	1.0364	1.2369	1.0396	1.0765	1.1879	1.0558
190	1.0401	1.2529	1.0418	1.0808	1.2060	1.0583
200	1.0440	1.2680	1.0440	1.0850	1.2249	1.0614
210	1.0481	1.2839	1.0462	1.0898	1.2447	1.0645
220	1.0524	1.8001	1.0484	1.0935	1.2652	1.0675
280c	1.0568	1.8165	1.0506	1.0978	1.2866	1.0706

592; 10, 284 [1877 and 1880]; Henrichsen, W. 8, 85 [1879]; Baumgartner, W. 8, 648 [1879]).

The following values for S.H. of water, from 0° to 35°, are given by Bartoli a. Stracciati (A. Ch. [6] 29, 285); the values in the column calculated were obtained by using the formula:

 $\begin{array}{lll} \text{S.H.} & = & 1 \cdot 006630 \\ & - & \cdot 000593962t \\ & + & \cdot 000004338650t^2 \\ & + & \cdot 00000425520t^3 \\ & - & \cdot 00000002819t^4 \end{array}$

The unit is the quantity of heat given out by 1 gram water at 15° in cooling to 14°.

t°	£.H. calod.	S.H. observed
0°	1.006630	1.00664
1	1.006041	1.00601
2	1.005463	1.00543
8	·004898	1.00489
4	1.004350	1.00435
5	1.003d20	1.00383
6	1.003307	1.00331
7	1.002824	1.00283
8	1.002362	1.00233
9	1.001927	1.00190
10	1.001522	1.00149
11	1.001146	1.00111
12	1.00080	1.00078
13	1.000496	1.00048
14	1.000224	1.00023
15	0.999990	1.00000
16	0.999795	0.99983
17	0.999642	0.99968
18	0.999530	0.99959
19	0.999462	0.99951
20	0.999439	0.99947
21	0.999463	0.99950
22	0.999533	0.99955
23	0.999652	0.99964
24	0.999821	0.99983
25	1.000040	1.00005
26	1.000311	1.00031
27	1.000633	1.00064
28	1.000967	1.00098
29	1.001488	1.00143
80	1.001921	1.00187
31	1.002459	1.00241
32	1.003054	
33	1.003668	
34	1.004408	-
85	1.005170	

The S.H. of ice is considerably less than that of water; Regnault's determinations gave '474 between -75° and 0°; the determinations of Person a. Desains gave '504 between -20° and 0°; Ramsay a. Young (T. 1884. 475) give the value '5 as the mean of various experiments. S.H. of water gas at 100° is given by Strecker (W. 17, 85) as '37 referred to an equal weight of water = 1, and 1'36 referred to an equal weight of water = 1; the ratio S.H.p. given by S. is 1'4, Jaeger (W. 36, 165 [1889] gave the value 1'33 to this ratio, and Cohen (W. 37, 628 [1889]) gave the value 1'287 for the temperature interval 144° to 300°.

The heat of vaporisation of water—i.e. the quantity of heat required to convert 1 g. of water at 100° into steam at 100°—is 535.77

gram-units, according to Favre a. Silbermann (A. Ch [3] 37, 461). The following values are given for the heat required to convert 1 g. of water at t° completely into water vapour; 606.5 when $t=0^{\circ}$, 637 when $t=100^{\circ}$, 676.6 when $t=230^{\circ}$ (Regnault, Acad. 21, 635); Dieterioi (W. 38, 1) gives 596.8 when $t=0^{\circ}$; Regnault (Lc.) gives the formula $\lambda=A+Bt$ for the total heat of vaporisation of water at different temperatures, and gives the values A=606.5, B=305.

According to Sakurai (C. J. 61, 495 [1892]), the temperature of the steam escaping from a boiling salt solution is exactly the same as that of the solution.

The heat of fusion of ice, i.e. the quantity of heat required to convert 1 g. of ice at t° into 1 g. of water at t° is given as follows by different observers (the values are in gram-units of heat):—79·24 and 79·06 when $t^{\circ} = 0^{\circ}$ (Regnault, A. Ch. [3] 8, 19); 79·25 when $t = 0^{\circ}$, 74·2 when $t = -10^{\circ}$, 80·02 when t varies from -2° to -21° (Person, A. Ch. [3] 21, 295; 30, 73); 77·85 at $-2\cdot8^{\circ}$, 76·75 at $-4\cdot995^{\circ}$, 76·11 at $-6\cdot28^{\circ}$, 76·0 at $-6\cdot5^{\circ}$ (Pettersson, J. pr. [2] 24, 129).

The thermal conductivity of water, from 10° to 18°, was determined by Winkelmann (P. 153, 481) to be '154; Bottomley (Pr. 31, 300) obtained nearly the same value; this figure means that heat sufficient to raise '154 mgm. water from 0° to 1° passes per second through a layer of water 1 mm. thick and 1 sq. mm. area, when the difference between the temperatures of the two surfaces of the layer is maintained at 1°. For electrical conductivity of water v. Reactions, No. 2.

The refractive indices of water at different temperatures, and for different lights of determinate wave-lengths, have been measured by many observers. The following values have been found for μ_{Ha} : 1·33120 at 19·9°, 1·33091 at 23·7°, 1·33050 at 26° (Brühl, B. 24, 644). For detailed tabulation of the results obtained by Fraünhofer, van der Willingen, Baille, Damien, Landolt, Wüllner, and Rühlmann, v. Landolt a. Börnstein's Physikalisch-Chemische Tabellen [Berlin, 1883] 205; cf. also Perkin (C. J. 61, 293), who gives values for μ for the lines A C D and F, at 15° and 83·7°.

Observations have been made on the absorption spectrum of water and water-gas, but the matter has not been thoroughly investigated (v. Vogel, P. 156, 326; Jansen, B. A. 1866. 11).

Water crystallises, as ice, in rhombohedral forms; snow is generally found crystallised in six-sided stars derived from six-sided prisms.

Molecular weight of liquid water. Several observations have been made which tend to show that the molecular weight of liquid water is greater than 18. Paterno (B. 21, 3180), from measurements of the lowering of the freezing-point of water dissolved in acetic acid, concluded that the mol. w. might be 18, or might perhaps be 36. From the depression of the freezing-point of paratoluidine by water dissolved therein, Eykman (Z. P. C. 4, 510) concluded that the mol. w. of liquid water is probably 36, and this conclusion was strengthened by Walker's experiments on the connection between heats of fusion and solubility (Z. P. C. 5, 194). From measurements of the surface tension of water Ramsay and Shields (C. J. 63, 1089 [1893]) con-

clude that the mol. w. of liquid water is probably 72 at the ordinary temperature.

Reactions. -1. In 1847 Grove (T. 1847. 1) showed that water was decomposed into H and O by heat. Grove formed a little ball on the end of a Pt wire, by fusing the Pt, heated the globule of Pt to whiteness by an electric current, and plunged it into a little air-free water, nearly boiling, in a small basin, with a test tube full of air-free water arranged to collect any gas that might come off. Deville (C. R. 56, 195, 322 [1863]) found that H and O were given off in considerable quantities when molten Pt was plunged under water. D. noticed no decomposition when steam was passed through a Pt tube heated to bright redness, but by passing a current of an indifferent gas, such as CO2, through the hot tube, and thus sweeping away the products of decomposition, H and O were obtained. 2. Water is scarcely decomposed by an electric current. Kohlrausch found the electrical conductivity of the purest water he could obtain by distillation in vacuo to be 2.5×10^{-6} in C.G.S. units, or c. 72 billionths of the conductivity of Hg (P. M. [5] 18, 542). By calculations based on this result, Ostwald concluded that in a litre of pure water the weight of water dissociated into H and OH ions, expressed in gram-molecules, is $\cdot 6 \times 10^{-6}$ (Z. P. C. 11, 521). By other methods of calculation, based on other data, Ostwald arrived at the value 2 to 9 × 10-6 for what has been called the dissociation constant of water the transfer that C is the constant of water $(Z.\ P.\ C.\ 11,\ 521)$; Wijs, by calculations based on the hydrolysis of methyl acetate, obtained the values $\cdot 1 \times 10^{-7}$ ($Z.\ P.\ C.\ 11,\ 492$), and $\cdot 14 \times 10^{-6}$ ($ibid.\ 12,\ 514$); Arrhenius obtained the value $\cdot 1125 \times 10^{-6}$ ($ibid.\ 11,\ 827$); and Bredig, the value $\cdot 6 \times 10^{-6}$ ($ibid.\ 11,\ 829$). Later experiments $\cdot 10^{-6}$ ($\cdot 10^{-6}$) ($\cdot 10^{-6}$). ments on the conductivity of water by Kohlrausch a. Heydweiller (Z. P. C. 14, 316 [1894]) with water that had been distilled in vacuo ten years ago, had then stood in a vessel filled with water, and been again distilled in vacuo, gave the following results:—conductivity (Hg = 1) $\cdot 014$ at 0°, $\cdot 04$ at 18°, $\cdot 058$ at 25°, $\cdot 089$ at 34°, $\cdot 176$; all these to be multiplied by 10⁻¹⁰. K. a. H. say that 1 mm. of this water at 0° had a resistance = that of 40 million kilometres of Cu wire of the same area. K. a. H. calculate that in 1 litre of the purest water at 18° there is 00008 mgm. H as free ions, and 000105 mgm. at 25°.-3. Steam is decomposed by electric sparks; for conditions and details of results v. Thomson (Pr. 53, 90).

The reactions of water are so many that an approximate classification of them into groups is all that can be attempted here.-4. Many metals react with water, at temperatures varying from the ordinary to a full red heat, forming oxides or hydroxides and giving off H. The following metals decompose cold water: Ba, Cs, Ca, (Ce?), (La?), Li, K, Rb, Na, Sr. Al, Fe, Pb, Mg, (? Mn), Mo, Ti react at c. 100°; and most of the other metals at temperatures from c. 100° to a full red heat.—5. Many non-metals react, generally slowly, with water, forming acids and giving cff O. F rapidly decomposes water at the ordinary temperature; Cl reacts slowly at the ordinary temperature, and somewhat more rapidly at a red heat; Br reacts more slowly than Cl; I has probably a very slight (? any) action. S and P react slowly at 100°; Se is said not to decompose water at 160°. Carbon gives off H at a red heat. Boron acts like a metal, giving off Hat a red heat .- 6. Many haloid compounds react with water, giving oxyhaloid compounds or oxides, and haloid acids.-7. Some metallic sulphides react with steam to form oxides and H.S. 8. A few of the lower oxides decompose water; e.g. CrO.xH2O at the ordinary comperature, and CO at c. 600°.-9. Water reacts with many oxides to form hydroxides which are either basic or acidic; in some cases hydrates are formed. Hydrates of various salts are also produced by combining the salts with water (v. Hydrates, vol. ii. p. 703; cf. Hydroxides, vol. ii. p. 733).-10. Water dissolves very many compounds of the most different properties (v. Solutions, vol. iv. p. 484).

Small quantities of water often bring about chemical changes that do not occur when the substances are perfectly dry; for instance, a mixture of dry CO and O is not exploded by sparks, but a trace of water suffices to start the change (v. Baker, C. J. 65, 611 [1894]).

The acidic and basic characters of water are so nearly brianced that the compound cannot be classed among either acids or basic bodies. The chemical relations of water to the compounds formed by reactions between it and other substances are determined chiefly by the chemical characters of the substances that react with it; thus the relations of HOH to $M_x(OH)_y$, formed by the interaction of water and metals, are those of an acid to its salts, whereas the relations of HOH to HX, formed by the interactions of water and non-metals, are those of a basic hydroxide to salts derived therefrom. The relations between both classes of derivatives of water and the parent compound are sometimes expressed by saying that the compounds belong to the water type (v. Types, vol. iv. p. 811). M. M. P. M. WAX. A term applied to various natural

WAX. A term applied to various natural solids more or less resembling bees'-wax. They are compound ethers, but differ from fats in yielding monovalent alcohols and not glycerin on saponification. They melt below 100°, are insol. water, sl. sol. or insol. alcohol, and sol. ether. They are not volatile.

Bees'-wax. [64°]. S.G. 965. Consists of a portion (about 5 p.c.) soluble in alcohol (cerin) and a portion insoluble in alcohol (myricin). Myricin is myricyl palmitate (Brodie, A. 67, 180; 71, 144). Cerin is chiefly composed of cerotic acid; but it contains small quantities of melissic acid $C_{so}H_{so}O_2$ [90°], and on acid melting at 73° (Schalfejeff, B. 9, 278, 1688; Nafzger, A. 224, 246). There is also present one or more acids whose lead salts dissolve in ether and whose Ba salts dissolve in alcohol, and which therefore probably belong to the cleic series. Myricin may be saponified by alcoholic potash, and the myricyl alcohol separated from potassium palmitate by extraction with ligroin. Crude myricin yields CCl, and C₆Cl₆ when heated with I and excess of SbCl₅ at 400° (Hartmann, B. 24, 1022). Bees'-wax contains two hydrocarbons, one of which [60°] (c. 275° at 11 mm.) is probably n-heptaicosane C₂,H₃₆, and the other [68°] (c. 300° at 11 mm.) n-hentriacontane C₁₁H_{c.} (Schwalb, A. 235, 106). According to Schwalt, the myricyl alcohol of bees'-wax has the formula. C₁₁H₆₄O [85°], and is converted by heating with.

soda-lime into an acid C₁₁H₄₅O₂ [89°], which forms a methyl ether [71°] and an ethyl ether [70°]. Among the products of saponification of bees'-wax, ceryl alcohol C₂, H₃₀O or C₂₆H₃₀O, and an alcohol C₂, H₃₂O or C₂₄H₃₀O occur. The last-mentioned alcohol when heated with soda-lime gives an acid $C_{25}\Gamma_{50}O_2$ or $C_{24}H_{48}O_2$ [75.5°].

Carnaüba wax v. vol. i. p. 710.

Chinese wax, which is produced by an insect, is almost entirely composed of ceryl cerotate (C₂₇H₃₅)C₂₇H₃₅O₂ (Brodie, A. 67, 199). Cork wax v. CERIN.

Pine wax v. CEROPIC ACID.

Sugar-cane wax v. Cerosin.

Japan wax. [42°-55°]. Obtained in the East from Rhus succedanea. It appears to be really a fat, since palmitin is its chief component It also contains the ether of a fatty acid of higher melting-point than stearic acid (Sthamer, A. 43, 343; Buri, Ar. Ph. [3] 14,

Wax of Ficus gummifius of Java contains an alcohol $C_{15}H_{30}O$ [73°], v. sol. ether, and a small quantity (5 p.c.) of isocervl alcohol $C_{27}H_{56}O$, sl. sol. ether (Kessel, B. 11, 2112).

Myrtle wax. [49°]. Got by boiling the berries of Myrica cerifera of North America with water (Moore, J. 1862, 506). Consists of palmitic and some lauric acid and (20 p.c. of) palmitin.

Opium wax. Contains ceryl cerotate and ceryl palmitate (Hesse, B. 3, 637).

Tobacco wax. Contains $C_{70}H_{140}O_2$ [63°], insol. cold alcohol, sol. ether and a small quantity of $C_{40}H_{120}O_2$ [64.5°], sl. sol. cold ether (Kissling, B. 16, 2433).

Coca leaf wax (Hesse, A. 271, 214). The wax from Trujillo coca is palmityl-(β)-amyrin $C_{46}H_{80}O_2$ [75°] [a]_D = 54.5°. On saponification it yields palmitic acid and (β)-amyrin C_{so}H_{so}O [196°] $[\alpha]_{\rm p} = 94 \cdot 2^{\circ}$, which yields an acetyl derivative [236°] and a benzoyl derivative [228°]. The wax from the broad-leaved coca of Peru and Bolivia melts at 70°; contains palmityl-(8)-amyrin and a ketone, (8)-cerotinone C₅₂H₁₀₆O [66°], m. sol. alcohol, ether, and ligroin. The wax from Java coca contains the same substances, and also some ceryl cerotate and ethers of myristic acid and of oxycerotic acid C27H34O3 [82°], which is v. e. sol. hot alcohol and ligroin, v. sl. sol. ether, and is converted by Aco at 100° into cerotolic acid C₂₇H₃₂O₂ [70°].

WHEY-PROIEÏD v. MILK.

WINE OIL. Light oil of wins. An oil obtained in the preparation of ether by distilling alcohol with H.SO. The ether is shaken with milk of lime and fractionally distilled, the successive fractions being ether, alcohol, and wet alcohol. Light oil of wine (.25 to .5 p.c. of the alcohol etherified) rises to the surface when the last fraction (90°-120°) is allowed to stand. When dried over CaCl₂ it has S.G. 17-5 903. It contains C. H. (157), EtOC, H. (112°), conessin. (q.v.).

Et.CO.C,H11 (154°), and CH3.CO.C,H12 (164°)

(Hartwig, J. pr. [2] 23, 449).

Heavy oil of wine, which passes over when the temperature is raised after the preparation of ether, consists of Et. SO, mixed with olefines (Claesson, J. pr. [2] 19, 259; Serullas, A. Ch. [2] 89, 152).

WINTERGREEN OIL contains methyl o-oxybenzoate

WINTERENE C₁₃H₂₄. (260°-265°.) S.G. 18:934. [a]₁ 11·2° at 16°. A dextrorotatory sesquiterpene obtained by distilling winterbark (from Drymis Winter-Forster) with water (Arata a. Canzoneri, G. 18, 527). Coloured green by Br in CHCl,

WOOD v. LIGNONE.

WOOD GUM v. XYLAN.

WOOD NAPHTHA v. METHYL ALCOHOL.

WOOD OIL. Gurjun Balsam. Flows from incisions in the stem of Dipterocarpus costatus. It contains an essential oil, which gives a splendid violet colour when its solution in CS, (20 pts.) is treated with a drop of a cold mixture of HNO. and H2SO4. The same reaction is exhibited by the balsam itself, and also, in a more transient manner, by cod-liver oil and copaïba balsam (Flückiger, Ph. [3] 7, 2). The essential oil consists chiefly of a terpene (255°) (Werner, J. 1862, 461), which composes 65 p.c. of the balsam (Guibourt, J. 1876, 907). The resin contains a neutral substance, $C_{2a}H_{4a}O_{3}$, which crystallises from light petroleum in triclinic prisms [126°— 130°]; it dissolves in conc. H2SO, and is reppd. from the resulting reddish solution by adding water. It is not affected by potash-fusion (Flückiger, Ar. Ph. [3] 12, 58). A substance $C_{20}H_{50}O_2$ [129°] is described by Brix $(M.\ 2,\ 516)$ as ppd. by adding water to an alcoholic extract of wood oil. It is neutral and insol. alkalis, and yields a diacetyl derivative [75°]. It is perhaps identical with the compound C_{.8}H_{.0}O_{.2} Wood oil also contains gurjunic acid C_{.2}H_{.4}O_{.4} [220°], which crystallises from alcohol, distils with decomposition at 260°, and forms Ag, A".

WOOD SPIRIT v. METHYL ALCOHOL.

WORMSEED OIL. Oleum Cina. tained by steam-distillation from wormseed, the flower-buds of Artemisia Vahliana, A. Sieberi, and A. inculta (Trommsdorff, Tr. N. J. 3, 812; Völckel, A. 38, 110; 87, 312; Hirzel, J. 1854, 591; 1855, 655; Kraut a. Wahlforss, A. 128, 293; Faust a. Homeyer, B. 7, 1429; Hell a. Ritter, B. 17, 2609; Wallach a. Brass, A. 225, 291). It consists chiefly of cineol C10H18O. Wormseed (A. Gallica) also contains betaine and choline (Jahns, B. 26, 1493).

WEIGHTINE C, H, N. [122°]. Occurs in the juice of Wrightia antidysenterica (Stenhouse, Ph. [2] 5, 493; Warnecke, B. 19, 60). Needles with bitter taste, sl. sol. water, v. sol. alcohol and ether. Conc. H,80, at 100° gives a dark-green colour, turned dark blue by adding water. Wrightine is probably identical with

EXAMPLE 1. A. H. Smith, Ph. [3] 23, 793). White crystalline powder, insol. water and alkalis, sl. sol. hot alcohol, m. sol. benzene, v. e. sol. chloroform. Weak base, forming yellow salts from which it is ppd. by hot water. H2SO4 forms a deep-orange solution, from which water ppts. the sulphate as yellow needles. Hot HNO, does not decompose it. Zinc and H2SO4 reduce it to hydroxanthaline C37H38N2O9, which forms white crystals [137°], nearly insol. water, v. sol. alcohol and benzene, and forms easily soluble crystalline salts. Hydroxanthaline is coloured deep-violet by H2SO4 (even free from HNO,), the colour being destroyed by water, but reproduced by H.SO4.-B'H2Cl24aq: needles. Gives off all its acid at 150°

XANTHAMIDE v. Ethyl ether of (β)-Thio-

CARBAMIC ACID.

XANTHATES. The salts RS.CS.OEt where

R is a metal; v. ETHYL DITHIOCARBONATE.

XANTHIC ACID v. ETHYL DITHIOCARBONATE. XANTHINE C, H, N, O, i.e.

NH.CH:C.NH CO.NH.C=N CO. Xanthic oxide. Mol. w. 152. S. .007 in the cold; .08 at 100°. Occasionally found in urinary calculi (Marcet, Essay on Calculi, London, 1819; Liebig a. Wöhler, A. 26, 340; Lebon, C. R. 78, 47) and in urinary deposits (Bence Jones, C. J. 15, 78). Occurs in small quantity in the urine of man, in the pancreas, spleen, and liver of oxen, in the thymus gland of the calf, in muscle of mammalia and fishes (Scherer, A. 107, 314; 112, 257; Städeler, A. 111, 28; 116, 102; Dürr, A. 184, 45; Kossel, H. 6, 422), in some kinds of guano (Unger a. Phipson, C. N. 6, 16), and in yeast (Schindler, H. 18, 432). Occurs also in Iupin seeds (Salomon, J. 1881, 1012; Schulze a. Barbieri, J. pr. [2] 27, 858), in pumpkin-seeds (E. Schulze, J. pr. [2] 32, 457), and in tea (Baginsky, H. 8, 396).

Formation.-1. By the action of nitrous acid on guanine (Strecker, A. 108, 141; 118, 151; 131, 121; Balke, J. pr. [2] 47, 542).—2. In small quantity by heating a mixture of HOAc and aqueous HCy (Gautier, Bl. [2] 42, 141).

Preparation.-1. Separates as a crystalline powder when NaNO₂ (8 g.) is added slowly to a solution of guanine (10 g.) in H₂SO₄ (20 g.) and water (150 g.) at 75° (Fischer, A. 215, 309).-2. The aqueous extract of sprouting lupin seeds is evaporated, the residue treated with alcohol, and the alcoholic filtrate evaporated. residue is dissolved in water and AgNO, and NH, added. The gelatinous pp. is dissc'ved in hot HNO, (S.G. 1.1), which yields on cooling a crystalline compound of hypoxanthine and silver nitrate, and on adding NH, to the filtrate silver-Lanthine is ppd. (S. a. B.).—3. Urine is ppd. with baryta-water; the filtrate is evaporated to a small bulk and boiled with cupric acetate. The pp. is dissolved in warm nitric acid and evaporated.-4. A solution containing xanthine is treated with Fehling's solution and hydroxylamine hydrochloride, and the ppd. copper compound decomposed by H_2S . The xanthine may be further purified by preparing its read salt and decomposing this with H₂S (Balke, J. pr. [2] 47, 552).

Properties.—Small scales (by evaporation) or powder composed of minute globules, nearly insol. cold water, insol. alcohol and ether. V. e. sol. KOHAq and reppd. by CO₂ and other acids. Weak base. Does not form an acetate. A cold saturated aqueous solution of xanthin, gives white pps. with HgCl2 and AgNO2, and a yellowishgreen, flocculent pp. with hot cupric acetate. An ammoniacal solution of xanthine is ppd. by HgCl2, ZnCl2, CdCl2, and AgNO3. Xanthine reduces ammoniacal cupric chloride (Drechsel, B. 25, 2454). Xanthine evaporated with nitric acid leaves a yellow residue turned orange by KOH (but not by NH_s), the colour becoming violet-red on warming. Solid xanthine added to a mixture of bleaching-powder and NaOHAq on a watch-glass forms a dark-green spot, changing to brown and finally disappearing. Xanthine warmed with chlorine-water and a trace of HNO, as long as gases escape, and then evaporated to dryness, yields a residue which is coloured rose-red by gaseous NH₂ (Weidel, A. 158, 365; Kossel, H. 6, 426).

Reactions.-1. Decomposed above 150°, giving off HCy, NH, cyanogen, and CO₂.—2. KClO₃ and HClAq at 60° form urea and alloxan.— 3. Conc. HClAq at 230° forms glycocoll, formic acid, NH,, and CO, (E. Schmidt, A. 217, 311).— 4. NaOH (2 mols.) and Pb/OAc), form a lead salt which, if dried and heated with MeI at 130°, yields theobromine (Fischer).-5. Slowly attacked by pure HNO, the gas evolved consisting of nitrogen (1 vol.), CO₂ (4 vols.), and N₂O (11 vols.) (Franchimont, R. T. C. 6, 223).
Salts.—B'HCl. Nodular groups of silky

needles.—B'H₂SO₄ aq. Scales, decomposed by water.—B'BaH₂O₂. Sl. sol. water.—B'₂Cu₄O₂. Formed by the action of Fehling's solution and hydroxylamine hydrochloride (Balke).-B'Ag₂O. Yellowish-white flocculent pp., got by adding AgNO, to an ammoniacal solution of xanthine. Blackens on boiling.—NaC, H, N, O, aq. Minute needles.

Bromo-xanthine C₅H₃BrN₄O₂. Formed by heating xanthine with bromine at 100°, and also by the action of nitrous acid on bromoguanine (Fischer a. Reese, A. 221, 343). Crystalline powder, sol. conc. HClAq and H2SO4, but reppd. by water; sl. sol. hot water and hot alcohol; sol. alkalis.

Isoxanthine C.H.N.O. i.e.

NH.CO.C.NH>N. Formed by reducing diazo-

i onitroso-methyl-uracil C_bH_aN_aO₄ with SnCl_a and HClAq (Behrend, A. 245, 223). Needles (containing laq), al. sol. hot water. Dissolves in Ac₂O without change. On evaporation with ppd. with AgNO,; the pp. is crystallised from (containing aq), sl. sol. hot water. Dissolves not diluted HNO, treated with ammoniacal in Ac.O without change. On evaporation with AgNO, decomposed by H₂S, and the solution HNO, it leaves a residue coloured orange by

KOHAq. Yields C, H, BrN, O, aq, crystallising

from water in six-sided tables.

Pseudo-xanthine C₅H₄N₄O₂. A product of the action of H₂SO₄ (2 pts.) on uric acid (1 pt.) at 120° (Schultzen a. Filehne, B. 1, 150). Formed also by the action of nitrous acid on adenine (Kossel, H. 10, 258). Powder, sl. sol. water, HClAq, and NH₃Aq, v. sol. KOHAq. Its aqueous solution is acid in reaction, and on evaporation with HNO leaves a lemon-yellow residue, which is turned orange on warming with KOHAq.

The name pseudo-xanthine is also given by Gautier (Bl. [2] 48, 19) to a substance C₄H₅N₅O occurring in musc lar tissue. This is a yellow powder, which forms a very soluble hydrochloride. Its aqueous solution is ppd. by HgCl₂, AgNC_a, and ammoniacal Pb(OAc)₂, but not by Pb(OAc)2. This nseudo-xanthine also gives an orange colour when the residue, after evaporation with HNO, is treated with potash.

Paraxanthine C,H,N,O, [c. 284°]. Occurs in human urine (Salomon, B. 16, 195; 18, 3406; H. 13, 187; Thudichum, H. 11, 415; Kossel, H. 13, 302). Silky needles or monoclinic tables, insol. alcohol and ether, sl. sol. cold water, v. sol. hot water. Sol. NH, Aq and HClAq. Poisonous, acting like caffeine and theobromine. AgNO. added to its solution in HNO, or NH, Aq gives a gelatinous or flocculent pp. Picric acid forms a yellow crystalline pp. when added to its solution in HClAq. Gives a red colour (like xanthine) when the residue after evaporation with chlorinewater is exposed to gaseous NH₂. Does not give an orange colour when KOHAq is added to the residue after evaporation with HNO. Conc. NaOHAq forms a crystalline salt. does the same. A solution of paraxanthine is ppd. by Cu(OAc), phosphotungstic acid, HgCl, and ammoniacal lead subacetate, but not by

mercuric nitrate.

Heteroxanthine C₆H₆N₄O₂. Occurs in urine of men and dogs (Salomon, B. 18, 3407; H. 11, 412). Amorphous powder, v. sl. sol. cold water, sol. NH, Aq, insol. alcohol and ether. Its solution in HClAq is not ppd. by pieric acid. AgNO, HgCl₂, Cu(OAc)₂, and ammoniacal lead sub-acetate give pps. When evaporated with chlorine-water and HNO₂ it leaves a residue which is coloured red by gaseous NH2, the colour changing to blue on addition of NaOHAq. NaOHAq forms a salt crystallising in tables, v. sol. water, sl. sol. NaOHAq. The hydrochloride forms sparingly soluble crystalline aggregates, which lose HCl on treatment with water.

Hypoxanthine v. vol. ii. p. 745.

XANTHININE C.H., N.O. S. 0025 in the cold; 025 at 100°. Formed by heating ammonium thionurate at 200° (Finck, A. 132, 298). Got also by heating pseudo-uric acid with H2SO4 at 150° (Grimaux, Bl. [2] 31, 535). White powder, nearly insol. water, sol. NH, Aq, forming a solution with blue fluorescence. Its solution gives a white pp. with HgCl, and a yellow pp. with AgNO. Not attacked by HNO. Sol. KOHAq and reppd. by CO₂. Sol. conc. H₂SO₄, forming a laminar sulphate, decomposed by water with separation of xanthinine. B'Ag2O. Bulky yellow pp., got by pouring an ammoniacal colution of xanthinine into excess of aqueous AgNO.

XANTHOCHELIDONIC ACID v. CHELIDONIC

XANTHOCREATININE v. CREATININE and LEUCOMAINES.

XANTHOGALLOL C18H4Br14O6 i.e.

C₆H_.Br₄O < 0.C₆HBr₅O > 0 (?). [122°]. Formed by adding pyrogallol (1 pt.) to bromine (10 pts.), leaving the mixture to stand for two hours, and then shaking with water and heating (Stenhouse a. Groves, C. J. 28, 1; A. 177, 191; 179, 237; Theurer, A. 245, 334). Tri-bromo-pyrogallol is an intermediate body in its preparation. Yellow laminæ (from CS₂ and ligroin), v. sol. ether and CS₂, m. sol. ligroin. Decomposed by boiling with water or alcohol. Very stable towards oxidising agents, even crystallising unaltered from conc. HNO. Not reduced by sodium-amalgam or by zinc and dilute H.SO. Aniline in HOAc yields an anilide [205°], to which Theurer assigns the impossible formula C18H4Br11(NHPh),O4. p-Toluidine forms a corresponding p-toluide. Reacts with phenyl-hydrazine acetate.

Reactions. -1. NaOHAq forms hexa-bromobenzene dihydride C_sH₂Br_s [139°], sl. sol. alcohol, which crystallises in prisms, while the motherliquor contains a sodium salt of an acid C₅H₄Br₂O₃ [124°], which forms BaA", crystallising from dilute alcohol in large white plates (Theurer). By the action of dilute NaOHAq on xanthogallol, Hantzsch a. Schniter (B. 20, 2033) obtained C₁₈H₄Br₁₁(OH)₂O₂, which yields Ba₄(C₁₄H₄Br₁₁O₂)₂ and crystalline C₁₈H₄Br₁₁Ac₂O₂. 2. Na₂CO₃Aq converts xanthogallol in the cold into $C_{18}H_{*}Br_{11}O_{b}$, crystallising from benzene in needles [72°] and prisms [131°], yielding the crystalline derivatives $C_{18}H_{*}Br_{11}O_{b}SNPhH_{*}$ and $C_{18}H_{*}Br_{11}O_{b}SNPhH_{*}$ and $C_{18}H_{*}Br_{11}O_{b}SNH_{*}C_{b}H_{*}Me.—3. HBr passed into$ a cooled solution of xanthogallol in MeOH forms C₁₈H₄Br₁₁O₃(OMe)_e, crystallising from MeOH in prisms [113°], and converted by boiling dilute NaOHAq into an acid C_sH₂Br₁O₂(OMe)₂ [105°], and by MeOH and hydrochloric acid into C. HBr. Cl(OMe), [77°]. The brominated acid C₀H₂Br₁O₂(OMe)₂ reacts with aniline, forming crystalline C₀H₂Br₁O₂(OMe)₂NH₂Ph. Conc. H₂SO₄ converts C₀H₂Br₄O₂(OMe)₂ into C₀H₂Br₄O₄ [65°].—4. Hydrochloric acid gas passed into a cooled solution of xanthogallol in MeOH forms C₁₈H₄Br₁₁Cl₂O₈(OMe)₆ [86°], which is insol. water, v. sol. alcohol, does not react with aniline, and is decomposed by dilute alkalis.—5. HCl passed into an alcoholic solution of xanthogallol forms C18H4Br11Cl2O2(OEt)8, crystallising from alcohol in colourless prisms [75°], converted by NaOHAq into a product [92°]. C18H4Br11Cl2Oe [104°] is a bye-product in the action of alcohol and HCl on xanthogallol. It forms large yellow crystals, v. sol. alcohol.

XANTHOGENIC ACID v. ETHYL DITHIOCAR-

XANTHOMETHYLIC ACID v. METHYL THIO-CARBONATES

XINTHONE is DIPHENTLENE RETONE OXIDE. XANTHOPURPURIN is m-DI-OXY-ANTHRA-

XANTHOQUINIC ACID p. Oxy-QuinoLine CARBOXYLIC ACID.

XANTHORHAMNIN C44H40029 (?). Obtained from Persian berries (the fruit of Rhamnus infactoria) by extracting with three times their

weight of 85 p.c. alcohol; the yield being 12 p.c. (Liebermann a. Hörmann, B. 11, 952, 1618; A. 196, 307; cf. Kane, P. M. [3] 23, 3; Gellatly, N. E. P. J. 7, 252; C. N. 8, 196; Hlasiwetz, A. 111, 108; Bolley, A. 115, 55; C. J. 13, 328; Stein, Z. [2] 5, 183, 568; Behrend, B. 11, 1353). Yellow needles (containing 2aq), v. sol. water and alcohol, insol. ether. Has little tinctorial power. Reduces Fehling's solution and ammoniacal AgNO, forming a mirror. FeCl, gives a dark-brown pp. Ppd. by ammoniacal lead acetate. Boiling dilute H.SO, splits it up into rhamnetin (2 mols.) and isodulcite (4 mols.).

Salts.—C₄₈H₄₂K₄O₂₈. Yellow powder, v. e. sol. water.—Pb₂A'v. Yellow pp.

derivatives Acetyl C48H84AC12O29 (Schützenberger, Z. [2] 4, 668).—C₄₈H₄₉Ac₁₇O₂₉. Powder, v. e. sol. alcohol (Liebermann a. Bergami, B. 20, 2245).

XANTHOROCCELIN v. Picro-roccelin.

EXANTHORRHEA RESIN is Acarold resin. **XANTHOXYLIN** $C_{10}H_{12}O_{1}$. [80°]. Occurs, together with xanthoxylene $C_{10}H_{10}$ (162°) in the essential oil from japan-pepper (Xanthoxylon piperitum) (Stenhouse, Ph. [2] 13, 423; 17, 19; A. 104, 237). Silky monoclinic crystals, insol. water, v. sol. alcohol and ether. Its alcoholic solution is not ppd. by AgNO, or lead acetate, even on addition of NH, Aq.

XENYLAMINE v. p-AMIDO-DIPHENYL.

KENYLENE-DIAMINE v. p-p-Di-AMIDO-DI-PHENYL.

XERONIC ACID v. DI-ETHYL-MALEÏC ACID. XYLAN $C_6H_{10}O_5$. Tree gum. Wood gum. S. 2 at 100° . [a]_b = $-69^\circ6^\circ$ (Tollens); -84° (Thomsen). Obtained from the bark of trees (T. Thomsen, J. pr. [2] 19, 146; Poumarède a. Figuier, A. 64, 388). Obtained, to the extent of 1.73 p.c., by extracting jute with dilute (5 p.c.) NaOHAq; and got also by extracting beechwood or pine-wood sawdust with 5 p.c. NaOHAq (Tollens, A. 254, 307, 320, 324, 326; Bl. [3] 1, 1102; Winterstein, H. 17, 381). Obtained also by extracting wheat-straw, first with 2 p.c. NH,Aq, and then with 5 p.c. NaOHAq (Tollens, A. 260, 291). The electric and HCl. The alkaline extract is ppd. with

Properties .- Porous mass, insol. cold, sol. hot, water; sol. NaOHAq. The hot aqueous solution becomes opalescent on cooling. Insol. alcohol, but the aqueous solution is not ppd. by alcohol unless an acid or the salt of an alkali is added. Insol. NH, Aq, lime- and baryta-water. Its aqueous solution is lævorotatory. furfuraldehyde when distilled with H2SO4 or HCl, and xylose when boiled with dilute H.SO. Gives no colour with iodine. HNO, oxidises

Gives no colour with iodine. HNO, exidises xylan to saccharic acid, but gives no mucic acid. o-XYLENE C_8H_{11} , i.e. $C_8H_{12}M_{01}[1:2]$. Dimethyl-benzene. Mol. w. 106. $[-28^\circ]$ (Colson, A. Ch. [6] 6, 128). (142°). S.G. § 8932. (Pinette, A. 248, 50); 24 876 (Gladstone, C. J. 59, 290). μ_A 14928, μ_B 1.5328. C.E. (0°-10°) 00098. S.V. 189°9 (Schiff); 137°6 (Finette). H.C. 1,084,274 [C,O₂=94,000; $H_{21}O=69,000]$ (Stohmann, J. pr. [2] 35, 41). M.M. 18-81 (Schönrock, Z. F. C. 11, 758. Critical temperature: 358° (Altschul. Z. P. C. 11, 590). Occurs ture: 358° (Altschul, Z. P. C. 11, 590). Occurs in coal-tar. The S.G. of crude xylone varies between 857 and 866; it contains 70 to 87 p.c. enic and xylyl m-xylene, 2 to 15 p.c. 10; 156, 236).

o-xylene, and 3 to 10 p.e. fatty hydrocarbons (Levinstein, B. 17, 444; cf. Fittig, A. 148, 10). If 100 c.c. of the mixture are boiled for 45 minutes with 40 c.c. of HNO₂ (S.G. 1.42) diluted with 60 c.c. of water, the p- and o-xylene are oxidised, leaving the m-xylene and fatty hydrocarbons. If the residual hydrocarbons, after washing with NaOHAc followed by steam distillation, be shaken for 80 minutes with 13 volumes of H2SO4, the m-xylene will be sulphonated and dissolved, while the fatty hydrocarbons remain. If 100 c.c. of crude xylene be shaken with 120 c.c. of H₂SO₄, the o- and mxylene dissolve, leaving the p-xylene and fatty hydrocarbons. On crystall sing the sodium salts of the dissolved sulphonic acids, sodium o-xylene sulphonate separates first, and may be converted into o-xylene by heating in a sealed tube with HClAq at 190° (Jacobsen, B. 10, 1009), or by heating with diluted sulphuric acid. Nölting, Witt, and Forel (B. 18, 2668) found 25 p.c. of p-xylene in commercial xylene. According to Nölting and Palmer (B. 24, 1955), crude xylene may contain 10 p.c. of ethyl-benzene. When a mixture of o-xylene and ethyl-benzene is treated with Br (20 pts.) and I, tetrs-bromo-xylene is formed, together with a less highly brominated ethyl-benzene (Crafts, C. R. 114, 1110).

Formation.—1. By distilling its carboxylic acids with lime.—2. From o-bromo-toluene, MeI, and Na (Jannasch a. Hübner, A. 170, 117; Reymann, Bl. [2] 26, 532). — 3. By heating cantharidin with P_2S_3 (Piccard, B. 12, 580).—4. By passing MeI through a mixture of toluene and AlCl, at 85° (Jacobsen, B. 14, 2628).

Properties.—Oil, solidifying in a freezingmixture at -28° . Unlike m- and p- xylene it does not yield a solid nitro- derivative with a cold mixture of H2SO4 and HNO3. It is completely oxidised by chromic acid mixture.

Reactions .- 1. Dilute HNO, forms o-toluic acid.—2. Boiling aqueous KMnO₄ oxidises it to phthalic acid.—3. PCl₅ at 200° reacts, forming C_oH₄(CCl₃).CHCl₂ (Colson a. Gautier, Bl. [2] 45, 507).-4. Bromine in the dark forms bromo-orylene C₆H₃Me₆Br [1:2:4]. In direct sunshine the products are C₆H₄Me₆CH₂Br and C₆H₄(CH₂Br)₂ (Schramm, B. 18, 1278).—5. AlCl₂ and gaseous HCl at 100° yield benzene, m- and a little pxylene, ψ-cumene and mesitylene (Hesse a. Töhl, A. 270, 168).

m-Xylene C₆H,Me₂ [1:3]. V.D. 3·68 (calc. 3·67) (Schiff, A. 220, 92). (139°). S.G. $\frac{9}{6}$ ·8812 (Pinette, A. 243, 50); $\frac{13\cdot 3}{4}$ ·8715 (S.); $\frac{9}{4}$ ·8655 (Brühl, A. 235, 13); 22.5 ·864 (Gladstone, C. J. (Brun, A. 25), 15); = $\frac{1}{100}$ (Gratistine, C. J. 59, 290). C.E. (0°-10°) 00096. $\mu_{\rm A}$ 1·495 (B.) $\mu_{\rm A}$ 1·4876, $\mu_{\rm B}$ 1·5277. S.V. 139·8. H.C. 1,084,168 (Stohmann, J. pr. [2] 35, 41). M.M. 12·73 at 20°-30° (Schönrock, Z. P. C. 11, 758). Critical temperature: 346° (A.). Occurs in coaltar (v. o-XYLENE).

Formation.—1. From m-iodo-toluene, MeL and Na (Wroblewsky, A. 192, 200).—2. From (1,8,4)-xylidine and amyl nitrite (Staedel a. Holz, Bt 18, 2919).—5. From toluene, McCi, and AlCl. (Friedel a. Crafts, A. Ch. [6] 1, 461; Ador a. Rilliet, B. 11, 1627).—4. By disti'ling mesitylenic and xylylic acids with lime (Fittig, A. 148, Properties.—Liquid. Not attacked by dilute HNO₂. Conc. HNO₂ on warming forms tri-

nitro-xylene [176°], sl. sol. alcohol.

Reactions.—1. Oxidised by chromic acid mixture to isophthalic acid.—2. PCl, at 200° forms a hexachloride (Colson a. Gautier, Bl. [2] 45, 508).-3. 3romine in the dark forms C.H.Me.Br [1:3:4], while in direct sunshine C.H.Me(CH₂Br) and C.H.(CH₂Br)₂ are produced (Schramm, 3. 18, 1277; M. 8, 805).—4. AlCl₂ and gaseous HCl at 100° form benzene, mesitylene, and some p-xylene and ψ -cumene (Heise a. Töhl, A. 270, 168). On boiling with AlCl, the products are benzene, toluene, a little p-xylene, ψ -cumene, mesitylche, and durene (Anschütz, A. 235, 182).—5. On heating with Mel and I at 250° it yields \u03c4-cumene, mesitylene, and C10H14 (Rayman . Preis, A. 223, 320).-6. Ethyl-malonyl chloride at 60° ir presence of AlCl, forms the ketone C₁₃H₁₄O₂ [63°] (Béhal a. Anger, Bl. [8] 8, 122).—7. Benzoyl per xide forms dixylylene C₁₆H₁₆ (265°), S.G. ²² ·9984 (Lippmann, M. 7, 528).—8. CrO₂Cl₂ added to its solution in CS₂ ppts. chocolate brown C₆H₄Me₂2CrO₂Cl₂, which is converted by water into m-toluic aldehyde and at 200° vields C_aH₄Me.CH(CrO₂Cl)₂ (Étard, A. Ch. [5] 22, 244).—9. CH₂Cl₂ and AlCl₄ form tetramethyl-anthracene [163°] (Friedel a. Crafts, A. Ch. [6] 11, 268).—10. PH₄I forms C₈H₁₄ on heating (Baeyer, Z. [2] 4, 455). HIAq and P at 280° form m-xylene hexahydride.

p-Xylene C₉H,Me₂ [1:4]. [13°] (Beissert, B. 23, 2242); [15°] (Jannasch). [138°). S.G. § :8601 (P.); ½? :860 (Gladstone, C. J. 59, 290). C.E. (0°-10°) ·00098 (Pinette, A. 243, 51). μ. 1·4854. μ. 1·5253. S.V. 140 (Schiff). H.C. 1,084,274 (Stohmann, J. pr. [2] 35, 41). M.M. 12·79 at 20°-30° (Schönrock, Z. P. C. 11, 753). Critical temperature: 344°. Occurs in coal-tar (v. supra) and in Galician petroleum (Pawlewski, B. 18, 1915). Formed by the action of MeI and sodium on p-bromo-toluene (Fittig, A. 136, 303; Jannasch, A. 171, 79) and on p-di-bromo-benzene (V. Meyer, B. 3, 753). Monoclinic prisms; a:b:c=2·32:1:2·34; β=69·5° (Baeyer, A. 245,

141).

Reactions.—1. Dilute HNO₃ forms p-toluic acid.—2. Chromic acid mixture yields terephthalic acid.—3. PCl₅ at 190° gives C₆H₄(CH₂Cl)₂ and at 200° C₆H₄(CCl₃)₂ (Colson a. Gautier, Bl. [2] 45, 6, 507).—4. Bromine in the dark forms C₆H₂BrMe₂, while in direct sunshine the products are C₆H₄Me.CH₂Br and C₆H₄(CH₂Br)₂ (Schramm, B. 18, 1276).—5. AlCl₂ and HCl at 100° act in the same way as with 0-xylene.

References.—Bromo-, Bromo-nitro-, Chloro-, Dichloro-nitro, Di-10Do-, Nitro-, and Oxx-

XYLENES.

XYLENE-AZO- compounds v. Azo- compounds.

XYLENE CARBOXYLIC ACID v. DI-METHYL-

DENZOIC ACID and MESITYLENIC ACID.

m-XYLENE DICARBOXYLIC ACID C₁₀H₁₀O₄
i.e. C₄H₂Me₂(CO₂H)₂ [1:3:4:6]. Dimethyl-iso-

phthalic acid.

(a)-Cumidic acid. Mol. w. 194. [above 320°]. Formed by heating di-bromo-m-kylene dissolved in ether under pressure with ClCo₂Et and sodium amalgam. Formed also, together with (β)-cumidic acid, by oxidation of durene with dilute HNO₂ or of durylic acid with KMnO₄ (Schnapauff,

B. 19, 2508). Minute prisms (from water) or plates (by sublimation).—Bah" 1\frac{1}{2}aq: crystals, v. sol. water. Yields m-xylene on distilling with lime.

Methylether Me.A". [76°]. Crystals.

m-Xylene dicarboxylic acid

 $C_sH_2M_{2_3}(CO_2H)_s$ [1:3:4:5]. Dimethyl-phthalic acid. Isocumidic acid. [280°]. Formed by oxidation of (β) - and (γ) -isodurylic acid (Jacobsen, B. 15, 1857). Small crystals. May be sublimed. Yields m-xylene on distillation with lime. The Ca salt crystallises in small plates. The Ba salt is amorphous.

p-Xylene dicarboxylic acid

 $C_cH_2Me_c(Co_aH)_2[1:4:2:5]$. Di-methyl-terephthalic acid. (\$\beta\$)-Cumidic acid. Formed, together with (\$\alpha\$)-cumidic acid, by oxidation of s-durene with dilute HNO₃ or of durylic acid with KMnO₄ (S.). Minute six-sided prisms (from alcohol). Sublimes in small plates without melting. Nearly insol. hot water. Yields p-xylene on distilling with lime.—BaA" $2\frac{1}{8}$ aq: tables, v. sol. water.

Methyl ether Me₂A". [114°]. (c. 297° cor.). Xylene dicarboxylic acid

by oxidation of C₆H₃Me(CO₂H)CH₂·CO₂H[S:2:1]. [178°]. Formed by oxidation of C₆H₃Me(CO.Et)CH₂·CO₂H (Béhal a. Auger, C. R. 109, 972). Gives off CO₂ on fusion.

Xylene di-ω-carboxylic acid v. Phenxiene-

DIACETIC ACID.

Reference.—Oxy-xylene carboxylic acid. XYLENE GLYCOL v. Di-oxy-xylene.

o-XYLENE DIHYDRIDE v. CANTHARENE.

m. Kylene dihydride C₅H₁₂. (133°). S.G. 22

·828. V.D. 3·74 (calc. 3·88). µ_D 1·4675. Formed, together with a polymeride C₁₆H₂₄ (280°-285°)
S.G. 2º ·832, by heating the ketone C₆H₁₄O with ZnCl₂. The ketone itself is obtained by distilling the anhydride C₁₀H₁₄O, which is got by the action of Ac₂O on cineolic acid, formed by oxidation of eucalyptol (Wallach, A. 258, 327). Liquid, smelling like xylene. Yields on nitration C₆H₄Me₂(NO₂) [1:3:4].

p-Xylene dihydride CMe CH. CH. CMe. (134°) at 720 mm. Formed by heating di-methyl-quinite dibromide C₈H₁₄Br₂ with quinoline (Baeyer, B. 25, 2122). Smells like turpentine. Forms a crystalline compound with HBr.

m-Xylene tetrahydride C.H.Me2. (119°).

S.G. 2 ·814; 14 ·794.

Formation.—1. By heating oxycamphoric anhydride C₁₀H₁₄O₄ with water at 180° or with HI at 150° (Wreden, A. 163, 386).—2. By heating camphoric acid with syrupy phosphoric acid at 200°.—3. By distilling camphoric acid with 2nCl₂ (Ballo, A. 197, 322).—4 By heating camphoric acid with HIAq at 200° (Wreden, A. 187, 171).

Properties.—Oil. Oxidised by chromic acid mixture to acetic, isotoluic, isophthalic, and terephthalic acids. HNO₃ forms tri-nitro-m-

xylene.

Xylene tetrahydride C_eH₁₄. (181°). S.G. 22 ·816. Occurs among the products of the distillation of colophony (Benard, A. Ch. [6] 1, 236). Slightly dextrorotatory liquid. Yields oily di- and tri- bromo- derivatives. Sulphuric acid forms an oily polymeride C_{1e}H₂₈, which is oxidised by fuming HNO₂ to succinic and oxalic acids. Absorbs oxygen. Does not reduce

ammoniacal AgNO₃. In ethereal solution it combines with Br, forming C₂H₁₄Br₂.

m-Xylene tetrahydride (?) C₂H₁₄. (120°).

Octonaphthylene. Formed by the action of moist Åg₂O on C₂H₁₄I, which is got by the action of CaI₂ at 60° on C₂H₁₅Cl (175°), a product of the chloriestics of extensible the Libertian of CaI₂ at 60° on C₃H₁₅Cl (175°). chlorination of octonaphthene (Jakowkin, J. R. 16, ii. 294). Oil, smelling like turpentine. Combines with Br (2 atoms). An isomeric octonaphthylene (123°-129°) is got by distilling C_sH₁₅Cl obtained by chlorination of iso-octonaphthene (Putochin, J. R. 16, ii. 295).

m-Xylene hexahydride CaH1s i.c.

C.H., Mo. [1:3]. Octonaphthene. (118°). S.G. § '7814 (Lossen, A. 225, 110); 2 '7706 (A.). C.E. (0°-26°) '001072. V.D. 3 '87. S.V. 164 '8. $\mu_{\rm D} = 1.419$ at 22° (Wallach, B. 25, 923). Occurs in Baku petroleum (Beilstein a. Kurbatoff, B. 13, 1820; Markownikoff a. Spady, B. 20, 1850), and among the products of the distillation of colophony (Renard, A. Ch. [6] 1, 229). Formed by heating heptanaphthene carboxylic acid with HIAq and P (Aschan, B. 24, 2718). Prepared by heating camphoric acid or m-xylene with HIAq at 200° (Wreden, A. 187, 157). Liquid. Not oxidised by aqueous KMnO₄. H₂SO₄ and HNO, form tri-nitro-m-xylene. Sulphur at 220° forms m-xylene.

Isomeride C, H16. Iso-octonaphthene. (122°). 8.G. 8 ·7767 17.8 ·7637. Occurs in Caucasian petroleum (Putochin, J. R. 16, ii. 295). Liquid.

p-Xylene hexahydride $C_eH_{10}Me_2[1:4]$. (138° cor.). V.D. 4.01 (obs.). S.G. 4.7956. Formed by heating bromo-camphor with ZnCl, (R. Schiff, B. 13, 1407; G. 10, 320). Liquid, yielding trinitro-p-xylene [127°] on nitration.

m-XYLENE PHOSPHINIC ACID

Chloride C.H.Me.PCl2 [1:3:4]. (257°). Got from mercuric dixylyl Hg(C₆H₄Me₂ [4:1:3]/₂ and PCl₄ at 235° (Weller, B. 20, 1720). Liquid, Liquid, converted by water into the acid C₆H₃Me₂P(OH)₂. Chlorine forms the compound C₆H₂Me₂PCl₄, which is converted by water into C₆H₃Me₂PO(OH)₂ [194°].

m-Xylene-phosphinic acid C_eH₂Me₂P(OH)₂ [98°] (Michaelis, A. 212, 237). Formed by the action of water on the chloride. Needles (from

alcohol). Probably a mixture.

Chloride C.H.Me.PCl. (256°). S.G. 19 1.24. Formed from m-xylene, PCl₃, and AlCl₃ (Weller, B. 20, 1720). It is a mixture of the preceding body and the 1,3,5 isomeride. Readily absorbs chlorine, forming C₀H₂Me₂PCl₄, whence SO₂ forms C₀H₂Me₂POCl₂ (280°-300°).

p-Xylene phosphinic acid C.H.Me.P(OH)2. p-Xylene phosphinous acid. Formed by treating the chloride with hot water (Weller, B. 21,

1494). Crystallises with difficulty.

Chloride C.H.Me.PCl. [-30°]. (254°).

S.G. 18 1-25. Formed by heating p-xylene (150 g.) with AlCl, (30 g.) and PCl, (200 g.) for 36 hours with inverted condenser (Michaelis a. Paneck, A. 212, 236). Strongly reflacting liquid.

m-XYLENE i-PHOSPHONIC ACID

C.H.Me. PO(OH). m-Xylene (a) phosphinic acid. [194]. S. 1.5 at 20°; 6.9 at 100°. Formed by boiling the chloride C.H.Me.PCl. (v. supra) with water (Weller, B. 20, 1721). Needles, v. e. sol. alcohol. Decomposed into m-xylene

and phosphoric acid on heating with alkalis. Br added to dilute solutions forms bromo- and di-bromo-m-xylene. Yields two nitro-derivatives [182°] and [100°].

Salts. — BaA" aq. Plates. — CdH₂A"₂ aq. Plates, more sol. cold than hot water. — NiH₂A"₂ aq.—Ag₂A"; white pp.

m-Xylene s-phosphonic acid

C_eH_aMe₂·PO(OH)₂. m-Xylene (β)-phosphinic acid. [161°]. S. 1·8 at 15°; 117·3 at 100°. Prepared by boiling with water the mixture of chlorides C_sH_sMe₂.POCl₂ (280°-300°) which is got by the action of SO₂ on the mixture of tetrachlorides C.H.Me.PCl. formed by combination of chlorine with the mixture of chlorides C.H.Me.PCl. obtained by heating m-xylene with PCl, and AlCl, (W.). Plates or needles, v. e. sol. alcohol and ether. Decomposed by lkalis into m-xylene and phosphoric acid.

p-Xylene phosphonic acid C₆H₃Me₂.PO(OH)₂. [180°]. Formed by the action of water on C₆H₂Me₂.PCl₄ [c. 60°] which is got by passing Cl through the product of the action of PCl₂ on pxylene in presence of AlCl, (Weller, B. 21, 1494). Needles, m. sol. water, v. sol. alcohol, sl. sol. Conc. HHO, forms the compound C.H.(NO.)Me.PO(OH). [224°]. KMnO. yields C.H.Me(OO.H).PO.H. [278°]. Salts.—KHA".—BaA". Pearly plates, more KMnO, yields

sol. cold than hot water.

Chloride C₆H₈Me₂.POCl₂. (281°). S.G. 18 1.31. Oil. Formed by the action of SO₂ on

C4H,Me,PCl4.

Reference.—Nitro-xylene phosphonic acid.

XYLENE PHTHALOYLIC ACID v. Phenyl.

XYLYL KETONE CARBOXYLIC ACID.

o-XYLENE SULPHINIC ACID C,H10SO2 i.e. C_cH₂Me₂·SO₂H [1:2:4]. [83°]. Formed by the action of zinc-dust on o-xylene sulphochloride (Jacobsen, B. 10, 1011). Plates (from water). m-Xylene sulphinic acid $C_eH_2Me_2(SO_2H)$

[1:3:4]. [c. 50°]. Formed by reducing the sulphonic chloride C_eH₃Me₂.SO₂Cl (Jacobsen;

cf. Lindoff a. Otto, A. 146, 233). Crystalline. p-Xylene sulphinic acid C₆H₂Me₂(SO₂H) [1:4:2]. [85°]. B. 11, 22). Groups of needles (Jacobsen,

o-XYLENE SULPHONIC ACID C.H. SO. i.e. C.H.Me. (SO.H) [1:2:4]. Formed by warming o-xylene with H.SO. (Jacobsen, B. 10, 1011; 11, 22). Tables (containing 2aq) (from dilute H₂SO₄).—NaA' 5aq.—BaA' 2aq. S. 5·8 at 0°; 83.6 at 100°.

Chloride C.H.Me.SO.Cl. [52°]. Prisms. Amide C.H.Me.SO.NH. [144°]. Yields Yields two acids C.H.Me(CO.H).SO.NH, on oxidation by KMnO4.

o-Xylene c-sulphonic acid C.H.Me.SO.H. [1:2:3]. Formed by the action of sodium-amalgam on $C_0H_2Me_2Cl(SO_2H)$ [1:2:6:3] (Krüger, B_1 18, 1760)

Amide C.H.Me. SO.NH. Granules.

m-Xylene c-sulphonic acid C.H.Me.(SO.H) [1:3:2]. m-Xylene (B)-sulphonic acid. Formed, together with a larger quantity of the (1,3,4)isomeride, by dissolving m-xylene in fuming H(SO, (Jacobsen, A. 184, 188; B. 11, 18). It remains in the acid liquor after ppg. the (1,3,4)isomeride by the proper quantity of water. Potash-fusion yields p-xylenol [74.5°] (Jacobsen, B. 21, 2828). Its K salt heated with sodium formate forms an acid C₉H₁₀O₂ [99°].—KA'.—

BaA'₂: minute needles.—CuA'₂aq.

Chloride C₆H₂Me₂.SO₂Cl. Oil.

Amide C₆H₂Me₂.SO₂NH₂. [96°]. Needles. m-Xylene sulphonic acid CaH3Me2(SO3H) [1:3:4]. m-Xylene (a)-sulphonic acid. Formed as above. Long flat prisms (containing 2aq). Its K salt heated with sodium formate yields xylylic acid.—NaA'.—BaA'. Small plates.— ZnA', 9aq. - CuA', 6aq. Chloride C,H,Me, SO,Cl. [34°].

Amide [137°]. Needles (from water). BzCl at 160° forms C.H.Me.SONHBz [151°] (Mahon, Am. 4, 194), which yields crystalline $Ca(C_{15}H_{14}NSO_3)_2$ aq and BaA'_2 , sl. sol. water.

p-Xylene sulphonic acid CaHaMe. (SOaH) [1:4:2]. Formed by sulphonating p-xylene (Beilstein a. Wahlforss, A. 133, 38; Fittig a. Glinzer, A. 136, 305; Jacobsen, B. 10, 1009; 11, 22; Remsen a. Emerson, Am. 8, 265). Plates or prisms (containing 2.4q). Oxidised by alkaline KMnO, to sulpho-terephthalic and two sulpho-ptoluic scids.—NaA'aq. [148°]. Trimetric plates; a:b:c=1.077:1:1488 (Moody a. Nicholson, C. J. 57, 978).—KA'aq.—BaA'₂. S. 2·27 at 0°; 5·53 at 100°.—CuA', 8aq. Light blue triclinic

prisms.—ZnA'₂10aq.

Chloride C_eH₂Me₂·SO₂Cl. [26°]. Prisms.

A mide [148°]. Needles, m. sol. hot water. o-Xylene disulphonic acid C6H2Me2(SO3H)2. Formed by heating (1,2,4)-xylene sulphonic acid with ClSO₃H at 150° (Pfannenstill, J. pr. [2] 46, 155).—K₂A"aq.—BaA"3aq.—PbA"3aq. Prisms. Chloride C₈H₂Me₂(SO₂Ol)₂. [79°]. Prisms. Amide C₈H₂Me₂(SO₂NH₂)₂. [289°].

m-Xylene disulphonic acid C.H.Me2(SO:H)2 [1:3:4:2]. Formed by heating m-xylene with fuming H.SO, at 150° (Wischin, B. 23, 3113), and by heating (1,3,4)-xylene sulphonic acid with SO₂ at 150° (P.). Deliquescent needles, converted by po'ash-fusion into C₄H₂Me₂(OH)₂ [146°].—(NH₂)A".—NaHA" 3aq.—KA" 2aq.— BaA" 8aq. Needles, v. sol. water. -PbA" 8aq.-CuA". Gummy mass.

Ethyl ether Et.A". Plates. Chloride C.H.Me.(SO.Cl). [129°]. Converted by PCl, at 180° (2,4,1,8)-di-chloro-xylene (220°).

[249°]. Amide C₈H₁₂N₂S₂O₄. Needles. Oxidised by KMnO₄ to $C_6H_2(<_{SO}^{CO}>NH)$ [225°].

Ethylamide C.H.Me. (SO2NHEt)2. [135°]. m-Xylene disulphonic acid C₆H₂Me₂(SO₃H)₂ [1:3:2:6?]. Formed in small quantity, together with the preceding isomeride, by heating (1,3,2)xylene sulphonic acid with ClSO,H at 150° (Pfannenstill, J. pr. [2] 46, 154). Needles. Chloride C.H.Me.(SO₂Cl)₂. Oil.

Amids. [210°]. More sol. water than its isomeride.

p-Xylene disulphonic acid C₆H₂Me₂(SO₄H), 4:2:6?].* Formed by heating (1,4,2) [1:4:2:6?].* [1:4:2:67]. Formed by neating (1,4:2)

C_eH₄Me₂(SO₂Cl) with fuming H₂SO₄ (Holmes,

Am. 18, 372; Pfannenstill, J. pr. [2] 46, 156).

Needles, v. e. sol. water.—BaA" 3aq.—CaA" 4aq.

—MgA" 7aq.—PbA" 3aq.—Ag₂L"aq.

Chloride C_eH₂Me₂(SO₂Cl)₂. [75°].

Amide C_eH₂Me₃(SO₂NH₂)₂. [295°].

References.—Bromo., Bromo.NITRO., NITRO.

said NITRO-AMIDO-, XYLENE SULPHONIC ACID.

XYLENE THIOSULPHONIC ACID. Xylyl ether C.H. SO, S.C.H. Xylyl disulphoxide. Formed by heating xylene sulphinic acid with water at 150°-160° (Otto a. Lindoff, A. 146, 239). Oil, v. sol. alcohol and ether.

XYLENIC ALCOHOL is TOLYL-CARBINOL.

XYLENIC DIBROMIDE v. DI-w-BROMO-XYLENE.

XYLENIC GLYCOL v. DI-W-OXY-XYLENE.

o-XYLENOL C₅H₁₆O i.e. C₅H₂Me₂·OH [1:2:4]. Mol. w. 122. [62°]. (225° i.V.). H.F.p. 61,566 (Stohmann, J. pr. [2] 34, 316). Obtained by potash-fusion from the sulphonic acid (Jacobsen, B. 11, 28), and by the diazo-reaction from (1,2,4). xylidine (Jacobsen, B. 17, 161). Long needles (from water). SiCl, yields Si(OC₈H₈), crystallising in prisms (350°-360° at 120 mm.) (Hertkorn, B. 18, 1691). Ammoniacal ZnBr₂ at 300° from xylidine and dixylylamine.—NaOC₂H₂. Flat needles, v. sl. sol. NaOHAq.

Benzoyl derivative [57°]. H.F. 77,768

(Stohmann, J. pr. [2] 36, 8). o-Xylenol C₆H₂Me₂(OH) [1:2:8]. [75°]. (218° i.V.). Occurs in coal-tar (Schulze, B. 20, Formed by the action of nitrous acid on c-o-xylidine (Töhl, B. 18, 2562; Nölting a. Forel, B. 18, 2673). Slender needles. Its aqueous solution gives a blue colour with FeCl,

c-m-Xylenol C₆H₃Me₂(OH) [1:3:2]. [49°]. Obtained from the crude xylidine, or by the action of HCl on p-oxy-mesitylenic acid (Nölting; Jacobsen, B. 21, 2829). p-Xylenol is got instead of m-xylenol by potash-fusion from (1,3,2). xylene sulphonic acid (Jacobsen, B. 11, 26; 21, 2828).

m-Kylenol C_cH,Me_c(OH) [1:3:4]. [26°] (Jacobsen, B. 18, 3463); [28°] (Staedel a. Hölz, B. 18, 2919). (211·5° i.V.). S.G. § 1·0362. H.F.p. 59,501 (Stohmann, J. pr. [2] 34, 316).

Formation.—1. From m-xylene sulphonic acid by potash-fusion (Jacobsen, B. 11, 24, 375; cf. Wroblewsky, Bl. [2] 10, 286; Wurtz, J. 1868, 459).—2. By heating oxy-mesitylenic acid with conc. HClAq at 200° (Jacobsen, B. 11, 2052).-3. By reduction of nitro-m-xylene and treatment of the resulting xylidine with nitrous acid Harmsen, B. 13, 1558; S. a. S.; Hodgkinson a. Limpach, C. J. 63, 104).

Properties .- Needles, v. sl. sol. water, Volatile with miscible with alcohol and ether. steam. FeCl, colours its aqueous solution blue and its alcoholic solution green, being changed to bright blue on dilution with water. Yields the corresponding xylidine and di-xylyl-amine when heated with ZnBr, ammonia, and NH, Br at 315° (Müller, B. 20, 1041). Potash-fusion forms (4,1,3)-oxy-toluic acid. Yields oily P)(OC,H,), (Kreysler, B. 18, 1703). SiCl, forms S(OC,H,), (455°) (Hertkorn, B. 18, 1690). Salt.—NaOC,H, V. sol. water and conc.

NaCHAq (unlike Na salts of other xylenols).

Methyl ether MeOC.H. (192° i.V.) (J.); (186°) (S.). H.F.p. 46,336 (Stohmann, J. pr. [2] 35, 24).

Acetyl derivative C.H.OAc. (226° i.V.). s-Xylenol C₆H₈Me₂(OH) [1:3:5]. [64°] (T.); [68°] (N. a. F.) (220°). Occurs in coal-tar (Schulze, B. 20, 410). Formed by the action of nitrous acid on s-xylidine (Thöl, B. 18, 362; Nöl'ing a. Forel, B. 18, 2679). Slender needles (from 872 XYLENOL.

water). Not coloured by FeCl.-NaOC.H.

Plates, sl. sol. NaOHAq.

p-Xylenol C_sH_sMe₂(OH) [1:4:2]. [74·5°]. (211·5° i.V.). H.F.p. 61,862 (Stohmann, J. pr. [2] 84, 316). S.G. 21·971. Formed by potashtusion from m- and from p-xylene sulphonic acid and from oxy-mesitylenic acid (Wurtz, A. 147, 378; Jacobsen, B. 11, 26). Formed also from p-xylidine by diazotisation followed by boiling with water (Nölting, Witt, a. Forel, B. 18, 2665). Long flat needles (from dilute alcohol). Volatile with steam. Its aqueous solution is not coloured by FeCl₂.

Potash-fusion gives (3,4,1)-oxy-toluic and oxy-terephthalic acids. KNO₂ and HOAc yield a nitroso- derivative $C_6H_2Me_2(OH)(NO)[1:4:3:6]$ [163°], which is the oxim of p-xyloquinone (Oliveri, G. 12, 162; Goldschmidt a. Schmid,

B. 18, 568).

Salt.—NaOC, H,. Large plates, al. sol. cold

NaOHAq.

Methyl ether MeOC₂H₂. (194° i.V.). Oil. Ethyl ether EtOC₂H₃. (199°) (S.); (205°) (N. W. a. F.). Oil. H.F.p. 54,150.

Acetyl derivative C,H,OAc. (237° i.V.).

S.G. 1 1.0264. Oil, not solid at -20°.

Xylenol. (220°). Occurs in beechwood-tar (Marasse, A. 152, 75; Tiemann a. Mendelssohn, B. 10, 57). Oil. Yields C.H.OMe (220°). On boiling with aqueous KMnO4 it yields an acid which is converted by potash-fusion into u-oxyisophthalic acid [300°].

References.—Bromo- and Nitro- xylenols.
XYLENOLIC ACID v. Oxy-DI-METHYL-BEN-

ZOIC ACID

o-XYLENOL SULPHONIC ACID C₅H₁₀SO₄ i.e. C₆H₂Me₂(OH).SO₅H. Formed by sulphonation of (1,2,4)-xylenol (Jacobsen, B. 11, 24). FeCl₃ gives a violet colour in neutral solutions.—NaA'. Slender prisms or small laminæ.—BaA'₂. Nodular groups of minute laminæ, sl. sol. cold water.

m-Xylenol (a)-sulphonic acid. Formed, together with the (β)-isomeride, by dissolving 1,3,4)-xylenol in H₂SO₄ (J.). FeCl₃ gives a violetblue colour.—NaA'. Tables.—KA'. Plates.—

BaA'₂. Leaflets. m-Xylenol (β)-sulphonic acid

C₆H₂Me_c(OH).SO₂H [1:3:4:2] (Jacobsen, A. 195, 283). Formed as above. Coloured violet-blue by FeCl₂. Converted by potash-fusion into (2,5,1)-oxy-toluic acid. — NaA' 4aq. Laminæ.—BaA'₂. Minute needles, less soluble than the Ba salt of the (α)-acid. By sulphonating (1,3,4)-xylenol by ClSO₃H, Hodgkinson (C. J. 63, 110) obtained a xylenol sulphonic acid yielding BaA'₂.

BaA', aq.

m-Xylenol sulphonic acid

C.H.Me. (OH).SO.H [1:3:6:4]. Formed from C.H.Me. (NH).SO.H by the diazo-reaction (Sartig, A. 230, 836). Slender needles, v. sol. water and alcohol. The acid and its salts give with FeCl. a bluish-violet colour turned green by alcohol.— KA'.—BaA'2 aq. Needles, v. e. sol. water.—PbA'2 2aq.

Ethyl derivative C.H.Me. (OEt).SO.H.

Ethyl derivative C₆H₂Me₂(OEt).SO₂H. Formed by heating diazo-m-xylene sulphonic acid with alcohol under pressure. Minute tables, v. sol. water and alcohol.—BaA'₂ Saq. Minute

plates, v. sol. water.

p. Evlenol sulphonic acid. Formed by dissolving p-zylenol in warm H₂SO₂. Hydrated

leaflets.—NaA'5aq. Tables.—BaA'2. Minute needles.

Reference.—NITRO-XYLENOL SULPHONIC ACID.
XYLENYL-AMIDOXIM v. DI-METHYL-BENZOIC ACID.

XYLENYL ALCOHOL v. DI-OXY-XYLENE.

XYLENYL CHLORIDE v. Tetra-ω-CHLORO-

YLENE.

XYLETIC ACID is OXY-DI-METHYL-BENZOIC ACID.

XYLIC ACID is DI-METHYL-BENZOIC ACID.

XYLIDIC ACID is TOLUENE DICARBOXYLIC

c-o-XYLIDINE C₈H₂Me₂(NH₂) [1:2:3].—Di-methyl-phenyl-amine. Amido-xylene. Mol. w. 121. (223° i.V.) at 739 mm. S.G. ¹⁵ 991, Obtained by reducing di-bromo-o-xylidine C₆HMe₂Br₂(NH₂) [1:2:4:5:3] with sodium-amalgam in the cold (Töhl, B. 18, 2562). Got also by heating (1,2,4)-xylenol with NH₄Br and ammonio-zinc bromide at 300° (Müller, B. 20, 1039). Prepared from o-xylene by nitration and reduction (Nölting a. Forel, B. 18, 2671; Wroblewski, B. 18, 2904). Oil. When oxidised by K₂Cr₂O₇ and H₂SO₄ ii yields a small quantity of o-xyloquinone [55°]. Converted by the diazo-reaction into o-xylenol [73°]. H₂SO₄ and HNO₂ give (4,1,2,8)-nitro-xylidine [114°] and (6,1,2,3)-nitro-xylidine [65°] (Nölting a. Stoecklin, B. 24, 567).—Salts: B'HClaq. Plates. Begins to sublime at 100°.—B'HNO₂. Needles.—B'₂H₂SO₄. Large plates, sl. sol. water.

Acetyl derivative C.H.Me.NHAc. [134°]
(N. a. F.); [131°] (T.; W.). Slender needles.

i-o-Xylidine C.H.Me.(NH₂)[1:2:4]. [49°].
(226°). S.G. 12° 1·0755. Prepared from oxylene by nitration and reduction (Jacobsen, B. 17, 159). Got also by heating m-toluidine hydrochloride with MeOH at 300° (Limpach, B. 21, 643). Monoclinic crystals, not coloured by exposure to light and air. Gives no colour with bleaching powder. Its solutions colour pinewood yellow. On nitration it yields (6,1,2,4)-nitro-xylidine [137°] and an isomeride [80°] (N. a. S.). Yields a saffranine on oxidation together with di-amido-di-phenyl-amine (Nietzki, B. 19, 8163).—Salt: B'HCl aq. Prisms, v. sol. water. Yields \$\psi\$-cumidine when heated with MeOH at

310° (N. a. F.).

Acetyl derivative C.H.Me.(NHAc). [99°].
c.m.Xylidine C.H.Me.(NH.) [1:3:2]. (6).

Xylidine (214° i.V.) at 739 mm. (N. a. F.).
S.G. 22° 9942 (W.). Formed by distilling pamido-mesitylenic acid with lime (Schmitz, A.
193, 179) and by reducing c-nitro-m-xylene (Grevingk, B. 17, 2430). Obtained from commercial xylidine by crystallisation of the sulphate; the last mother-liquor depositing crystals which are converted into the acetyl derivative by distillation with HOAc. The acetyl derivative is saponified by treatment with dilute (25 p.c.) H.2804 (Nölting a. Pick, B. 21, 3150; cf. Wroblewsky, B. 12, 1226). Gives m-xyloquinone [73°] on oxidation (Nölting a. Forel, B. 18, 2676). Yields (4,1,8,2)-nitro-xylidine [82°] on nitration (Nölting a. Stoecklin, Bl. [8] 5, 381).—Salts: BHCl.—B'HCl 2aq. S. 9.7 at 18°. Monoclinic plates. Yields mesidine on heating with MeOH at 310°—B'HNOs. S.
2-2. Needles.—B'H.28O4. Very soluble needles.—B'H.28O423aq. S. 60.

Acetyl derivative [174°] (G.); [176.5°] \ (N. a. F.). Needles.

Benzoul derivative [140°]. Needles. Yields benzoyl-nitro-xylidine [178°] on nitration

(Hübner, A. 208, 318)

s-m-Xylidine C_sH₃Me₂(NH₂) [1:8:5]. (222° i.V.). S.G. 15 972. Formed by reduction of s-nitro-m-xylene [71°] (Thöl, B. 18, 362; Nölting a. Forel, B. 18, 2678). On oxidation by K₂Cr₂O, and H SO, it gives m-xyloquinone [73°]. By boiling the diazo-compound with water xylenol [68°] is formed. ClCO_Et forms C_sH₃Me₂NH.CO₂Et [77·5°] (Frentzel, C. C. 1888, 1361). -B'HCl. MeOH at 250°-300° yields iso-1361).—B'HCI. MEUR at 200 -000 yiona accumidine C₄H₂Me₄(NH₂) [5:4:3:1] (Limpach, B. 21, 643).—B'H₂SO₄aq.—B'HNO₂. S. 4:66 at 13°. Acetyl derivative [140:5°] (N. a. F.); [138°] (T.); [144:5°] (W.). Colourless plates (from alcohol or ether).

i-m-Xylidine C₆H₃Me₂(NH₂) [1:3:4]. (a)-Xylidine. (215') at 745 mm. (G.). S.G. 35

·9184 (H.). `S.V. 148·3.

Formation.—1. By reducing i-nitro-m-xylene (Deumelandt, A. 144, 273; Tavildaroff, B. 2, 553; Wroblewsky, A. 192, 215; 207, 91; Grevingk, B. 17, 2430; Wallach, A. 258, 331). It may be isolated from commercial xylidine by conversion into hydrochloride and several crystallisations from water; the yield being about 40 p.c. (Nölting a. Forel, B. 18, 2677).—2. By heating o- and p-toluidine hydrochloride with MeOH at 300° (Hofmann, B. 9, 1295; Limpach, B. 21, 640).—3. By distilling (4,3,1,5)-amidomesitylenic acid with lime (Schmitz, A. 193, 177).—4. By heating i-m-xylenol with ammonium-zinc bromide and NH, Br (Müller, B. 20,

Reactions.-1. When dissolved in H2SO4 (10 pts.) it yields C₆H₂Me₂(NH₂)(NO₂) [1:3:4:6] [123°] on nitration (Nölting a. Collin, B. 17, 265).—2. The hydrochloride heated with MeOH at 310° yields mesidine.—3. CICO₂Et forms C_eH_sMe, NH.CO,Et [57°] (Frentzel, C. C. 1888, 1361).—4. When mixed with di-amido-di-phenylamine it yields a saffranine on oxidation (Nietzki, B. 19, 3163).—5. When m-xylidine (4 pts.) is heated with S (1 pt.) at 190° it gives off H₂S and forms a base C₁₆H₁₆N₂S [107°] (283° at 14 mm.) which crystallises from alcohol in yellowish-white prisms and yields C16H15AcN2S [227°] (Anschütz a. Schultz, B. 22, 582). By further heating with sulphur a homologue of primuline is formed.—6. Enanthol forms oily C₁₈H₂₈NO (Leeds, B. 16, 288).—7. Acrolein yields C, H, N, a reddish-yellow mass which on distillation yields cryptidine C₁₁H₁₁N (270°) (Leeds, A. C. J. 5, 1).

Salts.—B'HCl. Monoclinic prisms.-B'HCl $\frac{1}{2}$ aq. Plates (Staedel a. Hölz, B. 18, 2919).—B'₂H₂PtOl₂.—B'HBr.—Trimetric plates.
—B'HNO₂.—B'₂H₂SO₄4½aq. Cubes. Loses

xylidine on evaporation of its solution (Nölting Pick, Ei. [2] 50, 606).—B'H'PO. Only phosphate formed (Lewy, B. 19, 2728)—B'H,C.O. S. 3:32 at 18°.—B',ZnCl, (Lachovitch, M. 9, 514).—B'HgH,(SO,),2 aq.—B',Cu,H,(SO,),2 (Denigès, C. R. 112, 870).

Formyl derivative C.H.Me.NH.OHO. [111°] (G.); [114°] (G. a. M.). Got by heating the base (12g.) with formic acid (5g.) at 150° (Gasiorowski a. Merz, B. 18, 1011; Gudemann,

B. 21, 2549). Crystals (from water). Converted by P2S, at 100° into the thioformyl derivative [1050

Mono-acetyl derivative

 $C_cH_sMe_j(NHAc)$. [129°]. Needles or plates (from dilute alcohol). P_2S_s forms the compound $C_sH_sMe_2NH.CSMe$ [95°] (Jacobsen a. Ney, B.22, 907).

Di-acetyl derivative C₆H₃Me₂NAc₂. [60°]. Formed, together with the mono-acetyl derivative, by boiling the base with Ac2O (Wal-

lach, A. 258, 301).

C₆H₃Me₂NHBz. Benzoylderivative [192°]. Needles (Hübner, A. 208, 318). Converted by HNO, into the nitro-benzoyl derivative [184°] which is reduced by tin and HOAc to $C_6H_2Me_2 < NH > CPh [195°]$.

Phthalyl derivative [158°]. **Prisms**

(from alcohol).

p-Xylidine C₀H₁Me₁(NH₂) [1:4:2]. [15·5°] (Michael, B. 26, 89). (215° i.V.) at 789 mm. (N. a. F.); (213·5° uncor.) (M.); (220°) (S.). S.G. 18 ·980. Obtained from m-xylene by nitration and reduction (Schaumann, B. 11, 1537; Nölting a. Forel, B. 18, 2664; 19, 2680). Occurs to the extent of about 25 p.c. in commercial xylidine, , from which it may be isolated by dissolving in warm H₂SO₄ and pouring into hot water. The sparingly soluble m-xylidine sulphonic acid crystallises out, and the mother-liquor, after successive treatment with CaCO, and Na2CO, yields on evaporation sparingly soluble, pearly plates of sodium p-xylidine sulphonate, which on distillation with lime give p-xylidine almost without loss. May also be got from crude xylidine by means of its benzylidine derivative.

Reactions.—1. Oxidised by K2Cr2O, H₂SO₄ to p-xyloquinone [123°], the yield being 70 p.c. —2. Converted by the diazo-reaction into p-xylenol [75°] (210°).—3. On heating with sulphur it yields C₁₈H₁₈N₂S [144°], which crystellions from also be in well-with readless and the sulphur than t tallises from alcohol in yellowish needles, and yields an acetyl derivative [212°] (Anschütz a. Schultz, B. 22, 585).—4. The hydrochloride heated with MeOH at 290° under pressure forms ψ-cumidine (Limpach, B. 21, 646).-5. Bensoic aldehyde forms C,H,Me,N:CHPh [102°] (Pflug, A. 255, 168). The same body [96°] is got by adding benzoic aldehyde to an alcoholic solution of thionyl-p-xylidine (Michaelis, A. 274, 237).

Salts.—B'HCl aq. Leaflets, subliming at 125°-130° in needles.—B'HNO, Needles.— B'H2SO. Plates, sl. sol. water. -B'2H2C2O.

Formyl derivative C.H., NO.

Needles (Pflug, A. 255, 168)

Acetyl derivative C.H.Me.NHAc. [189]. sms. Yields on nitration acetyl-nitro-p-Prisms. xylidine [192°].

Bromo-acetyl derivative C₈H₁Me₂.NH.CO.CH₂Br. [145°]. Formed from bromo-acetyl bromide and p-xylidine in benz-ene (Abenius, J. pr. [2] 40, 435). Needles (from diluto alcohol). Boiling alcoholic potash converts it into an azine and the compound C₆H₃Me₂NH.CO.CH₂.OEt [50°].

Oxalyl derivative (C.H.Me.NH), C.O. Formed by heating the oxalate at 130°. Needles, subliming at 125°. An oxalyl xylidine (204°) was obtained by Genz (B. 3, 227) from crude,

commercial xylidine.

derivative C.H.Me.NHBz. Bensoyl[140°]. Needles.

References .- Bromo-, Chloro-, and NITRO-TYLIDINE.

XYLIDINE SULPHONIC ACID v. AMIDO-XYLENE SULPHONIC ACID and BROMO- and NITRO-XYLIDINE SULPHONIC ACIDS.

XYLIDINIC ACID v. TOLUENE DICARBOXYLIC

XYLINDEÏN. C, 65.5 p.c.; H, 4.7 p.c.; N, 1.0 p.c. A green colouring matter in green decayed wood (Rommier, C. R. 66, 108; Liebermann, B. 7, 1102). Extracted by phenol and ppd. by alcohol or ether. Four-sided blue plates, with coppery lustre. H,SO, forms a green solution. Rideal (C. N. 53, 277) finds that the blue substance in decayed birch-wood is soluble in chloroform, while the blue resin obtained from coniferin by successive treatment with H₂SO₄ and water is insol. chloroform.

(252° **XYLITONE** $C_{12}H_{16}O$. *Xylite oil*. (252°) (Pinner, *B*. 15, 589; 16, 1729); (c. 240°) (Claisen a. Ehrhardt, *B*. 22, 1013). S.G. 935. Xylite oil. A product of the action of HCl on acetone (P.). Formed also by the action of NaOEt on mesityl oxide in ether, and in the preparation of acetylacetone by the action of boiling EtOAc and NaOAc on acetone (C. a. E.). Oil, easily resini-

fied by strong acids.

(310°-320°) Dixylitone C, H20O2 duct of the action of HCl on acetone (Pinner).

XYLOHYDROQUINONE v. HYDROXYLO-QUINONE.

XYLOÏC ACID v. DI-METHYL-BENZOIC ACID.

XYLONIC ACID C.H.O. $[\alpha]_{\rm D} = 17.48^{\circ}$. Formed by oxidising xylose with bromine (Tollens, A. 260, 307). Dextrorotatory. Its Ca, Zn, and Ag salts are amorphous. - SrA'2. $[a]_D = +12.14^\circ$

XYLONITRILE v. Nitrile of DI-METHYL-BENZOIC ACID.

o-XYLOQUINONE CaHsO2 i.e.

C.H.Me.O. [1:2:3:6]. [55°]. Formed by oxidation of c-o-xylidine by K.Or.O., and H.SO., the yield being about 10 p.c. (Nölting a. Forel, B. 18, 2673). Yellow needles (by sublimation), v. sl. sol. water, m. sol. alcohol and ether.

m. Xyloquinone C₈H₂Me₂O₂ [1:3:2:5]. [78°]. Formed by oxidation either of c-m- or s-m-xylidine with K₂Cr₂O₇ and H₂SO₄ (Nölting a. Forel, B. 18, 2679). Got also by oxidation of mesidine (Nölting a. Baumann, B. 18, 1151). Yellow needles.

p-Xyloquinone C₆H₂Me₂O₂[1:4:2:5]. Prone. Metaphlorone. Mol. w. 136. [124°].

Formation.-1. Occurs, together with toluquinone and other bodies, among the products of the action of MnO2 and H2SO, on coal-tar and beechwood-tar (Rommier a. Bouilhon, C. R. 55, 214; Gorup-Besanez a. von Rad, Z. [2] 4, 560; Carstanjen, J. pr. [2] 23, 425).—2. By oxidation of p-xylylene-diamine (Nietzki, B. 13, 472; A. 215, 168).—3. By oxidation of ψ -currenol (Carstanjen, J. pr. [2] 23, 430).—4. By oxidation of ψ -cumidine by CrO₂ (Nölting a. Th. Baumann, B. 18, 1151; Sutkowski, R. 20, 977). 5. By distilling xylidine (213°) with H2SO4 and MnO₂; the yield being 5 p.c. (C.).—6. Py oxidation of p-xylidine with K₂Cr₂O₇ and H₂SO₄, the yield being over 70 p.c. (Nölting, Witt, a. Forel,

B. 18, 2667).—7. By heating diacetyl with dilute NaOHAq (Van Pechmann, B. 21, 1420).

Preparation .- 1. Commercial xylene (180°-140°) is heated with H2SO, at 100° for six days; the product poured into water; neutralised by CaCO₃; treated with K₂CO₃; and the K salts of the sulphonic acids fused with potash. The product is dissolved in water, acidified, and extracted with ether; and the mixture of xylenols so obtained is dissolved in warm H,SO4, and then diluted with water and distilled with MnO2. Crystals of p-xyloquinone appear in the condenser and receiver (Carstanjen, J. pr. [2] 23, 427) .- 2. 'Yellow carbolic acid' or 'creosote,' the mother-liquor from which phenol has crystallised, is dissolved in H2SO4, diluted with water, and then distilled with MnO2. The solid found in the receiver is crystallised from alcohol when the first crop of crystals is xyloquinone (E. Carstanjen, J. pr. [2] 23, 423).

Properties.—Long yellow needles (from benzene), sl. sol. water and cold alcohol, v. sol. ether. Volatile with steam. May be sublimed.

Reactions .- 1. Distilled with sinc-dust gives a hydrocarbon that oxidises to terephthalic acid.-2. Reduced by aqueous SO, to hydroxyloquinone.-3. Conc. HCl converts it into a mixture of mono- and di-chloro-hydro-xyloquinones,

C_eHClMe₂(OH)₂ [147°] and C_eCl₂Me₂(OH)₂ [175°].

Mono-oxim C_eH₂Me₂O(NOH). Nitroso-pxylenol. [168°]. Formed by heating p-xyloquinone (2 pts.) with an alcoholic solution of hydroxylamine hydrochloride (1 pt.) (Goldschmidt a. Schmid, B. 18, 568; Sutkowski, B. 20, 978). Obtained also by adding KNO₂ and excess of HOAc to an alkaline solution of p-xylenol (Oliveri, G. 12, 162). Yellowish needles (from alcohol), v. sol. alcohol, ether, and benzene, insol. cold water. Alkalis form a deep orangered solution. Reduced by SnCl2 to amido-pxylenol. Oxidised by alkaline K, FeCy, to nitrop-xylenol.

Di-oxim C₆H₂Me₂(NOH)₂. [272°]. Formed by boiling the quinone (1 mol.) with an alcoholic solution of hydroxylamine hydrochloride (2 mols.) (Sutkowski, B. 20, 977). V. sl. sol. alcohol, acetic acid, and benzene, insol. water. Dissolves in alkalis with a yellow colour. By tin and HCl it is reduced to xylylene-p-diamine. Its di-acetyl derivative C₆H₂Me₂(NOAc)₂ crystallises from acetic acid in small yellow prisms,

[170°], v. sol. alcohol and benzene.

References .- DI-BROMO- and CHLORO-XYLO-QUINONE.

XYLORCIN v. DI-OXY-XYLENE.

XYLOSE v. p. 538.

XYLOSTEIN. A glucoside in the berries of Lonicera Xylosteum (Hübschmann, J. 1856, 691; Enz. C. C. 1856, 393). Crystalline, insol. water, sol. alcohol and ether.

XYLOYL-BENZOIC ACID v. PHENYL XYLYL KETONE CARBOXYLIJ ACID.

XYLOYL-FORMIC ACID v. XYLYL-GLYOXYLIC ACID.

pp-DIXYLYL C, H, i.e. [4:1:3] C₆H₃Me₂.C₆H₃Me₂[3:4:1]. [125]. pared by distilling mercury di-p-xylyl (Jacobsen,

B. 14, 2112). Long needles (from alcohol).

Dixylyl. (290°-295°). Formed from bromem (?)-xylene and Na (Fittig, A. 147, 88). Liquid.

Di-xylyl (so-called). (295°) (Oliveri, G. 12, 158). Is got by warming commercial xylene with H2SO, but never from xylene which has been made from xylene sulphonic acid. Hence it is probably not a benzene derivative (Armstrong, C. J. 45, 150). It is a colourless, fluorescent liquid, smelling like copaiba balsam.

XYLYL-ACETAMIDE v. Acetyl derivative of XYLIDINE and Amide of XYLYL-ACETIC ACID.

XYLYL-ACETIC ACID

 $[102^{\circ}].$ [1:3:4] $C_6H_3Me_2.CH_2.CO_2H$. Formed from its amide, which is got by the action of yellow ammonium sulphide on xylyl methyl ketone (Willgerodt, B. 21, 534). Slender needles.

 \underline{A} mide $C_8H_9.UH_2.CO.NH_2$. [183°].

Reference.— OXY-XYLYL-ACETIC ACID.
XYLYL ALCOHOL is Tolyl-Carbinol.

XYLYL-AMIDO-ACETIC ACID $C_{10}H_{13}NO_2$ i.e. [1:3:4] $C_0H_3Me_2$.NH.CH₂.CO₂H. [134°]. Formed by boiling m-xylidine (2 mols.) with water and chloro-acetic acid (1 mol.) (Ehrlich, B. 16, 205). Prisms (from dilute alcohol). Insol. water, v. sol. acius.

Xylide C₈H₉.NH.CH₂.CO.NHC H₉. [128°]. Formed by boiling xylidine (2 mols.) with chloro-acetic ether (1 mol.). Thick needles chloro-acetic ether (1 mol.). (from alcohol), insol. water and HClAq.

XYLYL-AMIDO-CROTONIC ETHER

[1:3:4] C₆H₃Me₂.NH.CMe:CH.CO₂Et. Formed from m-xylidine and acetoacetic ether (Conrad a. Limpach, B. 21, 526). Decomposed on heating into (Py. 1, 3; B. 2, 4)-oxy-tri-methyl-quinoline, di-xylyl-urea, and alcohol.

XYLYL-AMIDO-ETHYL-PHTHALIMIDE [1:3:4] $C_6H_3Me_2.NH.C_2H_4.N:C_6H_4O_2.$ [123°]. Formed by heating bromo-ethyl-phthalimide with m-xylidine at 140° (Newman, B. 24, 2197). Small needles, v. e. sol. benzene, sl. sol. cold

DI-m-XYLYL-DI-AMIDO-OIAZTHIOLE

 $\mathbf{O}_{18}\mathbf{H}_{20}\mathbf{N}_{4}\mathbf{S}$ i.e. $\mathbf{S} \stackrel{\mathbf{C}(\mathrm{NHC}_{8}\mathbf{H}_{9}):\dot{\mathbf{N}}}{\mathbf{C}(\mathrm{NHC}_{8}\mathbf{H}_{9}):\dot{\mathbf{N}}}$. [79°]. Formed by oxidising (1,3,4)-xylyl-thio-urea with aqueous H.O. (Hector, B. 23, 368). Insol. water. Forms a very soluble hydrochloride.—B',H,PtCl, Decomposes at 214°-217°.—B',AgNO, Explodes on heating. Blackened by sunlight. - B'C, H, N, O, [218°]. Yellow crystals, v. sol. alcohol. Cyanogen passed into its alcoholic solution forms C₁₈H₂₀N₄SCy₂ [103°].

Acetyl derivative C18H19AcN4S. Needles. Benzoyl derivative C₁₈H₁₈BzN₄S. [212°]. Nitrosamine C₁₈H₁₉(NO)N₄S. [146°].

XYLYLAMINE v. XYLIDINE and METHYL-BENZYL-AMINE

Di-o-xylyl-amine $(C_8H_9)_2NH$. (330°-345°). Formed, together with xylidine, by heating (1,2,4)-xylenol with ammonio-zinc bromide and NH Br at 300° (Müller, B. 20, 1039). Liquid, solidified at a very low temperature.

Di-m-xylyl-amine. (305°-310°). Formed in

like manner from m-xylidine. Oil.

Di-xylyl-amine. [162°]. Formed, together with a liquid isomeride, by heating xylidine with its hydrochloride (Girard a. Vogt).

XYLYL-BENZYL-AMINE Und, Ni.e. C.H., Me. NH.CH.Ph. (c. 205° at 15 mm.). Formed by heating m-xylidine with benzyl chloride at 160° (Joblin-Gonnes, Bl. [3] 6, 21). Yellowish oil, sol. alcohol and benzene.

p-Xylyl-benzyl-amine [1:4:2]C,H,Me,NH.CH,Ph. (322°) (Pflug, A. 255, 168

o-XYLYL BENZYL KETONE C16H16O i.e. C₆H₃Me₂.CO.CH₂Ph. [95°]. (210°-220° at 25 mm.). Formed from o-xylene, phenyl-acetic chloride, and AlCl. (Wege, B. 24, 3540). Plates, sol. alcohol, v. sol. ether. Yields a crystalline NaOEt and benzyl chloride yield C₆H₃Me₂.CO.CH(CH₂Ph)Ph [75°].

m-Xylyl benzyl ketone

[1:3:4] C₆H₂Me₂.CO.CH₂Ph. (350°). Formed, together with a smaller quantity of an isomeride [93°], by the action of AlCl, on a mixture of m-xylene and phenyl-acetic chloride (Söllscher, B. 15, 1681). Yields C_sH₃Me(CO₂H)₂ [8:4:1] on oxidation. NaOEt and benzyl chloride form oily C_sH₃.CO.CHPh.CH₂Ph (365°-375°).

p-Xylyl benzyl ketone C,H,.CO.CH,Ph. $(22\hat{0}^{\circ}-230^{\circ}$ at 26 mm.). Formed in like manner from p-xylene (Wege). Yields an oxim [99°] and a phenyl-hydrazide [96°]. NaOEt and benzyl chloride yield C₈H_o.CO.CHPh.CH₂Ph [60.5°].

m-XYLYL-BENZYL-THIO-UREA

C₈H_eNH.CS.NHCH₂Ph. [85°]. Formed from benzyl-thiocarbimide and m-xylidine in alcohol (Dixon, C. J. 59, 558). Monoclinic prisms, v. sol. boiling alcohol and chloroform.

XYLYL BROMIDE v. BROMO-XYLENE.

XYLYL-tert-BUTANE C₆H₃Me₂.CMe₃. (201°). Formed from m-xylene, isobutyl bromide, and AlCl, (Baur, B. 24, 2840). Got also from isobutyl alcohol, m-xylene, and H2SO, or ZnCl2 (Nölting, B. 25, 791). Oil, yields mesitylenic and trimesic acids on oxidation.

XYLYL-BUTYL ALCOHOL

[2:4:1] C₆H₃Me₂.CHPr.OH. (above 270°). Got by reducing xylyl propyl ketone with zinc-dust and KOHAq, or with sodium-amalgam (Claus, J. pr. [2] 46, 474). Oil, v. sol. alcohol and ether.

Tri-xylyl-butyl alcohol (C,H,),C.CMe,OH. (above 300°). Formed from -xylene, tri-chlorotert-butyl alcohol, and AlCl, (Willgerodt a. Genieser, J. pr. [2] 37, 361). Oil, smelling like mushrooms.

m-XYLYL-n-BUTYRIC ACID [1:3:4]C₀H₂Me₂.CH₂.CH₂.CH₂.CO₂H. Formed by saponifying the amide (Claus, J. pr. [2] 46, 476). Needles, v. sol. hot water. May be sublimed.—BaA', 4aq.—CaA', 4aq. Prisms, v. sol. water and alcohol.

Amide CoH, Me2.C3H8.CO.NH2. [124°]. Got by heating xylyl propyl ketone with S and ammonium sulphide in sealed tubes at 250°.

Flat silky needles. May be sublimed.

p-Xylyl-n-butyric acid [1:4:2]C_uH₂Me₂.CH₂.CH₂.CH₂.CO₂H. [70°]. Needles, sl. sol. cold water. — BaA', 4aq. CaA'24aq. Needles, sol. water and alcohol.

1 mide C₈H₄,C₄H₆,CO.NH₂. [125°]. Formed from p-xylyl propyl ketone (C.). Flat needles, v.

sol, hot water.

n.-Xylyl-isobutyric acid [1:3:4] C_eH₂Me₂·CH₂·CHMe.CO₂H. [70°]. Flat pearly needles, sol. hot water, alcohol, and ether. May be sublimed.

Amide C.H., C.H., CO.NH. [120°]. Got by heating xylyl isopropyl ketone with 8 and ammonium sulphide at 235° (Claus, J. pr. [2] 46, 483). Needles, sol. hot water.

XYLYL-CARBAMIC ACID

[1:3:4] O.H.Mez.NH.COzH. Ethyl ether EtA'. Formed from m-xylidine and chloroformic ether (Hofmann, B. 3, 657; Frentzel, C. C. 1888, 1361). Needles.

Xylyl-carbamic acid [1:3:5]C_eH₃Me₂.NH.CO₂H. Ethyl ether EtA'. [77.5°] (Frentzel).

DI-p-XYLYL-CARBINOL

([2:5:1] C₀H₃Me₂)₂CH.OH. [131°]. Formed by reducing di-p-xylyl ketone with zinc-dust and KOH (Elbs, J. pr. [2] 35, 484; B. 19, 408). White needles (from alcohol), nearly insol. Aq.

XYLYL CYANATE [1:3:4] C₆H₃Me₂.N:CO.

(205°) (Frentzel, C. C. 1888, 1361).

Xylyl cyanate [1:3:5] $O_6H_3Me_2$. N:CO. (208.5°).

XYLYL CYANURATE

[1:3:4] (C₆H₃Me₂)₃N₃C₃O₃. [162°] (Frentzel, C. C. 1888, 1361).

XYLYLENE ALCOHOL v. DI-OXY-XYLENE.

XYLYLENE-DI-ALLYL-DI-THIO-DI-UREA $(CH_s)_2C_6H_2(NH.CS.NH.C_9H_s)_2[1:4:2:6]. [112.5°].$ Made from p-xylylene diamine and allyl thiocarbimide (Lellmann, A. 228, 252). Needles (from alcohol). Not decomposed on melting.

m-XYLYLENE-m-DIAMINE

 $C_6H_2(CH_3)_2(NH_2)_2[1:3:4:6]$. m-Di-amido-m-xylene [105°]. Formed by reduction of nitro-xylidine [123°] or of di-nitro-xylene [93°] (Fittig, A. 147, 18; Grevingk, B. 17, 2426; Witt, B. 21, 2419). White crystals. Sublimable. Does not yield a eurhodine on boiling with nitroso-dimethylaniline, NaOAc, and HOAc.— B"2HCl.— B"H₂SnCl₄.—B"H₂SO₄. Crystalline, v. sol. Aq. m-Xylylene-o-diamine

 $C_6H_1Me_2(NH_2)_2[1:3:5:6][78:5^\circ]$. Formed by reduction of $[2:4:1]C_6H_3Me_2.N_2.C_6H_2Me_2.NH_2$ [1:3:5:6] (Nölting a. Forel, B. 18, 2683) and also by the action of tin and HClAq on di-bromo-di-nitro-mxylene [191°] (Jacobsen, B. 21, 2821), and on (5,1,3,6)-nitro-xylidine (Hofmann, B. 9, 1298). Plates (from water), sl. sol. cold benzene, sol.

m-Xylylene-m-diamine

C.H.Me. (NH.), [1:3:4:2]. [64°]. Formed by reduction of nitro-xylidine [78°] or of di-nitro-m-xylene [82°] (Grevingk, B. 17, 2426). White needles. Gives a Bismarck brown with nitrous acid and a chrysoïdine with diazobenzene.

p-kylylene-o-utility $C_eH_2Me_2(NH_2)_2$ [1:4:2:3]. [75°]. Formed by reduction of c-di-nitro-p-xylene (Lellmann, A. White needless B. 19, 145). White needless A. p-Xylylene-o-diamine (by sublimation). Gives a deep-red colour with

FeCl. p-Xylylene-m-diamine p-Xylylene-m-diamine $C_0H_1Me_2(NH_2)_2$ [1:4:2:6]. [102°]. Formed by reducing (2,6,1,4)-di-nitro-xylene (L.; N.). Needles (by sublimation). Behaves like a m-

 $oldsymbol{p}$. Xylylene- $oldsymbol{p}$ -diamine

 $C_1H_1Me_2(NH_2)_2$ [1:4:2:5]. [142°] (S.); [147°] (N. a. F.); [150°] (N.). Formation.—1. By reduction of xylene-azoxylidine (amido-azo-xylene) [2:5:1] C₅H₃Me₇N₇C₅H₂Me₂NH₂[1:2:5:4] by zinc and HClAq (Nietzki, B. 13, 471; Nölting a. Forel, B. 18, 2685).—2. By reduction of nitro-paylidite [142°] (Nölting, Witt, a. Forei, B. 18, 2666; von Kostanecki, B. 19, 2818; Marckwald, D. 19, 2818; Marckwald, D. 19, 2818; Marckwald, D. 2818; Mar p-xyloquinone C.H.Me2(NOH), with tin and HCl (Sutkowski, B. 20, 979).

Properties. - Colourless needles, sol. hot water and alcohol, m. sol. benzene and ether. Yields xyloquinone [123°] on oxidation. Not oxidised by exposure to air. May be sublimed.

Salts.-B"2HCl. Colouriess leaflets. -B"H2SO4. Crystalline meal.

o-Xylylene exo-diamine

[1:2] C₄H₄(CH₂.NH₂)₂. Formed by heating the phthalimide C₄H₄(CH₂.N:C₂O₂:C₄H₄)₂ with conc. HClAq at 200° (Strassmann, B. 21, 579). Alkaline liquid, smelling strongly like ammonia. Absorbs CO₂. Gives with FeCl, a pp. of yellowish-red needles.—B"2HCl.—B' 2HAuCl4. Yellow plates.—B"2C,H,N,O,. Yellow needles, decomposing above 170° without melting.

Acetyl derivative C₈H₁₀Ac₂N₂ [14v°]. Benzoyl derivative C₈H₁₀Bz₂N₂. [168°].

m-Xylylene-exo-diamine [1:3] C₆H₄(CH₂NH₂)₂. (247°). Formed by the action of conc. hydrochloric acid at 220° on the $C_eH_4(CH_2.N:C_eH_4O_2)_2$ [237°], which is got by heating potansium phthalimide with di- ω -bromom-xylene (Brömme, B. 21, 2705). Liquid, sol. water, miscible with alcohol and ether. I bsorbs $CO_x - B''^2HCl.$ Needles (from water). -B"2H2PtCl6.—B"C6H3N3O7. Yellow spangles, decomposing at 187°.

Acetyl derivative C.H.(CH.NHAC). [119°]. Crystalline mass, v. sol. hot water.

Reference.—NITRO-XYLYLENE-DIAMINE.

XYLYLENE-BENZENYL-AMIDINE v. Benz-ENYL-XYLYLENE-DIAMINE.

XYLYLENE BROMIDE v. DI-BROMO-XYLENE. XYLYLENE-CHLORO-MALONIC ETHER v. CHLORO-XYLYLENE-MALONIC ETHER.

Xylylene-exo-di-chloro-malonic ether v. Di-CHLORO-PHENYLENE-DI-METHYL-DI-MALONIC ETHER. XYLYLENE-ETHENYL-AMIDINE C10H12N2

i.e. CMe $<_{
m N}^{
m NH}>$ C₆H₂Me₂. Formed by reducing C₆H₂Me₂(NO₂)(NHAc) [1:3:5:4] with tin and HCl (Hobrecker, B. 5, 922). Resinous mass.— B'HCl. Large needles.—B'_H_PtCl_.—B'HNO. Long needles.

XYLYLENE IODIDE v. DI-IODO-XYLENE.

XYLYLENE-DI-MALONIC ACID v. PHENYL-ENE-DI-METHYL-DI-MALONIC ACID.

o-XYLYLENE SULPHIDE C.H.S i.e.

C₆H₄<CH₂>S. [c. 0°]. Formed by heating di-w-bromo o-xylene with K₂S (Leser, B. 17, 1824; Hjelt, B. 22, 2904). Oil, smelling like mercaptan. Very unstable, quickly turning into a black resin.—B'₂HgCl₂. Long needles.

Methylo-iodide B'MeI. [155°].

Di-xylylene disulphide [1:3:4:5] C₆H₂Me₂:S₂:C₆H₂Me₂ [5:4:3:1]. [118°]. Formed by heating xylylene-diazosulphide at 200°-250° (Jacobsen a. Ney, B. 22, 911). Conc. H₂SO₄ forms a l'ue liquid, remainirg blue cu dilution.

u-DI-XYLYL-ETHANE C₁₂H₂₂ i.e. CH₂:CH_{(C2}H₃₎. (324°). S.G. ³² ·966. Formed from ethylidene chloride, m-xylene, and AlCl₄ (An ichütz, B. 18, 665; A. 285, 826).

Reference.—Chloro-di-xylyl-ethane. DI-XYLYL-ETHYLENE C1.H. i.e. 2666; von Kostanecki, B. 19, 2818; Marckwald, C.H.Me. CH:CH:CH:CH.O.H.Me. Tetra-rethyl-stilbens. B. 28, 1021).—8. By reduction of the di-oxim of [106°]. (325°-840°). Formed by distilling C.H.Me.CH.CH.Cl, which is got from di-chlorodi-ethyl oxide, m-xylene, and H2SO4 (Hepp, B. 7, 1416). Spangles (from alcohol), sol. ether and CS2 Unites with bromine. Oxidised by dilute HNOs to (1,3,4)-di-methyl-benzoic acid [122°

Di-xylyl-ethylene C,H,Me,CH:CH.C,H,Me, [157°]. Formed in like manner from p-xylene. Plates (from alcohol). Less soluble than the

preceding isomeride.

Tetra-xylyl-ethylene (C₆H₈Me₂)₂C:C(C₆H₈Me₂)₂. [245°]. Formed from xylene, CHCl₃, and AlCl₃ (Schwarz, B. 14, 1528). Yellow plates.

m-XYLYL-ETHYLENE-DIAMINE

C₆H₃Me₂NH.C₂H₄.NH₂. (274°). Formed from m-xylyl-amido-ethyl-phthalimide and HCl (Newman, B. 24, 2197).—B'HCl. [173°].—B'2H2PtCla. -B'C'H₃N₃O₇. [141°]. Reddish-yellow needles, v. e. sol. alcohol.

DI-m-XYLYL-ETHYLENE DIKETONE

 $C_2H_4(CO.C_8H_3Me_2$ [4:3:1]), [129°]. Formed from m-xylere, succinyl chloride, AlCl₂ and CS₂ (Claus, B. 20, 1375). Needles, insol. water, v. sol. alcohol and ether. Dilute HNC, oxidises it to (1,3,4)-di-methyl-benzoic acid.

 $Ox^{\frac{3}{4}m} C_2H_4(C(NOH).C_8H_9)_2$. [140°].

Phenyl-hydrazide C₂H₄(C(N₂HPh).C₅H₉)₂. [189°]. Crystals. Di-p-xylyl ethylene diketone

 $C_2H_4(CO.C_6H_8Me_2[2:4:1])_2$. [123°]. from p-xylene, succinyl chloride, AlCl, and CS2 Needles. (Claus).

m-XYLYL ETHYL KETONE

[1:3:4] C₆H₈Me₂.CO₄C₂H₆. (239°). Light oil, with aromatic odour, not solid at -10° (Claus, J. pr. [2] 43, 140). Yields (1,3,4)-di-methylbenzoic acid on oxidation.

Phenyl-hydrazide [126°]. Plates, v. sol.

hot water, alcohol, and ether.

p-Xylyl ethyl ketone [4:1:2] C₆H₈Me₂.GO.C₂H₈. (238° uncor.). Formed from p-xylene, AlCl, and propionyl chloride (Claus a. Fickert, B. 19, 3182). Oil. Oxidised by KMnO₄ to C₆H₄Me₂.CO.CH₂.CO₂H [132°] and some (4,1,2)-di-methyl-benzoic acid.

m-XYLYL ETHYL KETONE CARBOXYLIC

ACID [4:2:1] C₆H₃Me₂.CO.CH₂.CH₂.CO₂H. Di-methyl-bensoyl-propionic acid. [108°]. Formed from m-xylene, succinyl chloride, AlCl_s, and CS₂ (Claus, B. 20, 1876). Needles, v. sol. hot water, insol. cold.—NH₄A'.—KA' 4aq. V. sol. water.—NaA' 4aq. Needles.—BaA', 3aq. Needles.—PbA'2: white pp.—AgA': white crystalline pp.

p-Xylyl ethyl ketone carboxylic acid [4:1:2] C₆H₈Me₂CO.CH₂.CH₂.CO₂H. [84°]. Formed from p-xylene in like manner (C.). Colourless needles.

XYLYL-DI-ETHYL-PHOSPHINE C.H.Me..PEt. (260°). Formed by the action of ZnEt. on C.H.Me..PCl. (Czimatis, B. 15, 2016). Thick, colourless oil. Y.elds B'MeI [90°],

B', Me, PtCl, [202°], and B'EtI [136°]. m-XYLYL-GLYOXALINE C.H.N CH:N

[32°]. (279°). Formed by the action of HNO_a on $C_aH_aN < CH = CH = (192°]$, which is got by the action of HCl on the product of the action of amido-acetal on m-xylyl-thio-urea (Marck- hol.

wald, B. 25, 2368). Sl. sol. water. —Aurochloride [165°].-Picrate. [159°]. Yellow needles, m. sol. chloroform.

o-XYLYL-GLYOXYLIC ACID

[4:3:1] C₆H₃Me₂.CO.CO₂H. [92°]. Formed by oxidising o-xylyl methyl ketone with alkaline K_sFeCy_s (Buchka a. Irish, B. 20, 1766).—BaA'₂.

m-Xylyl-glyoxylic acid [4:2:1] C₆H₃Me₂.CO.CO₂H. $[54^{\circ}].$ Formed by oxidising m-xylyl methyl ketone with KMnO (Claus, B. 19, 231; J. pr. [2] 43, 142) Flat prisms. Yields (4,2,1) di-methyl-benzoic scid on boiling with dilute HNO3. Gives rise to C₈H₉.CH(OH).CO₂H [119°] on reduction.— CaA'22aq. Needles.—BaA'22aq.—AgA'. Needles, sol. hot water.

p-Xylyl-glyoxylic acid

[5:2:1] C₆H₃Me₂.CO.CO₂H. [70°-80°]. Formed by oxidation of p-xylyl methyl ketone by aqueous KMnO, (Claus a. Wollner, B. 18, 1859). Crystalline, v. sol. alcohol, ether, and HOAc, v. sl. sol. water. Splits up above 200° into CO. and di-methyl-benzoic aldehyde. Oxidised by HNO, to CoH, Me2.CO2H. Ac2O and NaOAc yield dimethyl-cinnamic acid.—BaA', 6aq: small felted needles.—CaA'₂ 3aq.—AgA': crystalline pp. Ethyl ether EtA'. Oil.

XYLYL GLYOXYLIC ALDEHYDE

[1:3:4] $C_8H_8Me_2$ CO.CHO. Oxim [95°]. Formed from xylyl methyl ketone, NaOEt, and $C_8H_{11}NO_2$ (Söderbaum, B. 25, 3463). Needles (from benz. ene-ligroin). Acetic anhydride gives the acetyl derivatives of the syn-oxim C₈H₉.CO.CH:NOAc [54°] decomposed by alkalis with formation of C_cH_sMe₂.CO_cH [126°] and by hot Ac_cO yielding C_sH_sMe₂.CO.CN [47°]. AcCl converts the oxim into C_cH_sMe₂.C(OH)_c.CH:NOAc [142°], whence cold NaOH forms C.H.Me2.CH(OH).CO2H [103°], while NaOHAq at 50° forms 'xylyl formoun' C_8H_9 .CO.CH(OH).CO.CO. C_8H_9 [155°].

DI-XYLYL-GUANIDINE

HN:C(NH.C,H, [1:2:4])₂. [158°]. Formed by the action of PbO and alcoholic ammonia on dixylyl-thio-urea (Hofmann, B. 9, 1295). Needles, (from alcohol).

XYLYL-HYDRAZINE

[4:2:1] C₂H₃Me₂NH.NH₂. [85°]. Formed from xylidine by diazotisation followed by reduction with Na, SO₃, the resulting C₂H₂, N₂H₂, SO₃Na and being then treated with zinc-dust and HOAc (Klauber, M. 11, 282). Pale-yellow needles, v. sl. sol. water, v. sol. alcohol. Reduces Fehling's solution on warming. On heating with acetoacetic ether it yields a crystalline body [203°]. reduced in alcoholic solution by Na to oxyxylyl-methyl-pyrazole.

Salt.—C'HCl 2aq. [183°]. Small needles.

Di-o-xylyl-hydrazine [3:2:1] C_eH_sMe₂NH.NHC_eH_sMe₂ [1:2:3]. [141°]. c-Hydrazo-o-xylene. Got by heating nitro-o-xylene (20 g.) with NaOH (20 g.), alcohol (100 c.c.) and zinc-dust (25 g.), and reducing the product with alcoholic ammonium sulphide (Nölting a. Stricker, Bl. [2] 50, 613). White needles, sol. alcohol and ether. Easily oxidised by air.

u-Di-o-xylyl-hydrazine N₂H₄(C₂H₄Me₂[1:8:4]), [107°]. Formed by reducing nitro-o-xylene with sodium-amalgam (N. a. S.). Yellowish white needles, sol. alcoDi-m-xylyl-hydrazine

 $N_2H_2(C_eH_3Me_{,}[1:8:5])_s$. [125°]. Obtained from s-nitro-m-xylene (N. a. S.). Needles, easily oxidised by air.

u-Di-m-xylyl-hydrazine

 $N_2H_2(C_eH_1Me_2[1:2:4])_2$. [122°]. Got by warming nitro-m-xylene (30 g.) with NaOH (30 g.), zinc-dust (40 g.), and alcohol (250 c.c.) and treating the xylene-azo-xylene so obtained with alcoholic ammonia and H.S (N. a. S.). White needles, sol. alcohol and ether.

Di-p-xylyl-hydrazine $N_2H_2(C_6N_3Me_2[1:2:5])_2$. [145°]. Formed from nitro-p-xylene (N. a. S.). Needles, not affected by air.

XYLYLIC ACID v. DI-METHYL-BENZOIC ACID. a-Xylylic acid v. ToLYL-ACETIC ACID.

p-XYLYLIDENE-DI-IMINE

C.H. (CH:NH)2 [1:4]. Formed by the action of alcoholic NH, or dry NH, gas upon terephthalic aldehyde (Oppenheimer, B. 19, 576). brittle crystals. V. sl. sol. alcohol and ether.

DI-XYLYL-KETONE $CO(C_6H_2Me_2)_2$. (c. 340°) Obtained from (m?)-xylene, COCl₂, and AlCl₂ (Ador a. Rilliet, B. 11, 399). Liquid, not solid at -60°. Split up by long boiling into H₂O and

C₁₇H_{1e}.
Di-p-xylyl ketone
Me_{ss}.CO.C_c [2:5:1] $C_aH_aMe_x$.CO. $C_aH_aMe_z$ [1:2:5]. (827° uncor.). Formed from p-xylene, CS_z , and $COCl_z$ in closed vessels; the yield being 55 p.c. (Elbs, J. pr. [2] 35, 481). Oil. Quickly decomposed by distillation into H₂O and (B. 2, 1',4')-trimethyl-anthracene [227°]. Reduced by zincdust and alcoholic KOH to di-p-xylyl-carbinol [131°].

Di-xylyl-tetraketone $C_{a}H_{a}.CO.CO.CO.CO.C_{a}H_{a}Me_{a}[4:3:1].$ [180°]. Formed by the action of cold HNO, (S.G. 1.4) on C_8H_9 .CO.CH(OH).CO.CO. C_8H_9 (Söderbaum, B. 25, 3475). Scarlet needles (from CS2). On dissolving in HOAc and ppg. by water it yields a

dihydrate [100°] C₂₀H₂₂(

XYLYL MERCAPTAN C.H.Me.SH. (214°). S.G. ¹³ 1.036. Got by treating xylene sulphonic chloride with Zn and dilute H₂SO₄ (Yssell, Z. 1865, 360), and also by heating xylyl ethyl xanthate with alcoholic potash (Leuckart, J. pr. 41, 192). Liquid. - Hg(S.C.H.)2. Pb(S.C.H.). Yellow powder. TRI-p-XYLYL-METHANE

 $\mathbf{CH}\{\mathbf{C_6H_3Me_2}\}_{\mathbf{3}}$ [188°]. (above 360°). From di-p-xylyl-carbinol, p-xylene, and P₂O₅ by boiling for four hours (Elbs, J. pr. [2] 35, 484). Crystalline grains (from alcohol), v. sol. ether and benzene.

o-XYLYL METHYL KETONE

[4:8:1] C₆H₃Me₂.CO.CH₃. (246°). Formed from o-xylene, AcCl, and AlCl₃ (Claus, B. 18, 1856; 19, 232; J. pr. 1890, 410). Oil, smelling like coumarin (Armstrong a. Kipping, C. J. 63, 81). Yields (4,3,1)-di-methyl-benzoic acid on oxidation. Conc. HOlAq forms C₂₀H₂₂O [114°]. P₂O₃ gives a body melting at 165°. Zinc-dust reduces gives a body melting at 165°.

the ketone to C₆H₄.CH(OH).CH₅ (257°).

Oxim C₆H₅.C(NOH).CH₅. [85°]. from dilute alcohol), converted by AcCl into

C.H. C(NOAc).CH. [72°].

Phenyl hydraside. Colourless prisms.

m-Xylyl methyl ketone [4:2:1] O₆H₄Me₂.CO.CH₂. (228°). Formed from m-xylene, AcCl, and AlCl₂ (Claus, B. 19, 280).

Obtained also by boiling m-xylene with HOAc, ZnCl₂, and POCl₃ (Frey, J. pr. [2] 43, 120). Oil, smelling like peppermint. Not condensed by smelling like peppermint. HClAq. Yields C₁₈H₂Br₃O [69°]. Reduction by HI yields C₂H₂Me₂.CH₂.CH₃ (184°) (Claus, J. pr. [2] 45, 380). HNO_3 gives $C_{20}H_{18}N_2O_4$ [108°] (Claus, J. pr. [2] 41, 492).

p-Xylyl methyl ketone [5:2:1] C₂H₃Me₂CO.CH₃. (225° uncor.). S.G. 12 9962. V.D. 4·93. Formed from p-xylene, AcCl, and AlCl₃ (Claus a. Wollner, B. 18, 1856). Oil, v. sol. alcohol and ether. Oxidised by cold aqueous KMnO, to p-xylyl-glyoxylic acid. Dilute HNO, forms di-methyl-benzoic and methylphthalic acids.

Oxim C₈H₉.C(NOH).CH₂. [58°].

XYLYL METHYL KETONE CARBOXYLIC ACID v. METHYL-BENZOYL-ACETIC ACID.

XYLYL-DI-METHYL-PHO?PHINE

C_eH₃Me₂.PMe₂. (230°). Formed from ZnMe₃ and C_eH₃Me₂.PCl₂ (Czin.atis, B. 15, 2016). Oil. Unites with CS₂, forming a compound [115°].

XYLYL-METHYL-THIOHYDANTOÏII

[1:3:4] $C_6H_3Me_2.N < {CO.CH_2 \atop CS.NH}$ [165°]. Formed from xylyl-thiocarbimide and alanine 'Marckwald, B. 24, 3282). Needles, v. sol. alcohol.

XYLYL PENTADECYL KETONE C_cH₃Me_c.CO.C₁₅H₃₁. [37°]. (269° at 15 mm.). Formed from m-xylene, palmityl chloride, and AlCl₁ (Kraftt, B. 21, 2269). Small plates, yielding di-methyl-benzoic acid [126°] on oxidation.

XYLYL PHENYLAMYL KETONE

C₆H₅Me₂·CO.CHPh.C₄H₅. [91·5°]. Formed by Friedel and Craft's method (Wege, B. 24, 3541). XYLYL-PHENYL- v. PHENYL-XYLYL-.

TRI-XYLYL PHOSPHATE PO(OC,H,Me,),. The o- and p- compounds are formed by heating o- and p-xylenol with POCl, (Kreysler, B. 18, 1702). They are oils, v. sol. ether, alcohol, and benzene.

m-XYLYL-PHTHALIDE

 $C_0H_4 < CO < CO < (C_0H_3Me_2) > O.$ [84°]. Got by heating C₆H₃Me₂.CO.C₆H₄.CO₂H with zinc and HCl (Gresly, A. 234, 237). Needles (from alcohol), sl. sol. benzene.

XYLYL-PROPIONIC ACID

[1:3:4] C₆H₈Me₂·CH₂·CH₂·CO₂H. [105°]. Formed by saponifying the amide. Silky needles, v. sol. hot water.—BaA'₂6aq. Plates, m. sol. water.—CaA'₂4aq.—AgA'. White powder.

Amide C.H.Me, C.H.CO.NH. [107°]. Got by heating the oxim of m-xylyl ethyl ketone with yellow ammonium sulphide under pressure (Claus, J. pr. [2] 46, 477). Needles, sol. hot

water

m-XYLYL PROPYL KETONE C12H16O i.e. [4:2:1] C₆H₂Me₂.CO.Pr. (251°). Formed from m-xylene, butyryl chloride and AlCl, (Claus, (J. pr. [2] 46, 474). Oil, v. sol. alcohol and ether. Yields m-xylyl-glyoxylic ac... [54°] on oxidation with KMnO.

Oxim. Crystals, v. e. sol. alcohol.

p-Xylyl propyl ketone [5:2:1] C.H.Mer.CO.Pr. (249°). Formed in like manner from p-xylene. Light oil.

Oxim [47°]. Colourless needles.

o-Xylyl isopropyl ketene [4:8:1] C.H.Mo. CO.Pr. (258°). Dil, smellingdike turpentine (Claus, J. pr. [2] 46, 484). Yields di-methyl-benzoic acid [163°] on oxidation.

Oxim [68°]. Large prisms. m-Xylyl isopropyl ketone

[4:2:1] C₆H₄Me₂.CO.Pr. (245°). Oil, yielding m-xylyl-glyoxylic acid on oxidation by KMnO₄.

Oxim [97°] Prisms (from alcohol).

Phenyl-hydrazide [129°]. Crystals.
p-Xylyl isopropyl ketone

[5:2:1] C₆H₂Me₂.CO.Pr. (240°). Formed from p-xylene, isobutyryl chloride, and AlCl₄ (Claus, J. pr. [2] 46, 484). Oil, smelling like mushrooms.

Oxim [76°]. Plates and prisms.

TETRA-o-XY'-fL SILICATE Si(OC₆H₃Me₂)₄. (c. 460°). Formed by heating o-xylenol with SiCl₄, the yield being 85 p.c. of the theoretical amount (Hertkorn, B. 18, 1691). Large prisms. Tetra-m-xylvl silicate Si(OC₆H₃Me₂)₄. (453°-

457°). Formed, in like manner, from m-xylenol. Oil.

m-XYLYL THIOCARBIMIDE C₂H₂NS i.e. [4:2:1]C₄H₃Me₂.N:CS. Formed from di-xylylthio-urea and P₂O₅ (Hofmann, B. 9, 1295). Crystals. Converted by PhOEt and AlCl₂ into C₂H₃NH.CS.C₄H₄.OEt [140°] (Gattermann, B. 25, 3530).

XYLYL ETHYL DITHIOCARBONATE
CS(OEt).SC,H,Me, Formed from potassium

xanthate and m-diazo-xylene chloride (Leuckart, J. pr. [2] 41, 192). Oil, converted by heating with alcoholic potash into xylyl mercaptan (214°).

DÍ-XYLYL-THIO-UREA

CS(NH.C₆H₃Me₂[1:2:4])₂. [158°]. Formed by digesting xylidine with CS₂ as long as H₄S escapes (Hofmann, B. 9, 1295). Crystals (from alcohol).

XYLYL-UREA NH₂.CO.NHC₂H₃Me₂[1:2:4]. [186°]. Formed from crude xylidine sulphate and potassium cyanate (Genz, B. 8, 226). Needles (from alcohol).

s-Xylyl ures NH_2 -CO.NH.C₆H₃Me₂[1:3:5].

[162°] (Frentzel, C. C. 1888, 1361)

Di-s-xylyl-urea CO(NHC,H,Me,[1:3:5]), [275°] (F.); [251°] (G. a. C.). Formed from s-xylidine and COCl, (F.). Formed also from m-xylyl cyanate and xylidine (Gattermann a. Cantzler, B. 25, 1089). Needles.

Di-m-xylyl-urea CO(NH.C_eH₂Me₂[1:2:4])₂. [263°]. A product of the distillation of m-xylyl-amido-crotonic acid (Conrad a. Limpach, B. 21,

527). Sublimes when melting.

Di-xylyl-urea CO(NHC₈H₂Me₂)₂. Formed by heating urea (1 pt.) with crude xylidine (3 pts.). Felted needles (from hot alcohol), not melted at 250° (Genz, B. 3, 226).

Reference. - METHYL-BENZYL-UREA.

Y

YTTERBIUM. Yb. At. w. c. 173. This name is given to the metal, not yet isolated, of an earth separated by Marignac, in 1878, from crude erbia obtained from gadolinite or euxenite. The claim of ytterbia to rank as a definite homogeneous compound is not yet satisfactorily established.

The discovery of a new earth in a mineral from Ytterby in Sweden was announced in 1788 by Gadolin; in 1797 Eckeberg confirmed the dis-The new earth was named yttria, and the mineral from which it was obtained was called gadolinite. Researches into the nature of yttria were conducted by Berzelius in 1819 (v. Lehrbuch [5th ed.] 2), Mosander in 1839 and 1843 (J. pr. 30, 27), Scheerer in 1842, Cleve and others. In 1878 Marignac made a careful examination of erbia (A. Ch. [5] 14, 247); he fractionally decomposed erbium nitrate by heat, and obtained an earth which he called ytterbia. The erbia used by Marignac was pink, and gave, in solution, an absorption spectrum; the new earth was white and showed no absorption spec-Nilson (B. 12, 554; 13, 1439 [1879-80]) prepared ytterbia, but found a reacting weight for the earth different from that assigned to it by Marignac. By repeated fractional decomposition by heat of the nitrate of ytterbium, Nilson separated another earth which he called scandia (cf. Scandium, this vol., p. 131). Nilson (B. 13, 1430) in 1879 made a fuller examination of ytterbia, and determined the at. w. of the metal of this earth, and the properties of several salts of the metal. In this memoir Nilson asserted

that erbia was separable into seven distinct earths: erbia proper, scandia, terbia, thulia, ytterbia, yttria, and Soret's X (afterwards called holmia). (cf. Erbium, vol. ii. p. 456; and Metals, rare, vol. iii. p. 245).

Occurrence.—Along with erbia, scandia, yttria, &c., in very small quantities in a few rare Scandinavian minerals, chiefly in gadolinite and euxenite. Nilson (l.c.) obtained c. 20 g. ytterbia from 6-7 kilos. of the crude earths prepared from c. 15

kilos. gadolinite.

Preparation of ytterbia.—Nilson (B. 13, 1430) mixed finely powdered euxenite or gadolinite, c. 400 g. at a time, with four times its weight of KHSO,, fused over a powerful burner, pulverised the fused mass, and thoroughly exhausted with cold water; he ppd. the aqueous solution by ammonia, washed the ppd. hydrated oxides, and dissolved in HNO, Aq; after boiling the solution for some time and filtering, he ppd. by oxalic acid, washed the pp., dried it, and heated strongly until it was decomposed. The crude earths thus obtained were freed from K.CO. by boiling with water, and were then dissolved in HNO, Aq; the solution was evaporated to dryness, and the residue was fused until red vapours begr 1 to come off. The reddish-yellow, opaque solid was then treated with boiling water, whereby a reddish-yellow pp. was formed, which was filtered of by means of a suction-pump. (This pp. contained oxide of Th, along with oxides of Ce, Fe, and U.) The reddish filtrate was eveporated to dryness, and the solid nitrates thus obtained were partly decomposed by heating, the soluble (undecomposed) portion removed by solution in water, the residue dissolved in HNO₂Aq, the solution evaporated to dryness, the residue partly decomposed by heating, the undecomposed nitrates were removed by washing with water, the residue was dissolved in HNO, Aq, and so on. After thirty repetitions of this process a solution was obtained that showed merely a trace of red colour after thirty-five repetitions the solution was quite colourless, and showed only two faint absorption bands, one in the green and the other in the red; after forty repetitions a solution was obtained which gave only the red band; and when the process of partial decomposition by heat had been repeated sixty-eight times a solution in HNO, Aq was obtained which showed no trace of an absorption spectrum. This solution was saturated with H2S, the small yellowishbrown pp. of PtS₂ was filtered off, the filtrate was ppd. by pure oxalic acid, the pp. was washed, dried, and decomposed by heating strongly, and pure ytterbia was thus obtained. For a modification of this method of separating ytterbia from accompanying earths v. Auer von Welsbach (M. 4, 630).

Atomic weight of ytterbium.—The metal has not been isolated; the at. w. was determined by Nilson (B. 13, 1433) by dissolving pure Yb,O, (v. supra) in HNO, Aq, adding a suitable quantity of H,SO,Aq, evaporating on a water-bath to drive off HNOs, then on a sand-bath, and finally over a flame until all free H_2SO_4 was removed, weighing the $Yb_2(SO_4)_s$ thus obtained, and analysing it. Nilson dissolved pure Yb_2O_8 in HNO.Aq, evaporated to dryness, partially decomposed the nitrate by heat, treated with water, &c., as described supra, and in this way obtained seven fractions, each of which was converted into sulphate. The results are presented in the following table :-

Weight of	Weight of sulphate	Petge. co. of sul	Atomio w.	
taken	formed	Yb _a O _a	80,	of Yb.
1.0068	1.6186	62.171	37.829	173-21
1.0139	1.6314	62.149	37.851	173.03
.8509	1.3690	62.155	87.845	178.08
•7371	1.1861	62.145	37.855	173.00
1.0005	1.6099	62.147	37.853	178.01
·8090	1.3022	62.126	87.874	172.84
1.0059	1.6189	62.134	87.866	172.91
Mean 62·147 37·853				173-01

Chemical relations of ytterbium.—If Yb is really a homogeneous substance, it must be placed in Series 10 of Group III. in the periodic classification of the elements. This group contains the following elements:

A comparison of the position of Yb with the

ought to be decidedly more metallic than B, and somewhat more metallic than Al, Ga, or In; the data, however, are too meagre to allow a detailed comparison to be made. Little is known of the properties of compounds of Yb: but what is known broadly confirms the conclusions drawn from the position of the elemen, in the periodic scheme of classification (cf. Earths, METALS OF THE, vol. ii. p. 424).

Ytterbium oxide Yb2O2. (Ytterbiu.) A very heavy, white, infusible powder; slowly acted on by acids when cold or gently warmed, but readily dissolved by dilute acid solutions when boiling. Solutions in acids are colourless, and show no absorption spectra. The emission spectrum of a solution in HClAq, obtained by help of the spark, shows several lines, the most conspicuous of which have the wave-lengths 6221, 556, 5476, 5352, and 5334 (v. Lecoq de Boisbaudran, C. R. 88, 1342). S.G. 9:175. S.H. (0°-100°) ·0646 (Nilson, l.c.).

Hydrated ytterbium oxide. A white, gelatinous, but heavy pp. obtained by adding NH, Aq to a solution of a salt of Yb; shrinks much when dried, and absorbs CO₂ from the air. Easily soluble in acids (Nilson, l.c.).

Ytterbium salts. Nilson (l.c.) prepared an oxalate, an acid selenite, and a sulphate; and

Marignac (C. R. 87, 578) prepared a formate.

Ytterbium oxalate Yb₂(C₂O₄)₃, 10aq. A white,
microscopically crystalline powder; insoluble in water, scarcely soluble in dilute acids; unchanged in air; gives up part of its water at 100°. Prepared by adding oxalic acid to a warm

solution of the sulphate.

Ytterbium selenite Yb2O3.4SeO2.5aq. tained by adding Na, SeO, Aq to a solution of Yb₂(SO₄)₃, washing the ppd. normal selenite, dissolving in water containing a large excess of H,SeO, evaporating nearly to dryness, and washing the residue with water. A white crystalline

salt, insoluble in water. Loses 4H,O at 100°.

Ytterbium sulphate Yb,(SO,),. Prepare d by dissolving Yb,O, in HNO,Aq, adding H,SO,Aq. evaporating on a water-bath to drive off HNO. then on a sand-bath, and finally over a flame at low redness till all H₂SO₄ is removed. An opaque, white solid. Dissolves in water when a large quantity is added at once; if a little is added much heat is produced, the salt combines with the water, and then dissolves very slowly when more water is added. Decomposed at a red heat, fully at a white heat. S.G. 3.793. S.H. ·1039 at 0° to 100°.

A hydrated sulphate Yb. (SO.). Saq was obtained by Nilson in large, white, lustrous prisms by evaporating a solution of Yb,(SO,), in water at a gentle heat.

M. M. P. M.

YTTRIUM. Y. At. w. c. 89.6. The name yttrium is given to the metal of an earth obtained from a few rare Swedish minerals. separation of a new earth from a mineral found at Ytterby, in Sweden, was made by Gadolin in The existence of the new earth was con-1788. firmed in 1797 by Eckeberg, and the earth was called by him yttria. Subsequent researches carried on by Berzelius, Mosander, Scheerer, Berlin, Delafontaine, Popp, Bahr a. Bunsen, Marignac, and others showed that the yttria of positions of elements placed in Series 10 (v. Gadolin and Eckeberg was a mixture of many table on p. 811 of vol. iii.) shows that Yb oxides (cf. Errum, yol. ii. p. 456; Scandium, vol.

YTTRIUM. 881

iv. p. 431; YTTERBIUM, this vol. p. 879; and v. !

infra, Homogeneity of Yttria).

Occurrence.—About 80 to 85 p.c. of yttria, in combination with SiO₂, is found in the exceedingly rare Swedish mineral gadolinite; some specimens of orthite also contain yttria, and the earth has also been found, in combination with Nb₂O₅ and Ta₂C₅, in specimens of euxenite. The oxides that generally accompany yttria in these rare minerals are ceria, didymia, erbia, lanthana,

lime, magnesia, soda, and oxide of iron.

Preparation of yttria.-The crude oxides of yttrium, erbium, &c., are separated from gadolinite by the method of Bahr a. Bunsen described under Erbrum (vol. ii. p. 457). Marignac (A. Ch. [5] 14, 247) converted the oxides into nitrates, heated the nitrates in a Pt basin until red fumes came off freely; treated the residue with water, and separated the solution, which contained much Yt nitrate, from insoluble basic nitrates of Er, &c. He then evaporated the solution to drynes, heated again until partial decomposition was effected, again treated with water, poured off the solution, evaporated it to dryness, partially decomposed the solid nitrates by heat, and so on. After some hundred repetitions of these processes, approximately pure yttria was obtained. Auer von Welsbach (M. 4, 630) separated approximately pure yttria from the crude earths obtained from gadolinite by a long-continued series of operations based on the facts (1) that yttrium nitrate is less readily decomposed to basic salts than erbium nitrate when a fairly conc. solution of these nitrates is heated with the oxides of the metals made into a paste with water, and (2) that basic yttrium nitrates dissolve in a solution of the normal nitrates of Y and Er more readily than basic erbium nitrates.

References.—Gadolin (Crell's Annal. 1796 [1] 313); Eckeberg (Scher. J. 3, 187); Klaproth (Scher. J. 5, 531); Vauquelin (Scher. J. 5, 552); Berzelius (Scher. J. 16, 250, 404; and Lehrbuch [5th ed.], 2, 177); Berlin (P. 43, 105); Scheerer (P. 56, 483); Mosander (P. M. 23, 251); Popp (A. 131, 179); Delafontaine (Ar. Sc. [2] 21, 97; 22, 30; 25, 112; 51, 48; 61, 273); Bahr (A. 135, 376); Bahr a. Bunsen (A. 137, 1); Cleve a. Höglund (Bl. [2] 18, 193, 279); Cleve (Bl. [2]

21, 344).

Preparation of yttrium.—The metal has not been obtained pure. Wöhler in 1828 (P. 13, 580) obtained impure yttrium by reducing the chloride by sodium. In 1864 Popp (A. 131, 179) dissolved yttria in HClAq, added NH,ClAq, evaporated, separated and dried the double chloride of yttrium and ammonium thus formed, mixed this with Na, the double salt and the Na being arranged in alternate layers, heated the mixture in a closed crucible till action began and then allowed the reaction to proceed; washed the fused mass with water, separated unreduced yttria by levigation, washed the dark-grey powder that remained with water, and then with alcohol, and dried it over H₂SO₄. In 1890 Winkler (B. 23, 787) obtained a black powder that decomposed water by heating yttria and finelydivided Mg in the ratio Y₂0₃:8Mg (cf. YTTYA, p. 882). Popp describes yttrium as a dark-grey powder resembling ferrum redactum, decomposing cold water slowly and hot water rapidly, VOL. IV.

easily dissolved by dilute acids, including acetic acid, decomposing boiling KOHAq, and NH, ClAq at the ordinary temperature, with evolution of H and NH,; the metal burns brilliantly when heated on Pt foil; when heated in O it burns with a very dazzling light.

An examination of the emission spectrum of yttrium was made by Thalen: the most prominent lines observed had the wave-lengths 6191, 6131, 5987, 5971, 5662, 5496, 5402, 5205, 5200, 5088, 4900, 4881, 4854, 4374, and 4309 (Kongl. Sw. Vetens. Acad. Handl. 12).

The atomic weight of yttrium was determined by Berzelius, Delafontaine, Popp, Bunsen a. Bahr (v. supra, References) by analyses of the sulphate, or by transforming yttria into the sulphate; the values varied from 92.2 to 102.3. In 1873 Cleve a. Höglund (v. B. 6, 1467) made more accurate determinations by analysing purer specimens of Yt₂(SO₄)₂, and obtained the value

Chemical relations of yttrium.—Assuming yttrium to be a homogeneous element with at. w. c. 89, it must be placed in Series 6 of Group III. in the periodic arrangement of the elements. This group contains the earth metals; yttrium is preceded in the even series family by B and Sc. and is succeeded by La and Yb. A consideration of the position given to Y in the periodic classification shows that the element ought to closely resemble the other metals of the earths. Too little is known of the properties of many of the earth metals to allow of a detailed comparison of them one with another. A general account of the relations of these metals will be found in

EARTHS, METALS OF THE (vol. ii. p. 424).

The homogeneity of yttria.—The properties described as belonging to yttria before 1885 were the properties of a mixture of at least five, and perhaps six, different bodies, according to Crookes C. N. 54, 13, 155). Crookes subjected 'yttria' to a prolonged process of fractionation by ammonia (for a description of this method v. EARTHS, vol. ii. p. 423), and he then examined the phosphorescence spectra of various fractions. Crookes concludes that yttrium is a compound, or perhaps a very intimate mixture, of simpler bodies. For details of the phosphorescence spectra of the substances obtained by fractionating 'yttria' v. METALS, RARE (vol. iii. p. 248). By thirty-two fractionations of 'yttria' by ppn. with NH,Aq, followed by twenty-six fractionations by ppn. with oxalic acid, Lecoq de Boisbaudran (C. R. 103, 649 [1887]) obtained a white specimen of yttria which gave no phosphorescence spectrum when mixed with lime, and which, when converted into chloride, gave a spark spectrum, showing only the lines of yttrium.

Yttrium, bromide of, YBr, 9H,O. Colourless, very deliquescent tablets; obtained by dissolving Y₂O₃ in HBrAq and concentrating (Cleve, Bl. [9] 18, 193). The anhydrous salt is obtained by Duboin (C. R. 107, 99, 243) by heating Y,O, to redness on a support of gas-carbon in a current of CO and Br vapour. Very soluble in

water or alcohol; insoluble in ether.

Yttrium, chloride of, YCl. 6H O. Obtained by dissolving yttria in HClAq, evaporating to dryness on a water-bath, dissolving in alcohol, and evaporating over H2SO, (Cleve, l.c.). Crystallises in large, colourless, very deliquescent,

chombic prisms (C., I.c.). The anhydrous compound is formed by heating yttria to redness on a support of gas-carbon in a slow stream of CO and Cl (Duboin, l.c.). By evaporating a solution of yttria in HClAq with addition of NH Cl a double compound of YCl, and NH Cl is obtained; double compounds with KCl and NaCl are formed by fusing the constituent chlorides together. Popp (A.131, 179) described the double compound YCl₃.3HgCl₂.9H₂O. Cleve (Bl. [2] 31, 195) described YCl. SnCl. 8H2O.

Yttrium, fluoride of, 2YF3.H2O. Obtained as a gelatinous pp. by adding HFAq, or solution of an alkali fluoride, to solution of a salt of yttrium

(Cleve, Bl. [2] 18, 193).

Yttrium, iodide of, (? YI.). Deliquescent crystals, soluble in alcohol; obtained by evaporating a solution of yttria in HIAq (Berlin, P. 43, 105; no analyses given).

Yttrium, oxides of. Besides yttria, Y2O3, there probably exists another oxide containing

more oxygen.

YTTRIA Y₂O₃. (Yttrium sesquioxide.) A hydrate, probably Y₂O₃.6H₂O (Popp, l.c.), is obtained, as a gelatinous pp. closely resembling hydrated alumina, by ppg. the solution of a salt of yttrium by ammonia. The oxide is formed by strongly heating this hydrate; also by decomposing by heat the carbonate, nitrate, or oxalate of yttrium in presence of air. Yttria is described as a heavy, white or almost white powder, S.G. 5-078 (Cleve, l.c.); 5-046 (Nilson a. Pettersson, B. 13, 1459). S.H. (0°-100°) 1026 (N. a. P., l.c.). By heating amorphous yttria with CaCl₂ for some hours, and washing with warm water, Duboin (C. R. 107, 99, 243) obtained highly refractive crystals of Y2O3, scarcely acted on by acids or by fused Na₂CO₃. Yttria is a markedly basic oxide; it absorbs CO. from the air, and decomposes solutions of ammonium salts, giving off NH₂. Yttria is insoluble in water; it dissolves slowly in cold HClAq, HNO, Aq, or H, SO, Aq, but more rapidly on warming. By heating a mixture of yttria and finely-powdered Mg, in the ratio Y, O,:3Mg, Winkler (B. 23, 787) obtained a black powder which slowly gave off H from cold water, rapidly from hot water, and dissolved in dilute HClAq, with violent evolution of H.

YTTRIUM PEROXIDE. By adding H2O2Aq and NH, Aq to a solution of sulphate or nitrate of yttrium, Cleve (Bl. [2] 43, 53) obtained a white gelatinous pp. to which he gave the composition Y₄O₉; but this formula cannot be regarded as

Yttrium, salts of. The salts which are formed by replacing the H of acids by yttrium belong to the form Y_2X_2 , where $X = 2ClO_2$, $2NO_2$, SO₃, SO₄, CO₃, ²PO₄, &c. Almost all the yttrium salts of oxyacids that have been prepared are normal salts; a basic nitrate, and an acid orthophosphate and selenate, are known. Many of the salts are obtained by diss lving yttria in acids and evaporating; those salts which are insoluble, or but slightly soluble, in water are formed by ppn. from the nitrate or sulphate.
The salts formed by ppn. are the arsenate, borate, carbonate, chromate, iodate and periodate, oxalate, phosphates, selenite and sulphite; besides these salts, the following salts of oxyacids are known, and are soluble in water: bromate, chlorate and perchlorate, nitrate, selenate, sulphate, thiosulphate and tungstate. A few double salts are known; the principal are $Y_2(CO_3)_3.M_2CO_3.xH_2O$, $Y_2(C_2O_4)_3.3M_2C_2O_4.xH_2O$, and

 $Y_2(SO_4)_*mM_2SO_4xH_2O$, where M=an alkali metal. Duboin (C.R. 107, 99, 243) obtained a silicate Y_2O_4 .SiO₂ by very strongly heating a mixture of 3 pts. yttria with 30 pts. CaCl₂, and washing with water. The yttrium salts of oxyacids have been examined, chiefly by Cleve (Bl. [2] 18, 193, 289; 21, 344; v. also Berlin, P. 43, 105; Högbom, Bl. [2] 42, 2; Popp, A.

131, 179).

Yttrium, sulphide of. No compound of yttrium and sulphur has been isolated with certainty. Addition of (NH4)2SAq or NH4HSAq to a solution of a salt of yttrium ppts. hydrated oxide. By fusing yttria, S, and an alkali carbonate, and washing the fused mass with water, a yellowish-green solid is obtained that is insoluble in water, but is partly decomposed thereby to hydrated oxide and H_2S ; this solid dissolves in acids, giving off H_2S . The same solid seems to be obtained by strongly heating yttria in H laden with CS. No compound of yttrium and S was obtained by heating yttria with Na2S. By heating YCl, in a stream of H2S, HCl is given off, and a yellow powder remains, which is decomposed by water, with evolution of much H₂S (v. Popp, A. 131, 179). According to Popp (l.c.), the composition of this substance is Y2S3.

By passing dry H2S over a mixture of YCl, with excess of NaCl, heated to c. 1000° in a boat of gas-carbon, Duboin (C. R. 107, 99, 243) obtained a greenish, crystalline solid, which when washed with water left transparent, greenish lamelle of the double compound M. M. P. M.

Y.S.Na.S.

Z

ZANZALOÏN v. ALOÏN. ZEORIN C, H2O. [231°]. Occurs in the ethereal extract from Zeora sordida (Paterno, G. 7, 281, 508). Small pyramids with hexagonal base (from ether-alcohol), v. sl. sol. alcohol and ether, insol. water. Neutral. Not attacked by alkalis or dilute acids.

ZINC. Zn. At. w. c. 65.3 (v. infra). Mol. w. c. 65.3 at c. 1400°; probably the same in solution in Hg (v. infra). Melts at 419° (determined by a.r thermometer; V. Meyer a. Riudle, B. 26, 2443); at 417.57° (determined by platinum thermometer, Callendar a. Griffiths, C. N. 63, 1); for older determinations, giving

m.p. from 840° to 450°, v. Carnelley's Meltingand Boiling-point Tables (1, 14). Boils between 980° and 954° (Deville a. Troost, C. R. 90, 773); at 930° (Violle, C. R. 94, 720; cf. Troost, C. R. 94, 788). S.G. c. 6.9 to 7.2 (v. Schiff, A. 107, 59; Kalischer, B. 14, 2750; Quincke, P. 135, 642; Spring, B. 16, 2724). V.D. 34·3 at c. 1400° (Mensohing a. v. Meyer, B. 19, 3295). S.H. (0° to 100°) ·0935 (Bunsen, P. 141, 1); (19° to 47°) ·0932 (Kopp, T. 155, 71; cf. Schüller a. Wartha, B. 8, 1016). C.E.; if length of bar at $0^{\circ} = l^{\circ}$, then length at $t^{\circ} = (1 + \beta t)$; if volume at $0^{\circ} = \nabla_{0}$, then volume at $t^{\circ} = (1 + 3\beta t)$; $\beta =$ *00002976 from 0° to 100° (Matthiessen, P. M. [4] 32, 472). Heat of fusion for 1 kilo = 28.130 (Person, A. Ch. [3] 24, 129). T.C. (Ag = 100)28·1 (Wiedemann, P. M. [4] 19, 243). E.C. (Hg at 0° = 1) 16·1 at 0°, 16·92 at 100° (Bénoit, P. M. [4] 45, 314); 14-83 at 15° (Kirchoff a. Hausemann, W. 13, 406). Chief lines in emission spectrum 6360, 6100, 4924, 4911, 4809, 4722, 4679 (Huggins, 1. 154, 139; cf. Thálen, Détermin. des Lonqueurs d'Onde des Raies Métalliques [Upsala, 1868]; and Ames, P. M. [5] 30, 33); for wave-lengths of lines of high refrangi-bility v. Hartley a. Adeney (T. 1884 [1]. 97). Zinc crystallises in tabular hexagonal plates, a:c=1.356425 (Williams a. Burton, Am. 11, 219; cf. Stolba, J. pr. 89, 122; 96, 178; Sharples, Am. S. [3] 7, 223; Nöggerath, P. 39, 324; Nicklès, A. Ch. [3] 22, 37; G. Rose, B. B. 1852. 26; P. 107, 448; Storer, P. Am. A. 6; Cooke, Am. S. 81, 191). The metal is probably dimorphous. H.C. [Zn,O] = 85,430 (Thomsen, Th. 3, 275). Refraction equivalent = 9.8 (Gladstone, Pr. 18, 49).

Historical.—That copper acquired a reddish colour, and that its properties were modified, by melting with certain ores, was known to the ancients and the alchemists; the ore with which copper was generally melted to form brass was called cadmia fossilis. The fact that a lustrous solid could be obtained by heating cadmia fossilis seems to have been known to the ancient writers on mineralogy. According to Kopp (Geschichte der Chemie, 4, 116), the earliest use of the word sinc is found in a writing of the fifteenth century attributed to Basil Valentine. Paracelsus, in the sixteenth century, speaks of zinc as a definite metal-like substance, and assigns it to the class of bastard or semi-metals. The name ' zinc' was applied during the seventeenth and eighteenth centuries alike to zinc ores and to the metal-like subscance obtained from these ores. Boyle speaks of zinc, and also uses the word spelter, or spiauter, a term of Indian origin according to Kopp. Tolerably pure zinc seems to have been obtained from zinc ores about 1720; probably by Henckel (v. Percy's Metallurgy, 1, 520).

Occurrence. — Zinc is found in small quantities; Becker (J. M. 1857. 698), and Phipon (P. 8. 55, 218) noticed the occurrence of native zinc in Victoria. Considerable quantities of compounds of zinc occur in many places; the commonest ores are calamine (carbonate), siliceous calamine (silicate), blende (sulphide), and red zinc ore (oxide); aluminate, arsenat, phosphate, and sulphate of zinc are also found, but in smaller quantities. Small quantities of Cd compounds occur in most zinc ores. The

ash of a plant that grows on the waste heaps of the zinc works in Rhenish Prussia (Viola calaminaria) is said to contain compounds of zinc (Braun, P. 92, 175). Zinc compounds have been found in some springs (v. Hillebrand, U. S. Geolog. Survey Bull. No. 113 [1893]). Traces of zinc compounds have been found, according to Lechartier a. Bellamy (C. R. 84, 687), in the human liver, in calves' liver, in beef, hens' eggs, wheat, barley, maize, beans, and vetches.

Formation.—Zinc ore, generally carbonate or sulphide, is roasted in reverberatory calciners; the roasted ore is mixed with half its weight of powdered charcoal, coke, or anthracite, and heated in crucibles or retorts arranged so that the reduced metal distils from the impurities; the metal is condensed and collected in suitable vessels, and CO passes off. For details and descriptions of the different forms of apparatus v. DICTIONARY OF APPLIED CHEMISTRY, vol. iii. p. 1042.

Preparation.—Commercial zinc generally contains from 5 to 3.3 p.c. lead, with c. 2 to 5 p.c. iron and cadmium, and small traces of arsenic.

Pure zinc was prepared by Reynolds a. Ramsay (C. J. 51, 854 [1887]) by dissolving zinc sold as free from common impurities in dilute H₂SO₄Aq, evaporating, crystallising the sulphate several times, electrolysing a solution of this salt, the metal being deposited on an electrode of Pt wire, dissolving the metal in dilute HClAq, ppg. ZnO₂H₂ by NH₃Aq, dissolving the pp. in excess of NH₃Aq, and ppg. ZnS by H2S; the ZnS was dissolved in HClAq, and again ppd. from an ammoniacal solution by H2S; the pp. was well washed, dissolved in pure dilute H2SO4Aq, and the sulphate obtained on crystallisation was re-crystallised several times; an aqueous solution of this sulphate was then electrolysed, using a Pt wire as electrode, and the metal was melted in a tube of hard glass from which the air had been exhausted. By distilling this metal in vacuo in a tube of hard glass, very pure zinc was obtained. Pullinger (C. J. 57, 816 [1890]) distilled zinc sold as 'chemically pure' in a hard glass tube, bent slightly towards the closed end, placed in a combustion furnace, the open end being connected with a good water-pump. By melting the distilled zinc in a tube of hard glass with a small bulb blown on the end, the part of the tube between the bulb and the wider portion containing the zinc being capillary, and the open end being connected with a water-pump, he obtained the zinc in perfectly bright, smooth, lustrous spheres free from hollows; the pump was stopped while the zinc was molten, and the increased pressure caused the molten metal to filter through the capillary tube into the bulb. For the preparation of pure zinc by electrolysis of an ammoniacal solution of the sulphate, v.

Myers (C. R. 74, 195).
Stolba (C. C. 1884. 419) says that zinc can be obtained free from As and nearly free from iron by the combined interaction of S and water vapour. He mixes burnt gypsum with one-fourth its weight of S powder, moulds the moistened mixture into spheres about 5 cm. diameter, and sinks these to the bottom of a crucible containing molten zinc; vapours of

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H2S and S are given off, and the molten metal is thereby briskly agitated. When the reaction is completed, the little balls are taken out, the upper crust is removed, and the operation is

repeated if necessary.

Zinc-dust is obtained either in the distillation of the metal, or, in not such fine division, by crushing and powdering the metal in an iron mortar at a temperature somewhat below the m.p. of zinc. The ZnO₂H₂ always present in specimens of zinc-dust may be removed by digesting with NH, ClAq and then with NH, Aq, and drying on a porous tile in vacuo (Carnegie, C. J. 58, 471).

Properties.—Zino is a white metal, with a slight shade of blue; it is very lustrous when polished. Commercial zinc is brittle at the ordinary temperature, but it becomes malleable between 100° and 150°; at 210° it again becomes brittle, and at that temperature it can be finely powdered in a mortar. Pure zinc is said to be malleable at the ordinary temperature. mercial zinc that has been heated to 100°-150° retains its malleability when it cools. Kahlischer (B. 14, 2747) noticed that rolled zinc ceased to give a ringing sound when struck after it was heated to c. 160°-300°, that it could then be bent easily, and that when bent it emitted a sound like the 'cry' of tin; he found that the zinc became crystalline, and the crystalline structure was more decided the higher the temperature; the S.G. was very slightly increased, and the electrical resistance was reduced by c. 8 p.c. a softer metal than copper; its hardness is increased by rolling. The tensile strength of zinc varies much with the mode of preparation. Zinc crystallises easily, especially when pure; the form is that of hexagonal plates (for references v. supra, beginning of this article). Zinc melts at c. 420°, and boils somewhat below 1000° Molten zinc expands somewhat (v. supra). during solidification. The metal can be distilled at a full red heat; the vapour takes fire in the sir, and burns with formation of ZnO and the production of a bright white light. Zinc exposed to the ordinary air becomes very gradually covered with a thin film of a basic carbonate which protects the mass of the metal from further corrosion; in pure dry air zinc is unchanged. Zinc dissolves in dilute acids, generally with evolution of H (v. infra, Reactions with sulphuric and nitric acids); it also dissolves in solutions of caustic potash or soda. When impure zinc is amalgamated it resists the action of acids. Zinc is not acted on by mineral lubricating oils; some of the commoner animal oils attack it slightly (v. Redwood, C. S. I. 5, 862). Zinc ppts. most of the other metals from their solutions; if a piece of impure zinc is dissolved in an acid, most of the metallic impurities (As, Cd, Cu, Pb, &c.) remain undissolved as long as undissolved zinc is present. Zinc combines directly with O, S, Se, Te, the halogens, and P; it forms alloys with many metals. Pure zinc does not decompose boiling water, but the commercial metal reacts and gives off H. Zinc-dust usually contains some ZnO₂H₂; it is said also to contain H, produced by the interaction of Zn and ZnO₂H₂ (v. Greville Williams, C. N. 52, 205, 268).

Stomic and molecular weights of sinc. In 1809 Gay-Lussac (Mém. S. d'A., 2, 174) deter-

mined the weight of ZnO obtained by dissolving a determinate weight of zinc in nitric acid, evaporating to dryness, and strongly heating the residue; Berzelius in 1811 repeated the experiment of Gay-Lussac (G. A. 87, 400; v. also P. 8, 184); and in 1843 Erdmann made similar experiments (v. Berzelius' Lehrbuch, 3, 1219). The values obtained for the at. w. or zinc were 65:39 (G.-L.), 65.41 (B.), and 64.94 and 64.88 (E.). In 1844 Favre decomposed ZnC₂O₄ by heating in air, and determined the weights of ZnO and CO. produced; the value 65.85 was thus obtained for the at. w. of zinc. In the same year Favre obtained the value 65.78 by dissolving zinc in dilute H2SO,Aq, burning the H given off by passing it over hot CuO, and weighing the water thus produced (A. Ch. [3] 10, 163). In 1884 Marignac determined the Cl and the Zn in ZnCl₂.2KCl, and obtained the value 65.18 for the at. w. of zinc (A. Ch. [6] 1, 509; v. also Baubigny, C. R. 97, 908 [1883]). By dissolving zinc in dilute H2SO Aq and measuring the H given off, van der Plaats concluded that the at. w. of zinc is 65·18 (C. R. 100, 52 [1885]). Reynolds a. Ramsay, in 1887, by measuring the H given off by the interaction between very pure zinc and H2SO, Aq, obtained the value 65.43 (C. J. 51, 854). By oxidising zine to ZnO, by HNO₂Aq, in 1888, Morse a. Burton obtained the 51, 854). value 65.1 (Am. 10, 811). In 1889 Gladstone a. Hibbert (C. J. 55, 443) determined the ratio of zinc dissolved to silver deposited by one and the same electric current; taking the at. w. of silver as 107.66 (the value adopted in this Dictionary) the at. w. of zinc was found to be 65.29.

In 1886 Mensching a. V. Meyer (B. 19, 8295) found the V.D. of zinc at c. 1400° to be 34.3, a number which shows that the gaseous molecule of zine is monatomic. Ramsay, in 1889, determined the lowering of the vapour pressure of Hg produced by dissolving zinc therein (C. J. 55, 521); assuming that equa' volumes of dilute solutions contain equal numbers of molecules, and that the molecular weight of liquid Hg is 200, Ramsay's results indicate that the molecules of zinc in dilute solutions of this metal in Hg are monatomic. The experiments of Heycock a. Neville (C. J. 57, 376 [1890], and 61, 888 [1892]) on the lowering of the freezingpoints of Bi, Cd, Pb, Sn, and Na by solution of zinc in these metals tend to confirm the conclusion that the molecule of zinc in dilute solu-

tions in these metals is monatomic.

Zinc is the Chemical relations of zinc. second odd-series member of Group II. in the periodic classification of the elements. Zinc is preceded in the odd series of this group by Mg and is followed by Cd, —, and Hg; the members of the even series of Group II. are Be, Ca, Sr, -, -. Zinc is the second member of series Ba, -5; it is preceded in this series by Cu, and is succeeded by Ga, Ge, As, Se, and Br. The general chemical character of zin- in that of a metal; its oxide ZnO is basic, its chloride ZnCl. is volatilisable without decomposition. No acids are known containing zinc, and the molecule of the element is monatomic. The fact that comfounds of ZnO and KoO are obtained by dis-solving ZnO,H, in KOHAq and adding alcohol shows that the hydroxide of the metal has feebly marked acidic properties, and the isola-

tion of a number of oxychlorides and basic carbonates, nitrates, and sulphates illustrates the fact that zinc is less positive than Mg, which is the first member of the zinc family of Group II. For a general account of the properties of the family of which zinc forms a member v. MAGNESIUM GR UP OF ELEMENTS, vol. iii. p. 163. The atom of zinc is divalent in the gaseous molecules of its compounds.

Reactions and Combinations .- 1. Zinc is unchanged in pure dry air; in ordinary air the surface layers are slowly converted into a basic carbonate, which protects the metal beneath. Heated strongly in air or oxygen, zinc burns to ZnO.-2. Heating zinc in bromine, chlorine, or iodine produces ZnBr2, ZnCl2, or ZnI2-3. When zinc is heated in phosphorus vapour, phosphide of zinc (q. v., p. 890) is formed.—4. By heating zinc with selenim vapour, a compound of Zn and Se is formed (v. ZINC SELENIDE, p. 890) .-5. Zinc combines with tellurium when the elements are heated together (v. ZING TELLURIDE, p. 891).-6. Sulphur and zinc combine when a mixture of these elements is very strongly compressed (Spring, B. 16, 1000).-7. Zinc decomposes water at a red heat. Commercial samples of zinc decompose water at 100° (L'Hôte, C. R. 101, 1153).—8. ZnS is formed by strongly heating zinc in a stream of hydrogen sulphide; when hydrogen selenide is used the product is ZnSe, and by heating the metal in hydrogen telluride ZnTe is formed (v. Margottet, C. R. 84,1293).— 9. Zinc is said to reduce carbon monoxide at a very high temperature.—10. Zinc is superficially oxidised by heating to dull redness in nitric oxide (Sabatier a. Senderens, C. R. 114, 1429). Oxidation in nitrogen dioxide (NO₂) is effected at 300° (S. a. S., C. R. 115, 236).—11. Many metallic oxides are reduced by heating with zinc.-12. Zinc reacts with sulphurous acid solution; according to Schweizer (v. C. N. 23, 293), the products are ZnSO, along with H,S,O,Aq, H,S,O,Aq, and S.—13. Zinc dissolves in hydrochloric acid, giving off H and forming ZnCl₂. Reynolds a. Ramsay (C. J. 51, 857) say that pure zinc scarcely reacts with boiling hydrochloric acid. For reactions of zinc with sulphuric and nitric acids, v. infra.-14. Zinc dissolves slowly in hot solutions of caustic soda or potash, giving off H, and forming ZnO.xM₂O (cf. Zinc hydroxide, p. 888); the reaction is much hastened by adding iron or Pt along with the zinc.-15. Many metallic salts in solution are reduced by zinc, with ppn. of the metals, e.g. salts of As, Cd, Cu, and Pb.-16. When zincdust is shaken with an aqueous solution of ferric chloride, the whole of the ferric salt is very rapidly reduced to ferrous chloride (for details of the application of this reaction in the estimation of ferric iron, v. Carnegie, C. J. 53, 468 [1888]).—17. According to Siersch (J. 1867. 257), zinc dissolves in sodium chloride solution, forming InCl₂.2NaClAq, Zn_U, and H.—18. Poumarède (J. pr. 73, 496) says that nearly all chlorides and fluorides are reduced by reacting with vapour of zinc in an atmosphere of H .-19. By Leating to redness a mixture of sino filings and sodium metaphosphate, Hvoslef (A. 100, 99) probably obtained phosphides of sinc (v. Zinc PROSPHIDES, p. 890).—20. Zinc reacus with socium hydrogen sulphits solution, forming ZnSO₃, Na₂S₂O₄Aq, and Na₂SO₂Aq (v. HYPOSULPHITES, this vol. p. 592) .- 21. By heating zinc with sinc sulphide in an exhausted tube, some of the ZnS is carried forward and deposited in the front part of the tube; probably at a high temperature ZnS is decomposed, and there exist only Zn and S, and at a lower temperature these recombine to form ZnS (Morse a. White, Am. 11, 848). A similar phenomenon is noticed on heating zine with sinc oxide (M. a. W., l.c. p. 258).—22. ZnS is formed by strongly heating zine and mercuric sulphide or zine and potassium polysulphides (v. ZINC SULPHIDE, p. 890).

When slips of zinc-foil are immersed in CuSO,Aq, a deposit of finely-divided Cu is formed on the zinc. This copper-zinc couple is an energetic reducing agent; KClO,Aq is reduced to KClAq, KNO,Aq to KNO,Aq and NH,Aq, K, FeCy, Aq to K, FeCy, Aq, SO, Aq to S, As, O, Aq to AsH₃, C₆H₅NO₂ (in alcohol) to C₆H₅NH₂, &c. (v. Gladstone a. Tribe, C. J. 33, 306).

Reactions of zinc with sulphuric and nitric acids. Commercial zinc dissolves easily in dilute H2SO4Aq, with formation of ZnSO4Aq and evolution of H. In 1830 De la Rive noticed that very little action took place between approximately pure zinc and H₂SO₄Aq; this fact has been confirmed by other experimenters. Reynolds a. Ramsay (C. J. 51, 857 [1887]) found that zinc prepared by repeated electrolysis of the sulphate and distillation in vacuo scarcely reacted with H₂SO₄Aq; L'Hôte (C. R. 101, 1153) also asserted that pure zinc does not react with H_2SO_4Aq . Divers a. Shimidzu (C. J. 47, 598) [1885]) observed very great differences between the rate of action of the same H2SO, Aq on sheets of commercial zinc of the same size. Pullinger in 1890 (C. J. 57, 815), and Weeren in 1891 (B. 24, 1785), made somewhat elaborate investigations into the connections between the conditions and the rate of the interaction of zine and H.SO.Aq.

Pullinger used 'pure' sulphuric acid diluted with three times its weight of water; and zinc prepared by distilling that sold as 'chemically pure' in vacuo and casting in balls under pressure to prevent the formation of cracks or hollows (v. supra, Preparation of sinc, p. 885). He found that when the surface of the zinc was very smooth-this was best accomplished by immersing in aqua regia for 10 or 15 seconds, and washing with water-and the acid was boiled for some hours before the experiment, there was practically no reaction at 20°-25° (spheres of zinc weighing c. 2\frac{1}{2} g. lost from 5 to 6 mgms. in 20 hours). Pullinger found that the presence of small quantities of H2S, SO, or H.S.O.Aq did not affect the weight of zinc dissolved by the boiled H2SO, Aq; addition of some H₂O₂ materially increased the rate of action; and H2SO, Aq that had been electrolysed before use dissolved from four to ten times more zinc than some of the same acid that had not been electrolysed; addition of a few drops of HNO, Aq considerably increased the action; when a tew drops of HIAq were added to the H₂SO₄Aq prac-tically no action occurred. P. supposed that all the 'purp' acid used by him contained traces of an oxidising substance, probably H.S. .. and that the solvent action was due to this. P.

insists on the important connection between the rate of action and the smoothness or roughness of the surface of the zinc used. But he concludes that 'in all probability pure dilute sulphuric acid would, at ordinary temperatures, be entirely without action upon metallic zinc, whether the surface of the latter were rough or smooth.'

Weeren used zinc which he found to be chemically pure by analysis; he gives no account of his method of preparation nor any details of his analytical results. W. found that the weight of pure zinc dissolved by pure H.SO.Aq (1:20) at the ordinary temperature, and under a pressure of 10 mm., was c. ten times greater than the weight dissolved at 760 mm. pressure; but the weights of impure zinc dissolved at the two pressures were almost the same. The pure zinc used by W. dissolved fairly rapidly in boiling H.SO.Aq; in one case when 2.1 mgm. dissolved at 0° after thirty minutes' action, 122 mgm. dissolved in the boiling acid. W. found that the solubility of his pure zinc increased very slowly as temperature rose up to within 1° of the b.p. of the acid used, but that when ebullition actually began the solubility of the zinc suddenly increased; the weight of pure zinc dissolved by H₂SO₄Aq at 100° and boiling, was c. sixteen times greater than the weight of the same zinc dissolved by the same acid at 100° but kept from actually boiling by pressure. On the other hand. W. noticed that as much impure zinc dissolved at 100° when the acid was boiling as when it was prevented by pressure from boiling. Addition of oxidisers, CrO, or H₂O₂, enormously increased (c. 300 times) the solubility of pure zinc in H.SO, Aq at 18°-20°, but only slightly increased (c. six times) the solubility of impure zinc. W. supposes that when pure zinc is immersed in pure H2SO, Aq a slight reaction occurs, and that the H produced is attracted to and held firmly on the surface of the zinc, and that the reaction ceases because the surface of the metal is protected by the layer of H. Increase of temperature does not appreciably affect the rate of action, because the layer of H remains fixed to the surface of the zinc, but when the acid boils the H is removed and rapid dissolution occurs; any conditions which remove the layer of H increase the solubility of the zinc. W. supposes that the rate of dissolution of impure zinc in H2SO, Aq is not much affected by boiling, presence of oxidisers, &c., because H is not given off at the surface of the zinc, but at the surface of the more negative The rapid solution of zinc in impurities. HNO, Aq is connected, according to W., with the rapid exidation of the H produced at the surface of the zinc, and hence the production of a layer of water on the surface of the metal which destroys the attraction between the zinc and any H that may escape oxidation.

As regards the products of the interaction of sino and sulphurio acid, when the acid is thirly concentrated (c. H₂SO₄, H₂O to c. H₂SO₄) SO₂ is given off, and at moderately high temperatures H₂S and S are also produced (v. Calve t a. Johnson, C. J. [2] 4, 435 [1867]). A qualitative examination of the products by Pattison Muir a. Adie (U. J. 58, 47 [1888]) showed that ZnSO₄ is the only sait of zinc produced, with any concen-

tration of soid, and at any temperature up to the b.p. of the soid used; that with approximately pure zinc little or no SO₂ or H₂S is formed unless the temperature be high; and that the purer the zinc the less is the quantity of S produced (cf. Ditte, A. Ch. [6] 19, 68).

The products of the interaction of zine and nitric acid are Zn(NO₃)₂Aq, Zn(NO₂)₂Aq, NH₄NO₃Aq and NH₄NO₂Aq, and NO, N₂O, and N. Acworth found that when HNC, Aq reacts with zinc in presence of (NH4)NO3 the chief gaseous product is N (along with some N₂O and NO) (C. J. 28, 828 [1875]). Acworth a Armstrong studied the reaction of HNO, Aq and Zn more fully in 1878 (C. J. 32, 54), and found that the only gaseous products were N, N₂O, and NO (cf. Deville, C. R. 70, 22, 550 [1870]). According to Divers (C. J. 43, 443 [1883]), a little NF₂OH is formed when cold HNO₂Aq (8 to 4 p.c.) is poured on to granulated zinc and quickly poured off again. Divers a. Sh'midzu (C. J. 47, 597 [1885]) found that considerable quantities of NH₂OH are produced by the combined reaction of HNO₃Aq and H₂SO₄Aq, or HNO₃Aq and HClAq, on zinc (v. Hydroxylamine, Formation, vol. ii. p. 734). Divers (l.c.) thought that zinc does not form Zn(NO2)2 by a direct reaction with HNO, Aq, but by interacting with some of the Zn(NO₃)₂ formed by the primary action of the metal on the acid. The products of the interaction of zinc and HNOsAq have been examined recently by Montemartini (G. 22[1], 277[1892]). who says that free H is not produced (he also says that no NH2OH is formed), and that in addition to nitrate and nitrite of zine and ammonium, hyponitrite is also produced; the gaseous products, according to M., are N.O, NO, NO2, and N; there is no nitrous acid formed, says M., if the HNO, Aq contains more than 30 p.c. HNO, and no NO if less than 30 p.c. HNO is present. M. also gives determinations of the quantities of the various products at different temperatures, and of the rate of reaction with different concentrations of HNO, Aq (Abstract in C. J. 62, 1279 [1892]).
Zinc, alloys of. Alloys of zinc have been

formed with most metals by fusing the constituents together; the alloys are generally hard, some of them are brittle. With antimony, two crystalline alloys are formed by melting the metals together, in the ratios 8Zn:2Sb and Zn:Sb, and allowing to cool very slowly; Cooke (Am. S. [2] 18, 229; 20, 222) for nulates these alloys as Zn, Sb, and Zn, Sb, respectively. These alloys react with water and give off H. Alloys with arsenic are formed by heating together Zn and As, or Zn and As,O₃, or by heating Zn in vapour of As and H; by the last method Descamps (C. R. 86, 1022, 1065) obtained crystals agreeing in composition with the formula Zn_sAs_2 . Spring (B. 16, 324) formed an alloy by compressing Zn and As, in the ratio 3Zn:2As, at 6,500 atmospheres. Zinc alloys with oismuth when the metals are melted together, but on cooling two layers are formed, the upper containing zinc with c. 2 p.c. Bi, and the lower Bi with from 8 to 14 p.c. zinc. An alloy of zinc with bismuth and lead (3 pts. Zn, 5 pts. Bi, and 5 pts. Pb) melts at 94.5°. For descriptions of alloys with calcium v. Caron (C. R. 48, 440; 50, 547); Wöhler (A. 138, 253); and Norton a. Twitchell

(Am. J. 10, 70). Brass is composed of alloys of zinc with copper (v. vol. ii. p. 254; and for details v. DICTIONARY OF APPLIED CHEMISTRY, vol. iii. p. 1052). Alloys of zinc with copper and nickel form German silver; most modern bronzes are alloys of zinc with copper and tin (v. vol. ii. p. 254; and for details v. Dictionary OF APPLIED CHEMISTRY, vol. iii. p. 838). Zinc alloys with iron (for details v. DICTIONARY OF APPLIED CHEMISTRY, vol. iii. p. 1051). For alloys with lead v. vol. iii. p. 125 (also Krant, S. C. I. 5, 537). Alloys with lead and tin are mentioned in vol. iii. p. 125 (v. also Wright a. Thompson, Pr. 48, 25). For alloys with magnesium v. Parkinson (J. pr. 101, 375). Zinc and mercury form a number of amalgams, which are brittle when a little Hg is present and pasty when much Hg is present (for references v. S. C. I. 9, 512). For alloys with platinum v. this vol. p. 288. For alloys with silver v. Wright a. Thompson (Pr. 48, 25),

For an account of the formation of zinc alloys by immersing zinc in various metallic solutions v. Mylius a. Fromm (B. 27, 630

[1894]).

Zino, amide of, Zn(NH₂)₂. This compound was obtained by Frankland (Pr. 8, 502) by passing dry NH₂ into an ethereal solution of ZnEt₂. A white, amorphous powder; unchanged at 200°, but decomposed at a red heat to Zn₂N₂ (v. Zino nitraide, p. 889) and NH₂; reacts with water to form ZnO₂H₂ and NH₃.

Zinc, bromide of, ZnBr₂. Formula probably molecular, from analogy of ZnCl₂. S.G. 3·643 at 10° (Bödeker). Melts at 39.4° (Carnelley, C. J. 33, 277). Boils at 695° to 699° (C. a. Williams, C. J. 33, 283); at 650° (Freyer a. V. Meyer, B. 25, 622). H.F. [Zn,Br²] = 75,930 (Th. 3, 275).

Preparation.—1. Zinc filings are heated to dull redness in a stream of Br vapour.—2. Zinc, or ZnO₂H₂, is dissolved in a slight excess of HBrAq, the solution is evaporated to dryness, and the dry residue is sublimed out of contact with air.

Properties and Reactions.—White needles; very deliquescent. Easily soluble in water, alcohol, or ether (v. Berthemot, J. Ph. 14, 610). Thomsen gives [ZnBr₂,Aq] = 15,030 (Th. 3, 275). Aqueous solutions of ZnBr₂ of different concentrations have the following specific gravities according to Kremers (P. 108, 115):—

8.G. ZnBr.Aq at 19.5%	P.c.ZnBr.
1.1849	18.3
1.3519	81.7
1.5276	43.2
1.7082	52.6
1.8525	59.1
2.1027	68.0

Combinations.—1. With ammonia to form a series of compounds. These compounds were examined by Rammelsberg (P. 55, 240), and, more received, by André (Bl. [2] 39, 398 [1883]). By dissolving ZnBr, in NH, Aq under different conditions, A. obtained (1) ZnBr, 2NH, H2O, (2) 3ZnBr, 8NH, 2H2O, (3) 3ZnBr, 10NH, H2O, and (4) ZnBr, 5NH, ; by satura-ing hot NH, BrAq with ZnO, he obtained 3ZnBr, 6NH, H2O.—2. With casium bromide, to form ZnBr, 2CsBr and ZnBr, 2CsBr (Wells a. Campbell, Zeit. f. caurg. Chemie, 5, 273).

Zino, chloride of, ZnCl₂. Mol. w. c. 138. Melts at 262° (Braun, P. 154, 190). Boils at 708°-719° (Carnelley a. Williams, C. J. 33, 284); at 730° (Freyer a. V. Meyer, B. 25, 622). S.G. 2·758 at 13° (Bödeker). V.D. at 890° to 907° = 66 (F. a. M., l.c.). H.F. [Zn,Cl]=97,210 (Th. 3, 275).

Formation.—1. By heating zinc filings in Cl.—2. By heating a mixture of 2 pts. HgCl, and 1 pt. zinc filings in a retort.—8. By distilling a mixture of equal pts. ZnSO, and CaCl,, or a mixture of 1 pt. ZnO and 2 pts. NH₄Cl.—4. By dissolving zinc, or ZnO₂H₂, in HClAq, evaporating to dryness, and heating the residue in a retort.

Preparation.—1. Zinc filings are heated in a small retort in a stream of Cl; the product is distilled.—2. Zinc, or ZnO₂H₂, is dissolved in a slight excess of HClAq; the solution is evaporated to dryness, and the residue is distilled from a small retort.

Properties and Reactions.—A white, semi-transparent, crystalline, very eaustic solid; very deliquescent. Easily soluble in water, also in alcohol. Thomsen gives [ZnCl²,Aq]=15,630 (Th. 3, 275). Sublimes at a red heat. An aqueous solution is partly decomposed on evaporation, giving oxychlorides (q.v.). Oxychlorides are also formed by heating a solution of ZnCl₂ in HClAq with ZnO, also with HgO or PbO. Kremers (P. 105, 360) gives the following data;

S.G. ZnCl,Aq	P.c. ZnOl
1.1275	13.8
1.2466	25.8
1.3869	87.5
1.5551	49.2

A conc. solution of ZnCl₂ is often used as a bath for maintaining fairly high constant temperatures. Dry ZnCl₂ is used as a caustic; also as a dehydrating agent; a dilute aqueous solution is employed as an antiseptic.

Combinations.—1. With water. By keeping ZnCl₂Aq containing 70.5 p.c. ZnCl₂ at 0.5 for 24 hours, Engel (C. R. 102, 111) obtained large crystals of the trihydrate, ZnCl₂.3H₂O, melting at 7°. When ZnCl,Aq containing 79.9 p.c. ZnCl₂ at 0° is kept for a long time it solidifies; on heating to 16° a part melts and a part remains solid; the liquid part is the dihydrate ZnCl₂.2H₂O, and the solid part is the hydrate 2ZnCl₂3H₂O (E., l.c.). By heating a syrupy solution of ZnCl, with a little HClAq, Schindler (Mag. Pharm. 36, 45) obtained octahedral crys. tals of the monohydrate ZnCl₂.H₂O. According to Engel (l.c.), the hydrate obtained by S. was 2ZnCl₂.8H₂O.—2. With hydrogen chloride and water. Engel (M. 5, 432) obtained the compounds 2ZnCl₂HCl.2H₂O and ZnCl₂.HCl.2H₂O by passing HCl gas into ZnCl2Aq, and adding zinc from time to time, and, when the solution had the S.G. of 2.0, cooling to 0° .- 3. Various compounds of ZnCl, with ammonia have been desci bed; the formulæ ZnCl, NH, ZnCl, 2NH, 2(ZnCl2.2NH4).H2O, ZnCl,4NH,H2O, ZnCl₂.5NH₂.H₂O have been given to compounds obtained by dissolving ZnCl₂ in NH₂Aq, passing in NH2, and evaporating under different conditions (v. Marignac, Ann. M. [5] 12, 1; Divers, C.N. 18, 18; Priwoznik, P. 142, 467; Davis, C. N. 25, 265; André, A. Ch. [6] 3, 84, 98; Thoms, B. 20, 748).—4. Lang (B. 21, 1578) described a compound with pyridine, ZnCl, 2C, H, N.—5. ZnCl, equivalent of ZnSO, in solution produces no pp., combines with alkali chlorides. Compounds, but the solution gives a pp. of ZnS or hydro ZnCl, NH, Cl.xH,O, ZnCl, 2NH, Cl.xH,O, and ZnCl, 3NH, Cl are described by Schindler (Mag. Pharm. 36, 45), Marignac (J. 1857. 217), Rammelsberg (P. 94, 508), and Hantz (A. 66, 287). A compound ZnCl₂.2KCl is described by R. (l.c.) and M. (l.c.); and a compound ZnCl₂2NaCl.8H₂O by M. (l.c.). Wells a Campbell (Zeit. f. anorg. Chemie, 5, 278) describe two compounds with CsCl, to which they give the formulæ ZnCl₂.2CsCl and ZnCl₂.3CsCl.—6. With sinc oxide, v. Oxychlorides, p. 891.

A solution of zinc in conc. HClAq, to which as much NH,Cl has been added as the weight of zinc dissolved, is used for cleaning the surfaces of metals that are to be soldered together; any oxide on the metallic surfaces reacts with the NH,Cl present to form chloride which dissolves

in the ZnCl₂ solution.

Zinc, cyanide of, and double zinc cyanides; v. vol. ii. p. 347. For details regarding the conditions of formation and the properties of sincmercuric cyanide, v. Dunstan, C. J. 61, 666

Zinc, ferrocyanide of; v. vol. ii. p. 337.

Zinc, fluoride of, ZnF₂. Formula probably molecular, from analogy of ZnCl₂. Formed by heating zinc or ZnO to redness in a stream of dry HF; also by the interaction of fused ZnCl2 and dry HF at 800° to 900°; also by heating the hydrated salt to redness in dry HF (Poulenc, C. R. 116, 581 [1893]). The tetrahydrate ZnF2.4H2O is obtained by evaporating a solution of zinc in HFAq, and crystallising. The salt ZnF, crystallises in colourless needles, probably monoclinic. S.G. 4.84 at 15°; the crystals act strongly on polarised light (P., l.c.). Clarke (Am. S. [3] 13, 291) gives S.G. of ZnF₂ as 4.556 at 17°, and 4.612 at 12°; and the S.G. of ZnF₂.4H₂O as 2.567 at 10° and 2.535 at 12°. ZnF₂ is slightly soluble in cold, more soluble in hot, water; it is insoluble in alcohol at 95°; soluble in boiling HClAq, H,SO,Aq, or HNO,Aq (P., l.c.). Heated in air, or to redness in steam, it is wholly changed to ZnO; fusion with alkali carbonates produces ZnO and alkali fluorides; heating with H2S forms ZnS, and with HCl forms ZnCl₂. It is reduced by H at a red heat (P., l.c.). ZnF₂ forms double salts with AlF₃ and with alkali fluorides. Berzelius (P. 1, 26) described the salts ZnF, 2AlF, and ZnF, 2KF; and Wagner (B. 19, 896) the ZnF, 2NH, F. 2aq, ZnF, KF, and ZnF, NaF. The compound ZnF₂.ZrF₄. 6aq (= ZnZrF₆. 6aq) is described by Marignac (A. Ch. [8] 60, 257); S.G. 2.255 at 12° (Topse, C. C. 4, 76). For ZnF.SiF. v. Zinc silicofluoride, p. 890.

Zinc, hydrophosphide of, ZnPH. A white, friable solid having this composition was obtained by Drechsel a. Finkelstein (B. 4, 352) by passing dry PH, into a cooled ethereal solution of ZnEt. The compound can be keptein a closed vessel; in air it absorbs moisture and

rapidly decomposes, giving off PH.
Zinc, hydrosulphide of. According to Thomsen (B. 11, 2044 [1878]), the pp. obtained by mixing NaHSAq and ZnSO,Aq, in equivalent quantities, is probably a hydrosulphide of zinc (no analyses or formula given). Addition of

equivalent of ZnSO, in solution produces no pp., but the solution gives a pp. of ZnS or hydro sulphide on standing, or on addition of acid or soda (T., l.c.). According to von Zotta (M. 10, 807 [1890]), the pp. supposed by Thomsen to be zinc hydrosulphide has the composition ZnS,H,2ZnS. When four equivalents of NaSH are added to one equivalent of ZnSO, both in solution, and the clear solution is boiled, or treated with acid or alkali, a pp. of ZnS, H2.2ZnS is obtained (von Z., l.c). Linder a. Picton (C. J. 61, 130 [1892] obtained evidence in favour of the existence of compounds of Zn, S, and H, probably 7ZnS.H₂S and 12ZnS.H₂S, by passing H2S into water at 0° with ZnO2H2 in suspension; they failed to isolate a definite compound with

certainty.

Zinc, hydroxide of, ZnO₂H₂. (Zinc hydrate. Hydrated zinc oxide.) Obtained, as a white flocculent pp., by adding to solution of a salt of zinc rather less KOHAq than is sufficient to decompose the whole of the salt, washing thoroughly, and drying at a low temperature. According to Ville (C. R. 101, 875), ZnO_2H_2 is obtained in crystals by shaking very finely-powdered ZnCO, or basic carbonate, with twice as much kOH in 10 p.c. solution as is theoretically required for the decomposition of the carbonate; formation of crystals of ZnO₂H₂ begins at once, and is complete after twenty to thirty minutes. ZnO2H2 is said to be obtained in lustrous rhombic prisms by immersing zinc in contact with iron or copper in NH, Aq (Nicklès, A. Ch. [3] 22, 31; Cornu, Bl. 5, 64 [1863]). Bödeker (A.94, 358) obtained the monohydrated hydroxide ZnO,H,.H₂O in very lustrous octahedra by keeping a saturated solution of the hydroxide in NaOHAq in a closed vessel for some weeks. S.G. of $Zn\hat{0}_{s}H_{2}$ is given by Nickles (*l.c.*) as 2.677, and by Filhol (*A. Ch.* [3] 21, 415) as 3.053. Thomsen gives [3] 21, 415) as 3.053. Thomsen gives $[Zn,O,H^2O] = 82,680$ (Th. 3, 270). ZnO_2H_2 dissolves easily in acids, forming salts ZnX12

ZnO₂H₂ also dissolves in caustic alkali solutions, and in (NH,),CO,Aq. By adding alcohol to a solution of ZnO,H, in KOHAq, Laux (4.9, 183) obtained small lustrous crystals (?ZnO₂K₂); Fremy (C. R. 15, 1106) obtained (?) ZnO₂K₂. ZnO; by using NaOHAq Comey a. Jackson (B. 21, 1589 [1888]) obtained two compounds derived from ZnO,H₂ by replacing H by Na. They shook a solution of ZnO₂H₂ in conc. NaOHAq with alcohol, separated the upper (alcoholic) layer and allowed it to crystallise, and treated the lower (watery) layer repeatedly with alcohol until it solidified. The crystals from the lower

layer had the composition

 $Zn_{s}O_{s}Na_{4}H_{2}.17H_{2}O(-Zn_{s}(OH)_{2}(ONs)_{4}.17H_{2}O)_{5}$ agreeing, except in water of crystallisation, with an ammonium compound described by Malaguti (C. R. 62, 413 [1866]); this compound melts at c. 70°, rapidly absorbs CO₂ from the air, loses 12H₂O at 100°, is insol. in ether, and is uccomposed by water, but not in presence of excess of NaOH, with formation of ZnO and a little ZnO.H. The crystals from the upper layer had the composition

ZnO.NaH.7H₂O = (2Zn(OH)(ONa).7H₂O), corresponding with the compound described by Fremy (l.c.); this compound forms white radisting needles, which do not meet at 300°; it absorbs CO, from the air more slowly than the other compound, is decomposed by alcohol or water, and does not lose H2O until heated above 200° (v. Am. 11, 145 [1889]). C. a. J. failed to obtain compounds derived from ZnO₂H₂ by replacing H by NH, or Mg. Compounds of ZnO with BaO, CeO, and MgO are described by Bertrand (C. R. 115, 939); these compounds are of the forms 2ZnO.MO.xH₂O.

Zinc, iodide of, ZnI₂. Formula probably molecular, from analogy of ZnCl₂. Formed by heating together zine and iodine, and subliming; also by digesting I with water and excess of zinc until the liquid is colourless and evaporating over H₂SO, (Rammel berg, P. 43, 665). Crystallises in white octahedra; S.G. 4696 (Bödeker); melts at c. 446° (Carnelley, C.J. 33, 278); and boils at c. 694° (C. a. Williams, v. Carnelley's Melting-Boiling-point Tables, 1, 23). H.F. (Zn,I'] = 49,230; $[Zn,I^2,Aq] = 60,540$ (Th. 3,275). Heated in air ZnI_2 gives off I, and ZnO remains. Kremers (P.111,61) gives the following data:-

S.G. ZnI, Aq	P.c. ZnI.	S.G. ZnI, Ag	P.o. ZnI.
1.2340	23.1	1.9746	63.5
35121	42.6	2.3976	76.0
1.7871	56.3	,	

Hot ZnI, Aq is said to dissolve zinc in contact with the air, and on filtering an oxyiodide is said to be deposited (Müller, J. pr. 26, 441). ZnI₂ Aq also dissolves I; Baup (J. Ph. 9, 37 [1823]) says enough I is dissolved to form ZnI₄.

ZnI, combines with NH, By allowing a solution of ZnI, in NH, Aq to evaporate, Rammelsberg (P. 48, 152) obtained lustrous, four-sided prisms of ZnI, 4NH, decomposed by water with production of ZnO. Various double compounds with other iodides have been described: ZnI,2MI, where M=NH₄, K or Na (Rammelsberg, l.c.; also P. 43, 665); ZnI₂.2CsI and ZnI₂.8CsI (Wells a. Campbell, Zett. f. anorg. Chemie, 5, 273); 2ZnI, BaI, ZnI. 2Bil. 12aq (Linau, P. 111, 240).

Zinc, nitride of, Zn_2N_2 . Obtained by Frankland (Pr.~8,~502) by heating zinc amide ($Zn(NH_2)_2$; v.~p.~887) to low redness. A grey, pulverulent solid; unchanged by heating to redness out of contact with air; reacts violently with water to give ZnO2H2 and NH2.

Zinc, nitroprusside of, v. vol. ii. p. 841.

Zinc, oxides of. Besides the normal basic oxide ZnO, there seems to be another higher (? superoxide) of zinc.

ZINC OMDE ZnO. Formula not certainly molecular.

Occurrence.-With oxides of Mn, and Fe, as red sinc ore and franklinite.

Formation.—1. By burning zinc in air or O. By strongly heating ZnO₂H₂, Zn(NO₂)₂, ZnCO, or another salt of zinc with a volatilisable

Preparation.—Pure ZnSO, prepared as described under Zinc, Preparation (p. 883), is dissolved in water, and the solution is very slowly added to c boiling solution of pure Na₂CO₃ containing rather less Na.CO, than suffices to decompose all the ZnSO. The pp. of basic zinc carbonate is washed by decantation with boiling water until the washings are free from carbonates

and sulphates; it is then dried, powdered, and heated either in a Hessian crucible, or, better, in a flask of hard glass, until a portion taken from the middle dissolves, when cold, in acid without effervescence. The solid is now finely powdered, sifted through a fine sieve, again heated, and then rubbed up with water, and washed and dried. ZnO is prepared in crystals by heating ZnCl, to redness in steam (Daubrée, C. R. 89, 153); also by heating the amorphous oxide slowly in a stream of H (Regnault, A. Ch. [2] 62, 850; Deville, A. Ch. [8] 43, 477); also by heating the amorphous oxide to whiteness in a stream of O (Sidot, C. R. 69, 202); also by heating Zn(NO₂), to redness (Brügelmann, W. 4, 283).

Properties.—A slightly yellow, powdery solid; becomes darker yellow when heated, but recovers its original colour on cooling. S.G. 5.5 to 5.7 (v. Brügelmann, B. 13, 1741; Filhol, A. Ch. [3] (v. brugeliman, B. 15, 1741; Fillol, A. Oh. 15; 1415; Herapath, C. J. 1, 42; Karsten, S. 65, 394). S.G. crystalline (at 15°) 5·782 (Brügelmann, W. 4, 286). ZnO crystallises in slightly yellow, lustrous, hemimorphous pyramids belonging to the hexagonal system (B., L.c.). Thomsen (Th. 8, 275) gives H.F. [Zn,O] = 85,480. Richards a. Rogers (Am. 15, 567 [1893]) found that ZnO prepared by heating Zn(NO) gave off that ZnO, prepared by heating Zn(NO₂)₂, gave off some gas when dissolved in dilute acids; the maximum amount of gas from 10 g. ZnO was 20 c.c.; the gas consisted chiefly of N, with a little O; the higher the temperature to which the Zn(NO_s), had been heated the less was the quantity of gas obtained; but some N always came off, even from ZnO which had been heated to the softening temperature of porcelain. Moissan (C. R. 115, 1034 [1893]) states that ZnO volatilises readily when heated in an electric furnace, and condenses again in transparent crystals. ZnO is not decomposed at c. 1750° (Read, C. J. 65, 313 [1894]).

Reactions and Combinations.—1. ZnO is a basic oxide. It interacts with most acids to form salts ZnX(X=2NO, SO, \$PO, &c.) and water.-2. With fairly conc. caustic soda or potash, salts are formed which are most simply regarded as derived from ZnO2H2, with H replaced by Na or K (v. Zinc hydroxide, p. 888).-3. ZnO is reduced to Zn, with explosive violence, by heating with magnesium (v. Winkler, B. 28, 128).-4. Morse a. White (Am. 11, 258) heated a mixture of ZnO and sinc in an exhausted tube, and found that a deposit of ZnO formed in the front part of the tube. They suppose that some of the original ZnO is decomposed by the combined action of heat and zinc, and that zinc and O exist together in the hot part of the tube, but combine in the cooler parts of the tube to form ZnO.-5. Oxychlorides, oxybromides, and probably oxylodides of zinc are formed by heating ZnO with sinc chloride, bromide, and iodide respectively.-6. ZnO is reduced to zinc by heating with carbon or in carbon monoxide.—7. ZnO does not directly interact with water; ZnO.H. is produced by ppg. a sine salt solution by an alkali (v. Zinc hydroxide, p. 888).—8. By heating ZnO with oxides of aluminium, chromium, and iron along with B₂O₂ in a pottery furnace until the B₂O₃ was volatilised, Ebelmen (A. Oh. [3] 33, 34) obtained the compounds ZnC.Al.O. ZnO.Cr.Os, and ZnO.Fe.Os.

ZINC PEROXIDE. In 1818 Thénard (A. Ch. [2] 9, 55) obtained what he called 'deutoxide de zine by the action of H2O2Aq on gelatinous ZnO2H2, and also by adding KOHAq to a solution of ZnO2H2 in HClAq to which H2O2 had been added. In 1884 Haas (B. 17, 2249) obtained a white, tasteless, neutral, gelatinous pp. by adding H2O2Aq to solutions of zinc salts and then adding NH, Aq; after washing and drying at 100°, the solids had approximately the compositions Zn,O, and Zn,O,. H. thinks it probable that the substances obtained by him were compounds of ZnO and ZnO₂; the formula 2ZnO.3ZnO₂ agrees well with some of his analyses.

Zinc, exybromides of. Oxybromides of zinc are formed by heating solutions of ZnBr, with ZnO, and by partially ppg. ZnBr,Aq by NH,Aq. Several compounds have been described by André (Bl. [2] 39, 398 [1884]):

(1) $ZnBr_2.4ZnO.xH_2O$, x=10, 13, and 19;

(2) ZnBr₂.5ZnO.6H₂O; (3) ZnBr₂.6ZnO.35H₂O. Zinc, oxychlorides of. These compounds are

formed by adding to ZnCl2Aq a quantity of NHAq less than sufficient for complete ppn.; also by boiling ZnCl2Aq with ZnO, or with certain other metallic oxides. Schindler (Mag. Pharm. 36, 45) obtained ZnCl, 3ZnO.4H,O by boiling ZnCl, Aq with ZnO, and the same compound by partially ppg. ZnCl2Aq by NH3Aq. By adding to ZnCl,Aq so much NH,Aq that the pp. which formed at first partly dissolved, and also by treating ZnCl₂.4NH, with water, Kane (A. Ch. 72, 296) obtained ZnCl₂.6ZnO.10H₂O. André (C. R. 106, 854) obtained 2ZnCl₂.3ZnO.11H₂O by boiling ZnCl2Aq with HgO and letting the solution cool; using PbO he obtained another oxychloride, probably ZnCl_3ZnO, combined with ZnCl,.PbO.

Zinc, expiodides of. According to Müller (J. pr. 26, 441 [1842]), ZnI₂Aq dissolves zinc when warmed therewith in the air; on cooling,

an oxyiodide, ZnI,3ZnO.2aq, is ppd.
Zinc, oxysulphides of. The compound 4ZnS.ZnO occurs native as voltzite, as spherical globules in some specimens of galena. A compound having the same composition was found in a furnace flue at Freiberg by Karsten (S. 57, 186). By heating dry ZnSO, to redness in H, Arrivedson obtained a yellow-coloured oxysulphide ZnS.ZnO (P. 1, 59).

Zinc, perbromoplatinocyanide of, v. vol. ii. p. 845.

Zinc, phosphides of. Zinc and P combine when heated together, forming a greyish, lustrous, somewhat malleable compound. By heating finely powdered zinc in vapour of P, Schrötter obtained a compound, with S.G. 4.76, having the composition Zn,P2 (W. A. B. 1859. 801). Vigier obtained the same compound by heating zinc in H until distillation began, and then passing H laden with vapour of P over the minc. The same compound was formed by Regnault (A. Ch. [4] 9, 162) by heating to we iteness a mixture of MgHPO4, ZnS and C; along with the Zn, P2 were formed crystals of another phosphide, probably ZnP; and by treating Zn,P, with dilute acid, an amorphous, yellow solid remained which detonated when mixed Hvoslef (A. 100, 99) by strongly heating a mixture of ZnO, P.O., and C. By heating a mixture of NaPO, and zinc shavings in a retort, H. (l.c.) obtained a yellowish red, loose sublimate which contained Zn and P; and by heating the fused mass in the retort with HClAq, grey lustrous tablets of ZnP2 remained. Lüpke (C. C. 1890 [ii.] 642) formed zinc phosphide (? Zn,P2) by adding amorphous P to molten zinc covered with (NH₄)₂CO₃; this phosphide was not decomposed by water, but was slowly decomposed by dilute HClAq, giving off PH,

For the compound ZnPH v. ZINC HYDROPHOS-

PHIDE, p. 888.

Zinc, platinocyanide of, v. vcl. ii. p. 344.

Zinc, salts of. Many salts are obtained by replacing the H of acids by zinc. The greater number of the salts of zinc are normal selts; several basic carbonates, nitrater, and sulphates are known. The following are the chief salts of zinc derived from oxyncids:-Arsenite and arsenate, borate, bromate, carbonates, chlorate and perchlorate, chromates, iodate and periodates, molybdates, nitrates and nitrites, phosphates and phosphites, selenate and selenite, silicate, sulphates and sulphite, thiosulphate, tungstates, vanadates (v. Carbonates, Nithates, SULPHATES, &c.).

Zinc, selenide of, ZnSe. This compound is formed, in reddish-yellow, regular crystals, S.G. 5.40, by heating zinc to redness in a stream of H2Se, and then subliming in a slow stream of

H (Margottet, C. R. 84, 1293).

Zinc, selenocyanide of, v. vol. ii. p. 348. Zinc, silicofluoride of, ZnSiF₆.6H₂O. Obtained in hexagonal plates, easily sol. in water, by evaporating to a small bulk a solution of ZnO in H₂SiF₅Aq (Berzelius). S.G. 2·104 (Topsöe, C. C. 4, 76).

Zinc, stannifluoride of, ZnSnF.6H2O; v.

this vol., p. 724.

Zinc, sulphocyanide of, v. vol. ii. p. 353.

Zinc, sulphide of, ZnS. This compound is not formed by heating together zinc and S, as the S volatilises before the temperature of combination is reached.

Occurrence.—As zinc blende.

Preparation .- 1. By repeatedly subjecting a mixture of zinc and S, in the ratio Zn:S, to a great pressure; the product resembled zinc blende (Spring, B. 16, 1,000).—2. By rapidly heating a mixture of zinc filings and cinnabar in a retort; formation of ZnS occurs with detonation; Hg distils off.—3. By heating ZnO with S.-4. By heating ZnO, or ZnO2H2, in a stream of H,S; in the case of ZnO,H, formation of ZnS takes place very slowly (v. Wagner, D. P. J. 197, 334).—5. ZnS is also formed by heating dry ZnSO, with S and C.—6 ZnS is obtained by passing H2S into a neutral or alkaline solution of a zinc salt, or into a solution in a weak acid, such as acetic acid, washing the pp., drying it at 100°, and heating in a stream of u.y 11,00. The pp. dried at 100° is said to be 2ZnS.H₂O (Geiger a. Reimann, Mag. Pharm. 81, 178; cf. Souchay, Fr. 7, 78).

Properties and Reactions.—S.G. of ZnS = 8.93 Zn,P, with dilute acid, an amorphous, yellow (Kalsten, S. 65, 394); S.G. of blende = 4.08 to solid remained which detonated when mixed 4.07 (v. Neumann, P. 23, 1). Thornsen (Th. 3, with KOlO, and struck, and which was perhaps 276) gives H.F. of ZnS.xH₂O as [Zn,S,xH²O] ZnP_e. The compound Zn₂P₂ was obtained by -41,580. ZnS prepared by ppn. is a yellowish solid; easily sol. in acid, forming salts of Zn and giving off H₂S. Zinc blende is a grey to black crystalline solid; slowly acted on by acids. By repeatedly washing ppd. ZnS, also by passing H₂S into water containing pure ZnO₂H₂ in suspension, Winssinger (Bl. [2] 49, 452 [1888]) obtained colloidal "nS, sol. in water; the solution could be boiled until all excess of H₂S was expelled without ppn. of ZnS. ZnS prepared by adding excess of NH₂Aq to ZnCl₂Aq, ppg. all the zinc as ZnS by H₂S, washing and drying the pp., and then heating nearly to whiteness in a Hessian crucible inside a plumbago crucible, is very phosphorescent (v. Henry, C. R. 115, 503).

By heating ZnS with zinc in a vacuous tube, Morse a. White (Am. 11, 348) found that some of the ZnS was carried forward and deposited on the cooler part of the tube; they supposed that some of the ZnS was decomposed, and that Zn and S existed in the hottest part of the tube and combined in the cooler portions. By heating blends with I and a little water some ZnSO, is formed (Filhol a. Melhies, A. Ch. [4] 22, 58;

Lasaulx, J. 1870. 1272).

According to Schiff (A.115,74), a 2 alysulphide of sinc, probably ZnS₃, is formed by ppg. a neutral solution of a salt of zinc by K₂S₄Aq, and drying the pp. over H₂SO₄. Spring (B.16, 1000) thought that a polysulphide of zinc was probably formed by strongly compressing a mixture of zinc with excess of S.

Zinc, telluride of, ZnTe. Obtained, in large ruby-red, regular crystals, by strongly heating zinc in a stream of H₂Te, and then subliming in a slow current of H (Margottet, C. R. 84, 1293).

Zinc, titanifluoride of, ZnTiF.; v. this vol., p. 742. M. M. P. M. ZINC, ORGANIC COMPOUNDS OF.

Zinc methyl or Zinc methide ZnC₂H₆ i.e. ZnMe_x Mol. w. 95. V.D. 3·29 (calc. 3·30). [-40°] (Haase, B. 26, 1053). (46°). S.G. 105 1·386.

Formation.—1. By heating methyl iodide with zino in sealed tubes at 150°. The compound IZnMe is formed as an intermediate product. The reaction takes place more easily when the MeI is mixed with two-thirds of its volume of ether and heated to 100°, but it is then difficult to separate the ether from the ZnMe₂ (Frankland, A. 85, 346; 111, 62; Wanklyn, C. J. 13, 124; Butlerow, A. 144, 2; Ladenburg, A. 173, 147).—2. By heating finely granulated zino with HgMe₂ in a sealed tube for 24 hours at 120°, and distilling the product (Frankland a. Duppa, C. J. 17, 30).—3. IZnMe is formed in 24 hours if MeI is left in contact with a large amount of copper-zinc couple (Gladstone, C. J. 35, 109).

Preparation.—Zinc filings (9 pts.) are heated with reduced copper (1 pt.) till the copper zinc couple is produced, and the product treated with

Mel (Gladstone, C. J. 35, 569).

From Periods.—Colourless mobile liquid, taking fire on exposure to air Decomposed by water into CH, and Zn(OH). By very careful oxidation it appears to yield Zn(OMe), and crystalline ZnMe(OMc), which is split u, by water into Zn(OH), methyl alcohol, and CH.

Reactions.—1. Acetyl chloride (1 mol.) forms CH, CCI(OZnMe). CH₂, which is slowly converted, by further treatment with ZnMe₂, into

CH₁-CMe(OZnMe).CH₂+ClZnMe, crystallising in prisms, and converted by water into tertbutyl alcohol, CH₄, zinc chloride, and Zn(OH)₂. The compound CH₂.CCl(OZnMe).CH₄ yields acetone on addition of water (Butlerow, Z. 1864, 365, 702; Pawlow, N. Petersb. Acad. Bull. 22, 497; Bogomoletz, A. 209, 88).—2. Bromo-acetyl bromide, followed by water, yields methyl-isopropyl-carbinol (Winogradoff, A. 191, 127).—8. In ethereal solution secondary hexyl iodide has no action at 100°; but at 125° the products appear to be CH₄, ethyl iodide, and hexylene (Purdie, C. J. 39, 464).—4. Chloral followed by water forms CCl₂CHMe.OH, while butyric chloral yields C₃H₄Cl₃.CHMe.OH.—5. Nitricoxide passed into ZnMe₂ forms CH₃N₂O₂(ZnMe), which is decomposed by water, yielding crystalline (CH₂N₂O₂)₂Zn aq, from which the Na salt CH₃N₂O₂Na aq of 'di-nitro-methylic' acid may be obtained (Frankland, A. 99, 369).

Zinc ethyl or Zinc ethide ZnEt, Mol. w. 123. [-28°] (Haase, B. 26, 1053). (118°). S.G. 12 1:182. H.F. -31,800 (Guntz, C. R. 105,

673). V.D. 4.26 (obs. and calc.).

Formation.—1. By heating zinc or zincsodium alloy with EtI or a mixture of EtI and
ether (Frankland, C. J. 2, 297; 3, 44; Tr. 142,
431; 145, 259; A. 95, 28; Wanklyn, C. J. 13,
124; Pebal, A. 118, 22; 121, 105; Rieth a.
Beilstein, A. 123, 245; 126, 248; Chapman,
Laboratory, 1, 195; Alexejeff a. Beilstein, C. R.
63, 171; Rathke, A. 152, 220; Wichelhaus, J.
1868, 425).—2. By heating granulated zinc with
HgEt, in sealed tubes for 36 hours at 100°
(Frankland a. Duppa, C. J. 17, 3).—3. From
zinc filings and EtBr (Wichelhaus, A. 152, 321).

Preparation.—Clean zinc filings (180 g.) are mixed with copper (20 g.) reduced from the oxide at a dull-red heat by coal gas; the mixture is rapidly heated over a very large Bunsen flame, and is well shaken meanwhile. As soon as the filings begin to cake together and the copper is no longer visible, the copper-zinc couple has been made. The flask is now connected with an inverted condenser, inclined slightly upwards, and when it is cold ethyl iodide (175 g.) is added. Dry CO₂ is passed through the apparatus, and the flask heated by a water-bath; in half an hour the EtI ceases to run back from the condenser, for it has been converted into IZnEt. The apparatus is now tilted so that the condenser is inclined downwards, and heated in a bath of paraffin. The reaction 2IZnEt = ZnI₂ + ZnEt₂ now takes place, and ZnEt₂ (60 g.) is collected in a receiver kept full of CO₂ (Gladstone a. Tribe, C. J. 26, 445;

Properties.—Colourless, mobile liquid, fuming in the air, and quickly taking fire, burning with a luminous green-edged flame, and giving off clouds of ZnO. Immediately decomposed by water into Zn(OH)₂ and ethane. In like manner it attacks organic compounds containing hydroxyl or amidogen, with elimination of ethane. It causes india-rubber to swell remarkably.

Reactions.—1. Gradually treated with dry oxygen it appears to produce ZnEt(OEt) and Zn(OEt), successively (Frankland). According to Demuth a. Meyer (B. 23, 394), the passage of oxygen through a solution of ZnEt in ligroin

(7 vols.) yields EtZn.O.O.Et, which is decomposed by heat, sometimes with explosive violence. The compound EtZn.O.OEt sets free I from a solution of KI and yields alcohol when distilled with dilute H.SO₄.—2. Sulphur forms zinc mercaptide Zn(SEt)₂. Se appears to act in like manner (Chabrié, Bl. [3] 2, 788).—3. Iodine forms. IZnEt and ZnI₂ successively, together with Etl. Bromine acts in like manner. Zinc ethyl takes fire in chlorine.—4. Dry ammonia forms $Zn(NH_2)_2$ and ethane (Frankland, Pr. 8, 502).-5. Nitric oxide is absorbed by an ethereal solution of ZnEt, forming crystalline ZnEt, (NO), which melts below 100° and is split up by water Into ethane and CoH. N.O. ZnOH. Carbon dioxide passed into an aqueous solution of this salt ppts. zinc carbonate, while zinc 'dinitroethylate' (C₂H₃N₂O₂)₂Zn remains in solution (Frankland, Tr. 1857, 59; A. 99, 342; C. J. 37, 570). The salt (C₂H₃N₂O₂)₂Zn aq crystallises in prisms, yields NH₂ and NH₂Et on treatment with sodiumamalgam (Zuckschwerdt, A. 174, 302), and HNO and NH Et on treatment with alcoholic potash (Zorn, B. 15, 1008). The dinitroethylates NaA', BaA', CaA', Baq, MgA', CuA', agq, AgA', and Ag, A'(NO,) were prepared by Frank-Hand.—6. SO, forms zinc ethane sulphinate (Hobson, C. J. 10, 455).—7. PCl, yields PEt, (Hofmann a. Cahours, Tr. 1857, 578).—8. SiCl, gives SiEt, -9. Heated with EtI at 170° it forms n-butane and ZnI2. Ethane and ethylene are also formed (Brodie, C. J. 3, 405). ZnEt, does not act on amyliodide.-10. Displaces halogens in organic compounds by ethyl.—11. Adds itself to a carbonyl group. Thus with di-ethyl-ketone it forms Et₂C(OZnEt).Et, which is converted by water into tri-ethyl carbinol. Di-propyl ketone forms, in like manner, ethyl-di-propylcarbinol. Ketones containing the group CO.CH, act differently, forming condensation-products; thus acetone yields mesityl oxide and phorone (A. Saytzeff, J. pr. [2] 31, 319). Zinc ethyl has no action on paraldehyde (Wedensky, J. pr. [2] 39, 588). With acetic aldehyde it forms CH. CHÉt.OZnEt, which is converted by water into sec-butyl alcohol.—12. In some cases it can add ZnEt and H to a carbonyl group, C2H4 being evolved. Thus ZnEt, converts chloral, dissolved in pure ether, into crystalline CCl. CH. OZnEt, which is converted by water into tri-chloro-ethyl alcohol (Delacre, Bl. [2] 48, 784). Further action of zinc ethide on chloral yields (CCl_s.CH₂O)₂Zn, a white powder. In like manner butyric chloral is converted into C.H.Cl.CH.OH. In this splitting off of an olefine, ZnEt, differs from ZnMe2, but resembles zinc propyl and zinc isobutyl (Garzarolli, A. 223, 162).—13. Can displace oxygen by two ethyl radicles; thus it converts oxalic ether into CO2Et.CEt2.CO2Et (Frankland a. Duppa, A. 142, 1; Pr. 12, 396.—14. Can displace ethoxyl by ethyl. Thus it converts formic ether into HCEt(OZnEt).OEt and then into HCEt(OZnEt).Et, which is turned by water into di-ethyl-carbinol (Wagner a. Saytzeff, A. 175, 851).-15. Can displace chlorine by hydrogen. Thus with chloro-benzoic acid the first product is probably C.H.(ZnEt).CO.Et, which is converted by water into benzoic ether, C₂H_a, and ZnO. 16. With benzonitrile at 150°, and subsequent treatment with alcohol, cyaphenine is formed (Frankland a. Evans, C. J. 87, 568). It

polymerises phenyl-acetonitrile in the same way, forming cyanbenzine.—17. Cyanogen forms ZnCy, and EtCN (Frankland a. Graham, C. J. 87, 740).-18. Azobenzene is reduced by ZnEt, followed by water, to aniline (Frankland a. Louis, C. J. 37, 563).—19. Acts on primary and secondary amines, giving off ethane, but has no action on tertiary amines (Frankland, Pr. 8, 502; Gal, J. Ph. [5] 7, 484). ZnEt, has no action on nicotine or quinoline, but forms with many alkaloids containing O compounds from which the alkaloids are regenerated by water.—20. Acetamide gives (AcNH), Zn, while oxamide gives C₂O₂(NH), Zn.—21. Acts on HgCl₂, forming Cliget and HgEt, (Buckeen, A. 109, 218). Hg₂Cl₂ yields HgEt₂, mercury, and ZnCl₂—22. AgCl forms ZnCl₂, silver, and butane.— 23. CCl, forms EtCl, ethylene, and propylene (Rieth a. Beilstein, A. 124, 242) —24. Chloroform yields amylene (R. a. B.), while bromoform gives propylene and EtBr (Alexejeff a. Beilstein, C. R. 58, 172). CHCl₂I produces ethylene, butane,

and hydrogen (Borodine, A. 126, 239).

Zinc brome-ethide ZnBrEt. [62°]. Formed by the actic n of the copper-zinc couple on EtBr at 100° (Gladstone a. Tribe, C. J. 27, 410). Pearly scales, decomposed by heat paray into ZnBr, and ZnEt, and partly into Zn, ZnBr,

C₂H₄, and C₂H₄. Zinc iodo-ethide ZnIEt. Formed in like manner. Crystalline. Split up by heat into

ZnI, and ZnĚt,

Zinc propyl ZnPr. (148°) (S.); (146°) (Gladstone a. Tribe, B. 6, 1186); (150°) (Pape, B. 14, 1873). Formed by heating propyl iodide with zinc and a small quantity of an alloy of zinc and sodium (Stcherbakoff, Bl. [2] 37, 345). Heavy liquid, with disagreeable odour. Reacts with aldehydes, forming propylene and such compounds as RCH, OZnPr, which yield secondary alcohols on addition of water.

Zinc isopropyl ZnPr2 (116°) with decomposition; (96° at 40 mm.). Got by heating PrI (170 g.) with ether (100 g.), zinc (100 g.), and zinc-sodium (3 g.) at 100° (Ragozin, J. R. 24, 549). Fuming liquid. Yields crystalline Zn(OPr), on exposure to air. Split up at 132° into Zn and hydrocarbons (Gladstone a. Tribe, C. J. 39, 6).

Zinc isobutyl $Zn(C_4H_9)_2$. (166°) at 734 mm. (Garzarolli-Thurnlackh a. Popper, A. 223, 167). Formed from zinc and Pr.CH.I. Got also from Zn and $Hg(C_4H_9)_2$ (Cahours, Bl. [2] 21, 857). Zinc isoamyl $Zn(C_9H_{11})_2$. (220°). S.G. 2 1.022. V.D. 6.95 (calc. 6.87). Prepared by heat-

ing zinc with Hg(C₅H₁₁)₂ (Frankland, A. 85, 360; 130, 122). Liquid, furning in the air but not taking fire. Takes fire in oxygen. When added to cooled aldehyde it yields, after addition of ice-cold water, methyl-isoamyl-carbinol, isopropyl-ethylene, and ethyl and isoamyl alcohols (Sokoloff, C. C. 1887, 988).

ZINC HYDROXIDES, p. 888.

ZIRCONATES v. under ZIRCONIUM HYDROXIDE, p. 896.

ZIRCONIA; raide of sirconium (q v., p. 896). ZIRCONIUM. Zr. At. w. c. 90 (v. p. 894). Mol. w. not known. Melts above m.p. of Si (i.e. above c. 1800°) (Troost, J. 1865. 188). S.G. 4·15 (Troost, J.c.); 4·25 (Moissan, G. R. 116, 1222). S.H. (0°-100°) ·066 (Mixter a. Dana, A. 169, 388 [1873]).

Historical.—Klaproth in 1789 (Beiträge, 1, 203, 227) announced the discovery of a new earth, in combination with silica, in the rare mineral sircon. The same earth has been found in a few other rare minerals, such as eudialyte, polymignite, fergusonite, &c. Bright-coloured varieties of zircon are generally called hyacointh, the greyish or brown kinds are called sirconite, and a variety which is nearly colourless is known as jargon.

In 1845 Svanberg (A. 56, 223) stated that zirconia was a mixture of at least three oxides, but Berlin (J. pr. 58, 147), Marignac (A. Ch. [3] 60, 257), and Hermann (J. pr. 31, 75; 97, 331) failed to obtain any evidence of the composite character of zirconia. At a later time, Sorby (B. 2, 126, 193 [1869]), and also Forbes (B. 2, 337, 383), thought they had obtained spectroscopic indications of the presence of another earth in zircon; but it was shown that the abnormalities observed were due to traces of compounds of tranium (Sorby, C. N. 21, 73; Knop, A. 159, 44; cf. Hannay, C. J. 26, 703).

Occurrence.—Never uncombined. The principal naturally occurring compound of Zr is zirconits, which is more or less pure ZrSiO₄. This mineral is found in small quantities in the sand and alluvial deposits of certain rivers in Ceylon, and in the older rocks in many parts of the world; it has been found in the granites of Sweden, Switzerland, Tyrol, and North America

(v. J. M. 1877. 97).

Formation.—1. By heating K₂ZrF₆ with K (Berzelius, P. 4, 117 [1825]).—2. By passing vapour of ZrF₆ over Na in a red-hot tube (Troost, C. R. 61, 109 [1865]).—3. By heating ZrO₂ with Mg (Phipson, J. pr. 96, 353 [1865]).—4. By melting ZrO₂ in a carbon crucible in an electric furnace; or by mixing ZrO₂ with carbon, melting in an electric furnace, then mixing the Zr carbides thus formed (v. Carbides, p. 894) with excess of ZrO₂ and again melting (Moissan, C. R. 116, 1222 [1893]).

Preparation .- 1. Finely powdered sirconite (ZrO_xSiO₂) is thoroughly mixed with an excess of finely powdered carbon, the mixture is compressed into small cylinders which are placed in a closed carbon crucible and heated in an electric furnace (with a current of 30 to 85 ampères and 70 volts), while a stream of CO2 is sent through the crucible; the greater part of the SiO2 is thus volatilised (Troost, C. R. 116, 1428). The carbides of Zr in the crucible are now mixed with excess of ZrO₂ and melted in the electric furnace (Moissan, C. R. 116, 1222).—2. Finely powdered and sifted sirconite is mixed with 2 to 3 times its weight of KHF2, the mixture is heated to bright redness for c. 15 mins., and, after cooling, is boiled with water containing a little dilute HFAq; K₂SiF₆ is removed by filtration, the filtrate deposits K₂ZrF₆ on cooling. The K₂ZrF₆ is re-crystallised from water, then dried, mixed with K, and heated in a vessel of glass or iron; the product is washed with water, digested for a long time with conc. HClAq at 40°, then washed with water containing NH,Cl, and finally with alcohol (Berzelius, l.c.; Troost, A. 136, 553). This method yields amorphous sirconium. According to Bailey (Pr. 46, 74 [1889]), the only

method that gives nearly pure amorphous Zr is to heat ZrO₂ with Mg foil (not powder) and treat with HClAq.—3. One part dry K₂ZrF₂ is mixed with 1½ pts. Al, and the mixture is heated to the melting-point of iron in a crucible made of gas coke; the product is digested with conc. HClAq diluted with twice its volume of water, when the crystalline leaflets of Zr, that have formed on the surface of the Al, separate from the mass of Al before the latter is wholly dissolved. When the whole of the Al has dissolved an alloy of Zr and Al remains, besides the crystalline leaflets of Zr; if the temperature of reduction is not sufficiently high, the main product is the Zr-Al alloy (Troost, Lc.; cf. Franz, B. 8, 58). This method yields crystalline zirconium.

Various methods have been used for decomposing sirconite and obtaining compounds of Zr. Franz (B. 3, 58) fuses the powdered mineral with KHSO₄, treats the fused material with boiling water containing a little H₂SO₄, washes the insoluble 3ZrO₂·SO₂ thus formed, and adds it, in small portions at a time, to molten NaOH; he then lixivistes with cold water, dissolves the ZrO₂ that remains in hot conc. H₂SO₄, dilutes, ppts. by NH₂Aq, washes, dries, and decomposes the ZrO₂·xH₂O by heat. The method of fusing with KHF₂ used by Marignao (v. supra, No. 2) yields pure K₂ZrF₆ very readily; Zr(SO₄)₂ may be obtained by heating the K₂ZrF₆ with conc. H₂SO₄ (v. Hiortdahl, A. 137, 34; cf. J. pr. 83, 201). For other methods v. Hermann (J. pr. 97, 330); Scheerer (P. 59, 48); Stromeyer (A. 113, 727); Chancel (J. pr. 74, 471); Berlin (J. pr. 58, 145); Henneberg (J. pr. 88, 508); Bailey (Pr. 46, 74 [1889]).

Properties.—Zirconium has been isolated as black amorphous powder, and also as a greyish crystalline solid.

Amorphous sirconium (v. Preparation, No. 2) is a black powder, which is extremely porous and resembles powdered charcoal; when pressed with a burnisher it agglomerates to graphite-like laminæ. After being heated strongly, and then cooled, in vacuo, it is said to glow and burn when exposed to air. The porous amorphous metal is said not to conduct electricity. Amorphous Zr is stated to be insoluble in ordinary soids, including aqua regia; but, according to Bailey (Pr. 46, 74 [1889]), it dissolves in cold conc. H₂SO₄; HFAq dissolves it readily, H being given off. It is oxidised by heating in the air, when it burns to ZrO₂ with a brilliant white light; oxidation is also effected by fusion with caustic or carbonated alkalis, or with KNO₂ or KClO₂.

Crystalline sirconium (v. Preparation, No. 3) is a very lustrous, greyish, brittle, solid, which resembles Sb in appearance; very hard, easily scratching glass and ruby (Moissan, C. R. 116, 1222). S.G. 4·15 (Troost, J. 1865. 183); 4·20 (Moissan, l.c.). S.H. (0° to 100°) ·066 (Mixter a. Dana, A. 169, 888 [1878]). Crystalline Zr burns in the O-H flame; it is not oxidised in Ou. atil heated to whiteness; it dissolves slowly in hot conc. HClAq, and is very slightly acted on by HNO. Aq or H. SO., even when warm; cold HFAq dissolves it readily. Molten KClO., or KNO., is said to be without action; molten KOH dissolves the metal as long as any moisture is present.

According to Troost (J. 1865. 183), Zr melts above the m.p. of Si (i.e. above c. 1300°). The emission-spectrum of Zr has been mapped by Thalén; the chief lines are 6127 in the red, and in the blue, 4815, 4771, 4738, 4709, and 4686.

The at. w. of Zr was determined, (1) by Berzelius (P. 4, 126; 8, 186 [1825]), by finding the ratio of ZrO₂ to SO₃ in the sulphate; (2) by Hermann (J. pr. 31, 77 [1844]) by analysing ZrCl₄: (8) by Marignac (A. Ch. [3] 60, 270 [1860]), by analysing K₂ZrF₆; (4) by Bailey (Pr. 46, 74 [1889]), by decomposing Zr(SO₄), by heat and weighing the ZrO₂ produced; (5) by Mixter a. Dana (A. 169, 388 [1873]), by determining the S.H. of Zr; (6) by Deville a. Troost (P. 108, 636, 641 [1859]), by determining the V.D. of ZrCl. The results of Berzelius gave the at. w. of Zr as 89.2; those of Hermann, 88.5 to 90.7; those of Marignac, 90.3 to 90.6; and the results obtained by Bailey gave 90.4 as the at. w. of the metal.

Zirconium is the third member of the evenseries family of Group IV. in the periodic classification of the elements. This group contains the following elements:-

As is to be expected from its position in the classificatory system, Zr is a metal with some non-metallic properties; the oxide ZrO2 forms corresponding salts, many of which are basic salts; the haloid compounds, ZrX4, are probably all volatilisable without decomposition (the V.D. of ZrCl, only has been determined), but aqueous solutions yield oxyhalides on evaporation; hydrated ZrO, reacts as a weak acid towards strong bases, forming zirconates; salts are known derived from the acid H₂ZrF₆; there is an oxide higher than ZrO₂, probably ZrO₂. For a more detailed comparison of Zr with the other evenseries elements of Group IV. v. TITANIUM GROUP OF ELEMENTS (this vol. p. 749), and cf. Ting Group of ELEMENTS (this vol. p. 735). The atom of Zr is tetravalent in the gaseous molecule ZrCl.

Reactions and Combinations.-1. Heated in air, or oxygen, Zr burns to ZrO2, giving out dazzlingly white light; amorphous Zr burns easily, crystalline only at a white heat, and then merely superficially .- 2. Both amorphous and crystalline Zr dissolve readily in hydrofluoric acid, giving off H and forming ZrF4.-3. According to Bailey (Pr. 46, 74), amorphous Zr dissolves in cold conc. sulphuric acid. Both forms dissolve in aqua regia.—4. Conc. hydrochloric acid reacts with Zr, when warm, forming ZrCl, solution. Crystalline Zr is said to interact with gaseous hydrogen chloride, at dull redness. forming ZrCl₄.—5. Amorphous Zr is oxidised by molten potash or soda, also by molten nitre or potassium chlorate; the crystalline form of Zr is said not to be acted on by molten KNO, or KClO3, and by molten caustic alkalis only as long as there are traces of moisture present .-6. Zr combines directly with the halogens, to form ZrX, and probably with sulphur (v. Sulphide, p. 897).

Detection and Estimation.—Addition of hot

conc. K2SO, Aq to solutions of salts of Zr ppts. white basic sulphate of Zr, which is insoluble in water and nearly insoluble in much HClAq. Turmeric paper becomes reddish-brown when immersed in a solution of a salt of Zr in HClAq. the reaction is completely masked by the presence of boric acid (v. Brush. J. pr. 62, 7). Bailey (C. J. 49, 481 [1886]) found that addition of H₂O₂Aq to a slightly acid solution containing salts of Zr, along with salts of Fe, Nb, and Ti, ppd. the whole of the Zr as ZrO_s (or Zr₂O_s), but none of the other metals. Troost makes use of the comparatively small volatility of ZrO2, at a very high temperature, to separate this oxide from SiO, (v. G. R. 116, 1428).

Zr is generally estimated as ZrO₂; salts of volatilisable acids are decomposed by heat; or solutions may be ppd. by NH,Aq and the pp.

washed, dried, and strongly heated.

Zirconium, alloys of. An alloy of Zr with aluminium is obtained in the preparation of Zr by reducing K₂ZrF₆ by Al (v. Preparation of Zirconium, No. 3). Mellis (Z. [2] 6, 296) obtained crystals having the composition ZrAl, (or perhaps Zr₂Al₆Si) by heating to whiteness a mixture of 1 pt. powdered zirconite with 5 pts. cryolite powder, 10 pts. mixed KCl and NaCl, and 1 pt. Al, treating with dilute HClAq, and separating the larger crystals of the alloy from the finer crystals of Si.

Zirconium, bromide of, ZrBr. Formula probably molecular, from analogy of ZrCl₄. Obtained by mixing finely powdered ZrO2 with sugar charcoal, making into balls with starch paste, drying, and heating to bright redness in a current of dry CO₂ and dry Br (Mellis, Z. [2] 6, 296). ZrBr, is a white, microscopically crystalline, hygroscopic powder; it is easily volatilised; heated in H it is not reduced to a lower bromide. Contact with moist air or water produces the oxybromide ZrOBr₂; evaporation of an aqueous solution gives the heptahydrate of this compound ZnOBr2.7H2O in large needle-

shaped, transparent crystals (M., l.c.). Zirconium, carbides of. By heating an intimate mixture of ZrO₂ and C in the electric arc, in an atmosphere of CO₂, Troost (C. R. 116, 1227 [1893] obtained a very hard, steel-grey solid, approximating closely to the composition ZrC. By using less C, other carbides seem to be produced (cf. Moissan, C. R. 116, 1222), but analyses of other compounds are not given. ZrC2 dissolves easily in dilute HFAq, but is not acted on by other acids. Moissan (l.c.) says that carbides of Zr containing more than 5 p.c. C (ZrC₂ contains c. 21 p.c. C) take fire fairly easily when exposed to the air; Troost says that carbides with a small proportion of C are only superficially oxidised by heating to redness, but that those with much C burn brilliantly at that temperature.

Zirconium, chloride of, ZrCl4. Mol. c. 231.5. V.D. 1.7 at 440° (Deville ... P. 108, 636).

Formation.—1. By heating Zr in Cl.—2. By heating an intimate mixture of ZrO, and C in Cl.-3. By passing vapour of SiCl, over heated ZrO. (Troost a. Hautefeuille, C. R. 75, 1819).

Preparation.—A mixture of ZrO, with c. its own weight of lampblack is made into a paste with oil; little balls, c. the size f hazel-nuts,

are formed of this, and these are covered with powdered charcoal, and heated to full redness in B closed crucible until the oil is completely charred; the pellets are then heated to bright redness in a tube of hard glass, while a current of dry Cl is passed through the tube; the ZrCl, which forms on the cooler parts of the tube is sublimed, in a stream of dry Cl, into a tube drawn out into bulbs, which are sealed off when the operation is finished (v. Bailey, Pr. 46, 74).

Properties, Reactions, and Combinations. A white solid; volatilises unchanged at c. 400°. Dissolves in water, with production of heat; the solution on evaporation gives oxychlorides (q. v.). Does not interact with sinc ethide at 180° (Hinsberg, A. 239, 253). Combines with ammonia to form ZrCl, 4NH,; with sodium chloride to form ZrCl, 2NaCl (Paykull, B. 6, 1467).

Zirconium, forrocyanide of. According to Hornberger (A. 181, 232), the pp. formed by adding K, FeCy, Aq to a solution of a salt of Zr, when washed and dried, has the composition $Zr_s(FeUy_s)_2(?)$; it is a blue powder with a greenish tinge, insoluble in water, decomposed by acids, giving off HCN.

Zirconium, fluoride of, ZrF. Formula pro-

bably molecular, from analogy of ZrCl.

Formation.—1. The hydrated compound ZrF, 3aq is formed by slowly evaporating a solution of ZrO₂H₂, or ZrF, in HFAq.—2. By heating ZrO, with twice its weight of (NH,)HF, to dull redness, till all the ammonium salt is volatilised (Marignac, A. Ch. [3] 60, 263)

Preparation.—Deville (A. Ch. [3] 49, 84) obtained ZrF, by heating to whiteness a mixture of finely powdered sirconite and fluorspar in a stream of HCl (? ZrSiO, + 2CaF, + 2HCl =

 $ZrF_4 + CaCl_2 + CaSiO_3 + H_2O$).

Properties and Reactions.—Prepared as described, ZrF, is a colourless, transparent, crystalline (probably hexagonal) solid; volatilisable at white heat; insoluble in water and acids. ZrF, prepared by heating ZrO, with (NH,)HF, is said to be soluble in water and in dilute HFAq. The crystals of ZrF. Saq are partially decomposed on drying. Heated to full redness

in air, ZrF₄ gives ZrO₂ and HF.

Combinations.—1. With water (v. Formation, No. 1) to form ZrF4.3aq.-2. With various metallic fluorides to form compounds of the form ZrF4.xMF.yH.O. These salts are often called sircono-fluorides, or fluo-sirconates; they are better named sirconi-fluorides. The zirconi-fluorides have been examined chiefly by Marignac (A. Ch. [3] 60, 257); they are obtained either by gradual evaporation of solutions containing the constituent fluorides, or by adding hydroxides or carbonates of the various metals to solutions of ZrF, in HFAq, and evaporating.

Ammonium zirconi-fluorides. salts are described: (1) 2NH₄F.ZrF₄= (NH \ Z_TF, rhombic crystals, i omorphous with K₂ZrF'₆; (2) 3NH₄F.ZrF₄, small octahedra, showing simple refraction (Baker, C. J. 35,

Cadmium zirconi-fluorides. Two salts seem to exist: (1) 2CdF2.ZrF416aq, monoclinic crystals, isomorphous with the Mn salts; (2) CdF, 2ZrF, Caq, laminar, fan-shaped groups of crystals.

Manganese zirconi-fluorides. (1) MnF₂.ZrF₄.5aq; and (2) 2MnF₂.ZrF₄.6aq.

Nickel zirconi-fluorides.

(1) NiF₂.ZrF₄. 6aq, hexagonal prisms; S.G. 2·227 (Topsöe, C. C. 4, 76); and (2) 2NiF₂.ZrF₄.12aq, monoclinic crystals. The first salt forms a double salt with K₂ZrF_e, viz. NiZrF_e.K₂ZrF_e.8aq. Potassium zirconi-fluorides.

(1) K₂ZrF₆; obtained by strongly heating sirconite with two or three times its weight of KHF2, boiling with very dilute HFAq, filtering from K2SiF6, and allowing the filtrate to cool. Crystallises in rhombic crystals, terminated by six-sided pyramids; solubility = 78 at 2°, 1.4 at 15°, 1.7 at 19°, and 25 at 100°. S.G. 3.582 (Topsöe, l.c.); (2) KF.ZrF₄.aq (= KZrF₈.aq); obtained, in monoclinic crystals, by dissolving the preceding salt in water containing a large excess of ZrF. (3) 3KF.ZrF. (= K,ZrF.); obtained by crystallising K,ZrF. from a solution containing excess of KF; small, brilliant, regular octahedra, with simple refraction (v. Baker, C. J. 35, 762). Soluble in water; the hot conc. solution deposits K2ZrF, on cooling.

Sodium zirconi-fluorides. The salt 5NaF.2ZrF, is said to be always formed in whatever proportion the constituent fluorides are mixed. Small, monoclinic crystals; solu-

bility at $18^{\circ} = 38$, and at $100^{\circ} = c$. 1.6.

Zinc zirconi-fluorides. (1) ZnZrF. 6aq; isomorphous with ZnSiF. 6aq and ZnSnF. 6aq, crystallising in regular hexagonal prisms, easily soluble in water. S.G. 2.255 (Topsöe, l.c.). (2) 2ZnF₂-ZrF₄.12aq; deposited from a solution containing excess of ZnF₂, in monoclinic crystals, isomorphous with 2NiF₂-ZrF₄.12aq.

Zirconium, hydride of. By heating ZrO₂

and Mg powder, in the ratio ZrO2:2Mg, in an atmosphere of H, Winkler (B. 24, 888; cf. B. 23, 2666) obtained a black powder which burnt in O, and contained c. 73 p.c. H. On treating this with dilute HClAq, Mg and MgO dissolved, H being given off, and a black residue was left which W. regarded as ZrH. Supposing the reduction of the ZrO, had proceeded in accordance with the equation $ZrO_2 + 2Mg + 2H =$ $2MgO + 2ZrH_2$, the product would have contained 53.44 p.c. ZrH_2 , and, therefore, would have given 1.15 p.c. H; as only .73 p.c. H was found, W. concluded that c. 1 of the ZrO, had been reduced.

Zirconium, hydroxide of, (?) ZrO2H2. Addition of NH, Aq to a solution of a salt of Zr produces a white, bulky, gelatinous pp. of ZrO2.xH2O. According to Berzelius, the dry pp. has the composition ZrO₂.H₂O = ZrO₂H₃; Hermann (J. pr. 97, 318) gave the formula $ZrO_2.2H_2O = ZrO_4H_4$ to the pp. dried at 17.5°; Paykull (B. 12, 1719) said that after drying in vacuo the composition was ZrO2.3H2O. and after drying at 100° ZrO...H.O. A detailed examination of the dehyd ation of the pp. formed by adding NH,Aq to a cold solution of Zr(SO₄)₂ was made by Carnelley a. Walker (C. J. 53, 68, 82 [1888]). The loss if water by the air-dried pp. was fairly regular to c. 90°, when the pp. had approximately the composition ZrO, 2H,O; the loss was then less rapid, but very regular, up to c. 385°, whereat the composition was approximately 2ZrO₂.H₂O; somewhere between 385° and 415° there was a sudden increase in the percentage loss or water

for a rise of 10°, this increase being accompanied by a change of colour from grey to pure white; at 415° the composition was approximately 24ZrO2.H2O; water was then given off very regularly, until at c. 550° dehydration was completed. C. a. W. conclude that probably no definite hydrate of ZrO2, unchanged throughout a fair range of temperature, is formed by heating ZrO_xxH₂O ppd. from a salt of Zr; they think that the change of colour which accompanies the comparatively large percentage loss of water at c. 400° is an accompaniment of a condensation of molecules chiefly of the composition n(2ZrO₂.H₂O) to molecules chiefly of the composition n(24ZrO, H,O). There is a distinct resemblance between the phenomena of the de-hydration of ZrO₂xH₂O and those of the dehydration of SiO₂xH₂O, SnO₂xH₂O, and TiO.xH.O.

Hydrated ZrO, ppd. cold, and washed with cold water, dissolves easily in the ordinary acids, including oxalic acid (Bailey, C. J. 49, 481), forming salts ZrX1, and ZrX112; when ppd. hot, or washed with hot water, the pp. is only slowly dissolved by the ordinary acids. According to Paykull (B. 12, 1719), moist ZrO₂xH₂O absorbs CO, from the air. Digestion with Zr(SO₄)₂Aq produces basic sulphates 3ZrO, 4SO, 15aq and

6ZrO₂7SO₃, 19aq (P., l.c.). ZIRCONATES. ZrO₂xH₂O behaves as a weak acid towards the stronger bases. A few salts have been prepared which are most simply regarded as derived from $Zr(OH)_x$ by replacing H by alkali and alkaline earth metals; these zirconates are obtained by heating ZrO₂ with bases; they have been examined chiefly by Hiortdahl (A. 187, 34, 236 [1865]).

Na₂ZrO₃ Sodium zirconates. (1)(=Na₂O.ZrO₂); obtained by heating ZrO₂ with Na₂CO₂ in the ratio ZrO₂:Na₂CO₃; the crystalline mass thus formed is gradually decomposed by water with separation of amorphous ZrO₂. (2) Na₄ZrO₄ (=2Na₂O.ZrO₂); obtained by heating ZrO₂ with a considerable excess of Na₂CO₃ to whiteness for a considerable time; decomposed by water, with formation of (3) Na₂O.8ZrO₂. 12aq, which forms hexagonal tablets.

Zirconates of calcium and magnesium were obtained by H (l.c.) by heating to redness mix-tures of ZrO₂ with SiO₂ and CaOl₂ or MgCl₂. Ouvrard (C. R. 112, 1444 [1891]) describes lithium sirconate, LitZrO₂, obtained by fusing ZrO, with LiCl, and cooling slowly; when Li2CO, was used, and the fused mass was treated with boiling water containing a little acetic acid, pure

ZrO, remained.
Zirconium, nitride of. In fusing amorphous
Zr with Al, in a cracked lime crucible, Mallet (Am. S. [2] 28, 346 [1858]) obtained a dark-grey, porous solid, which gave yellowish lustrous microscopic cubes when treated with HClAq: the crystals were unchanged by aqua regia or alkali solutions; when fused with KOH they gave off NH. The yellow crystals were probably a nitride of Zr.

Zirconium, oxides of. Besides zirconia, ZrO, there exists a higher oxide, probably ZrO,; and, possibly, also a lower oxide (?) ZrO.

ZIRCONIUM DIOXIDE ZrO₂. (Zirconia. Zirconio anhyoride.) Mol. w. not known.

Preparation.—There are many methods for

preparing ZrO, from sirconits. 1. The mineral is heated to redness and thrown into water, and the least-coloured pieces are finely powdered and levigated; the powder is very strongly heated with four times its weight of Na₂CO₃, in a Pt crucible, and the product is heated with excess of HClAq; the whole is evaporated to dryness; the residue is digested with water, filtered from SiO₂, and ZrO₂xH₂O is ppd. by NH₂Aq. The pp. is washed and dissolved in HClAq, and the solution is nearly saturated with Na2COs, and boiled with Na₂S₂O₄Aq as long as SO₂ is given off, whereby ZrO₂xH₂O is ppd. (Berzelius, P. 4, 124; 8, 186; Hermann, J. pr. 31, 77; Chancel, A. 108, 237; Stromeyer, A. J13, 127).-2. Very finely powdered sirconite is boiled with conc. HClAq, and the residue is washed with water and dried; 100 g. of the dried powder are added to 400 g. molten NaOH, with which 20 g. NaF have been mixed, and the whole is heated in a nickel crucible until the action ceases, when the molten mass is poured on to a sheet of nickel, and, after solidification, is plunged into water; the portion insoluble in water is dissolved in HClAq, the solution is evaporated to dryness, and treatment with HClAq and evaporation to dryness are repeated reveral times to remove SiO₂ and HF. The solution in HClAq is then ppd. by NH₃Aq; the ZrO₂. α H₂O is thoroughly washed by decantation, dissolved in the smallest possible quantity of conc. HClAq, and the solution is evaporated to dryness; the residue is washed in a funnel with conc. HClAq mixed with 4 pts. of alcohol; the washed solid is then repeatedly crystallised from conc. HClAq. The white, lustrous crystals of ZrOCl₂αH₂O thus obtained are dissolved in water, and ZrO₂αH₂O is ppd. by NH₃Aq (Venable, C. N. 64, 315 [1891]; cf. Bailey, Pr. 46, 74 [1889]; cf. Preparation of ZIRCONIUM, p. 893). By heating ppd. ZrO_{2.}xH₂O to c. 550°, ZrO₂ is obtained.

Crystalline sirconia is obtained from the amorphous form (1) by fusing with borax at a very high temperature, and washing with HClAq (Nordenskjold, P. 114, 612; Knop, A. 157, 868); (2) by fusion with microcosmic salt (K., l.c.); (3) by heating to dull redness in dry HOl at a pressure of three atmospheres (Hautefeuille a. Perrey, C. R. 110, 1038); (4) by melting and volatilising in an electric furnace, using a current of 860 ampères and 70 volts (Moissan, C. R. 116, 1222 [1898]); (5) by heating, to dull redness, a salt of Zr with an acid that can be removed by heating, in a current of dry HCl (H. a. P., l.c.).

Properties.—Amorphous zirconia is a white, tasteless, inodorous powder; S.G. c. 5; insoluble in soids, except in conc. H.SO, (v. Reactions, No. 7) or HFAq. Crystalline zirconia is a transparent, colourless solid; S.G. 5-1 (Moissan, *l.o.*), 5-42 (Knop, *l.o.*), 5-7 (Nordenskjold, *l.o.*). The crystals are described by N. (*l.o.*) as quadratic prisms, isomorphous with SnO2 (tinstone) and TiO2 (rutile). The crystals are extremely hard; they scratch glass easily (Moissan, l.c.). It is generally said that when ZrO, is heated to incipient redness it glows brightly, and becomes denser; but the experiments of Carnelley a. Walker (C. J. 58, 82 [1888]) on the action of heat on ZrO₂xH₃c

make it probable that the glowing is an accompaniment of the dehydration from approximately 2ZrO₂.H₂O to approximately 24ZrO₂.H₂O. conia reacts towards acids as a basic oxide, forming salts ZrXI4 and ZrXII2 (many of which are basic salts); it also decomposes alkali carbonates, and some other alkali and alkaline earth-salts, when fused therewith, forming zirconates mZrO₂nM₂O (v. Zirconates, p. 896).

Reactions.—1. When ZrO₂ is heated to a very high temperature it melts, boils, and volatilises, condensing in crystals; Moissan (C. R. 116, 1222 [1893]) effected the volatilisation in a crucible heated in the electric furnace, using a current of 360 ampères and 70 volts. -2. When very strongly heated with excess of carbon, carbides are formed (v. Carbides, p. 894). Heating in a carbon crucible in the electric furnace: produces Zr (M., l.c.).-3. Heating with magnesium proc 10es Zr, perhaps also an oxide lower than ZrO₂ (Winkler, B. 23, 2642; 24, 888; Phipson, J. pr. 96, 85£; Bailey, Pr. 46, 74; cf. Zirconium monoxide). When heated with magnesium in an atmosphere of hydrogen, some ZrH₂ is perhaps produced (v. Zirconium hydride, p. 895).—4. ZrCl₄ is formed by heating ZrO2, mixed with carbon in a stream of chlorine; using bromine produces ZrBr, (v. Bromide and Chloride, p. 894). — 5. ZrF, is formed by heating ZrO, to dull redness with excess of ammonium fluoride.—6. K2ZrF, is formed by slightly heating ZrO2 with excess of potassium hydrogen fluoride, boiling with very dilute HFAq, and filtering.—7. $Zr(SO_4)_2$ is obtained by heating ZrO, with excess of a mixture of 2 pts. conc. sulphuric acid and 1 pt. water.

ZIRCONIUM PEROXIDE. By adding H2O2Aq and NH, Aq to a solution of Zr(SO₄)₂, Clève (Bl. [2] 43, 53) obtained an oxide containing more Othan ZrO2; to this oxide he gave the formula ZrO. Bailey in 1886 (C. J. 49, 149, 481) examined the pp. obtained by adding H2O2Aq to solution of Zr(SO₄)₂; according to B., the moist pp. has the composition Zr₂O₅, and this composition is unchanged after keeping (moist) for some months. In Bailey's paper in 1889 (Pr. 46, 74) he gives the composition ZrO, 3H₂O to the oxide ppd. by H₂O₂Aq, either from an acid or an alkaline solution of Zr(SO₂), after drying over P₂O₅; and the composition Zr₂O₅ after drying at 100°. The oxide ZrO₇xH₂O was obtained (C. J. 49, 485) by preparing a solution of $Zr(SO_1)_2$ by heating K_2ZrF_4 with H_2SO_4 , and at once adding H_2O_2Aq . Zr peroxide is insol. in very dilute (1 p.c.) H, SO, Aq or H.C, H, O, Aq (B., l.c.). Bailey (l.c.) employed the ppn. of ZrO, xH, O to separate Zr from Fe, Nb, and Ti.

ZIBCONIUM MONOXIDE (?) ZrO. By heating ZrO, and powdered Mg, in the ratio ZrO, Mg, in a stream of H, Winkler (B. 23, 2668 [1890]) obtained a black powder, which he allowed to cool in H, then kept for some time in CO, freed from Ma by treatment with dilute HClAq, washed with water containing HCl, then with alcohol, and finally with ether, and dried in vacuo. He thus obtained a deep-black powder, which was un-acted on by HClAq, HNO, Aq, or H, SO, Aq, even on warming, and which burnt to ZrO, when heated in air. W. supposed this black powder | Zr(SCN), remained in solution. to be ZrO. Vol. IV.

Zirconium, oxyacid of. Hydrated zirconia, ZrO₂xH₂O, reacts as a weak acid towards strong. bases (v. ZIRCONATES, p. 896).

Zirconium, oxybromide of, ZrOBr₂.xH₂O. ZrBr, dissolves in water; on evaporation, transparent needle-shaped crystals are formed, to which Mellis (Z. [2] 6, 296) gave the composition $ZrOBr_2.7H_2O$. By dissolving $ZrO_2.xH_2O$ in HBrAq and evaporating, Weibull (B. 20, 1894) [1887]) obtained lustrous needles of ZrOBr. 8H.O. isomorphous with ZrOCl₂.8H₂O.

Zirconium, oxychlorides of. Various oxychlorides have been described. (1) $ZrOCl_2xH_2O$; prepared by dissolving ZrO_2xH_2O in HClAq and evaporating (for details as to the best conditions v. Weibull, B. 20, 1394). Hermann (J. pr. 31, 77) gave the formula as $ZrOCl_2.9H_.O$; Mellis (Z. [2] 6, 296) gave $2ZrOCl_2.9H_.O$; and Weibull (Lc.) $ZrOCl_x.9H_.O$ (v. also Paykull, B. 6, 1467; and Bailey, Pr. 46, 74). (2) Zr_2OCl_1 ; prepared by Troost a. Hautefeuille (C. B. 73, 563) by passing steam and vapour of ZrCl, through a red-hot tube. (8) 2Zr.O.Cl..15H.O; prepared by Endemann (J. pr. [2] 11, 219) by adding ether to a warm alcoholic solution of ZrOCl..xH.O.

Zirconium, oxyiodides of. By adding the calculated weight of BaI2, in solution, to Zr(SO,)2 dissolved in a slight excess of H2SO, Aq, filtering, evaporating over H2SO, and removing free I from the residue by CS₂, Hinsberg (A. 239, 253 [1887]) obtained a white, amorphous powder, easily soluble in water, giving off I in moist air. to which he gave the formula ZrIO, H, xH2O (x probably = 3).

Zirconium, oxysulphide of: v. ZIRCONIUM SULPHIDES (infra).

Zirconium, salts of. The salts formed by the replacement of H of acids by Zr belong to the forms ZrX₂¹¹ and ZrX₄¹; many basic salts are known. The following are the chief salts of Zr derived from oxyacids: -Arsenates, borates, carbonates, nitrates, oxalates, phosphates, selenites, silicates, sulphates, and sulphites.

Zirconium, silicofluoride of, (?)ZrSiF.; probably obtained by Berzelius by evaporating a solution (? of ZrO, xH,O in H,SiF,Aq) (v. Lehr. buch, 3, 505 [5th ed.])

Zirconium, sulphides of. According to Berzelius (v. Lehrbuch, 2, 189 [5th ed.]), a compound of Zr and S is formed by heating the elements together in vacuo, or with H. The elements together in vacuo, or with H. compound, to which no formula is assigned, is described by B. as a dark-brown coloured powder; unacted on by H,SO,Aq, HNO,Aq, or HClAq; slowly attacked by boiling aqua regia; dissolved easily by HFAq, giving off H₂S; insol. in KOHAq, and decomposed by fusion with KOH to K,S and ZrO₂. By strongly heating ZrO₂ in vapour of CS₂, Fremy (A. Ch. [8] 88, 326 [1858]) obtained a graphite-like solid, which was rapidly acted on hy HNO,Aq, with separation of S; Fremy supposed that the substance might be an oxysulphide.

Zirconium, sulphocyanide of. By adding Ba(SCN), to Zr(SO,), solution, Hornberger (A. 181, 232 [1876]) obtained a solution which, he said, contained Zr(SCN),; this solution quickly decomposed, depositing a yellow solid, while

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ADDENDA.

[DESCRIPTIVE INORGANIO CHEMISTRY ONLY.]

ALUMINIUM (vol. i. p. 141). Preparation. By electrolysing a molten mixture of AlF. 3NaF with NaCl (Minet, C. R. 111, 603 [1890]). The best results are obtained by using the salts in the ratio AIF .. 3NaF:6NaCl; details of the quantities to be used, the strength of current, &c., are given by M.

Properties.—Minet (C. R. 111, 603 [1890]) gives m.p. of Al as 675°; and S.G. at 820° as 1.76. The S.H. of Al is given by Richards (C. N. 65, 97 [1892]) as .224 at 20°, .232 at 1005, and $\cdot 2845$ at the m.p. = 625° ; the S.H. of molten Al at $650^{\circ} = .808$, according to Pionchon (C. R. 115, The molecule of Al in dilute solu-162 [1892]). tion in tin is perhaps diatomic (Heycock a.

Neville, C. J. 57, 376 [1890]).

Reactions.—Volatilises in electric furnace, with current of 250 ampères and 70 volts (Moissan, C. R. 116, 1429 [1893]). Not appreciably acted on by NO, at 500° (Sabatier a. Senderens, C. R. 115, 236 [1892]). An amalgam is formed by bringing Al into a solution of a mercuric salt (Klaudy, C. C. 1893 [1] 201; abstract in C. J. 64 [11] 376 [1893]). Heated with NaPO, in a current of H, from 28 to 31 p.c. of the P distils over; P is also obtained by heating Al with any phosphate of Ca or Mg (Rossel a. Frank, B. 27, 52 [1894]). According to Lunge (C. N. 65, 110 [1892]), vessels made of Al may be used for holding articles of food, at least at the ordinary temperature, without danger of the metal being dissolved; this result is confirmed by Balland (C. R. 114, 1536 [1892]).

Aluminium, alloys of (vol. i.p. 143). For an alloy with gold, Al, Au, v. Heycock a. Neville (C. J.

65, 74 [1894]).

Aluminium, chloride of (vol. i. p. 144). Nilson a. Pettersson (Z. P. C. 4, 206 [1889]) made a number of determinations of the V.D. of AlCla; they find that the compound gradually dissociates above its b.p. until c. 800°, when its V.D. becomes constant and corresponds with the formula AlCl₂. AlCl₂ may be prepared by heating an alloy of Al and Cu nearly to redness in dry HCl; AlCl, distils over unmixed with any Cu compound (Mabery, B. 22, 2658 [1889]). Sabatier (Bl. [3] 1, 88 [1889]) says that crystals of the hydrated chloride after being in vacuo for three months at 20° over H2SO4 have the composition AlCI, 6H,O.

Aluminium, exides of (vol. i. p. 146): By heating Al₂O₃ with Mg, in the ratio Al₂O₃:Mg, Winkler (B. 23, 780 [1890]) obtained a brownishblack powder which, he says, contained & considerable quantity of an oxide lower than Al.O. probably AlO. By the incomplete combustion of Al, Pionchon obtained a dark-grey powder, said by P. to have the composition $Al_6O_7 = 2Al_2O_3 \cdot Al_2O_3$ J. R 117, 828; abstract in O. J. 64 [11] 572 [1893]). By heating amorphous Al,O, to in-

cipient redness in HCl gas at a pressure of three atmos., Hautefeuille a. Perrey (C. R. 110, 1038 [1890]) obtained crystals of corundum. According to Read (C. J. 65, 818 [1894]), Al₂O₈ is unchanged at c. 1750°.

Aluminium, phosphide of (vol. i. p. 146). A grey crystalline powder, Al, P, was obtained by Rossel a. Frank (B. 27, 52 [1894]) by heating Al in vapour of P, and then heating the product

till P ceased to come off.

AMMONIA (vol. i. p. 196). Formation.— According to Loew (B. 23, 1443 [1890]), traces of NH₃ are formed by shaking Pt black with very dilute NaOHAq in the air. NH_s is also formed by the combined action of Pt black and dextrose

on KNO3Aq (L., l.c., p. 675).

Properties.-Ludeking a. Starr (Am. S. [8] 45. 200 [1892]) give S.H. of liquid NH, as 8857. For measurements of the wave-lengths of the lines in the emission spectrum of NH, v. Magnanini (Z. P. C. 4, 435 [1889]). Perkin (C. J. 55, 689, 728 [1889]) gives M. M. of NH, in aqueous and alcoholic solutions; also Mol. R. of various NII, salts in solution. Tables showing the solubility of NH, in alcohol of various concentrations, at 10°, 20°, and 80°, are given by Delépine (J. Ph. [5] 25, 496; v. abstract in C. J. 62, 1049 [1892]). For the freezing-points of aqueous solutions of NH, v. Pickering (C. J. 63. 181 [1893]).

Reactions. - Dry NH, does not combine with dry hydrogen chloride when these gases are mixed (Baker, C. J. 65, 611 [1894]). Nor do dry NH, and dry carbon dioxide combine (Hughes a. Soddy, C. N. 69, 138 [1894]). Michel a. Grandmongin (B. 26, 2565 [1893]) say that the products of passing NH, over heated peroxide of Ba, Pb, Mn, or Na, are N with a little HNO2 and HNO3, and oxide or hydroxide of the metal. The reactions of liquid NH, with CrCl, and FeCl, are described by Christensen (Zeit. anorg. Chemie, 4, 227; abstract in O. J. 64 [11] 469 [1893]).

AMMONIUM COMPOUNDS (vol. i. p. 200). Ammonium amalgam (vol. i. p. 201). From measurements of the polarisation given by this substance in a solution of NH, Cl against zinc amalgam, and a comparison of these measurements with the polarisations given by metallic amalgams, Le Blanc concludes that ammonium amalgam really exists (Z. P. C. 5, 467; abstract in C. J. 58, 1204 [1890]).

Ammonium chloride (vol. i. p. 202). Baker (C. J. 65, 611 [1894]) asserts that the V.D. of the dry salt at 350° is 28.7, corresponding with that required by the formula NH₄Cl. B. also says that no reaction occurs when dry NH Cl and

dry CaO are heated together.

ANTIMONY (vol. i. p. 282). Reactions .-Montemartini (G. 22, 884, 426) finds that no ARSENIC. 899

appreciable quantity of NH, is formed by the interaction of Sb and HNO, Aq varying from 2 to 70.8 p.c. HNO₂; NO₂ is practically the only gaseous product of the reaction; when 70 p.c. HNO.Aq is used, the residue probably has the

composition (SbO)NOs.

Antimonates (vol. i. p. 285). Several salts have been prepared, by double decomposition from the K salt, by Peilstein and O. v. Bläse (C. C. 1889. 803; abstract in C. J. 56, 1123). All the compounds are salts of HSbO,; B. a. B. conclude that the only well-marked antimonates are derived from this acid. Ebel (B. 22, 3044 [1889]) describes several metantimonates.

Antimonites (vol. i. p. 285). For antimonites of K and Na v. Cormimbouf (C. R. 115, 1805;

abstract in C. J. 64 [11], 171 [1893])

Antimony, bromide of (vol. i. p. 286). The b.p. of SbBr, is given as 275° by Freyer a. V. Meyer (Zeit. f. anorg. Chemie, 2, 1 [1892]).

Antimony, trichloride of (vol. i. p. 286). Boils at 228° (Freyer & V. Meyer, L.c.).

Antimony, pentachloride of (vol. i. p. 287). The V.D. at 218° and 58 mm. pressure was found to be 144.7 (corresponding with the formula SbCl_s) by Anschütz a. Evans (A. 25^, 95 [1890]). Double compounds of SbCl_s with RbCl and CsCl and described by Saunders (Am. 14, 152 [1892]), and by Wheeler (Am. S. [3] 46, 269 [1893]).

Antimony, fluoride of (vol. i. p. 287). The double compound SbF, 2AmF is described by H. von Helmholt (Zeit. f. anorg. Chemie, 3, 115

[1893]).

Antimony, hydride of (vol. i. p. 288). temperature whereat decomposition of SbH, begins is given by Brunn (B. 22, 8202 [1889]) as c. 150°. For the reactions of SbH, with AgNO, Aq v. Vitali (abstract in C. J. 64 [11], 206 [1893]).

Antimonic oxide (vol. i. p. 290). When heated to dull redness Sb₂O₅ gives off O, forming Sb₂O₄, which at c. 1750° is further reduced to

Sb₂O₃ (Read, C. J. 65, 313 [1894]).

Antimonious sulphide (vol. i. p. 291). Black Sb,S, is formed, according to Mitchell (C. N. 67, 291 [1893]), by passing CO₂ into boiling water with ordinary Sb.S. in suspension. Picton (C. J. 61, 142 [1892]) obtained a colloidal Sb₂S₃ in very fine suspension in water, by allowing solution of tartar emetic to flow into saturated H2SAq.

Antimony, thio-chlorides and thio-iodides of (vol. i. pp. 292-3). Ouvrard (C. R. 116, 1516 [1893]) describes SbSCl and Sb₄S₆Cl₂, obtained by the action of H2S on SbCl.; and SbSI formed by heating SbI, to 150° in dry H2S, and Sh.S.I. formed by heating Sh.S. with I and

crystallising from CS2.

ARSENIC (vol. i. p. 301). A full account of the different forms of As obtained, along with As O₃, by subliming As in a tube is given by Return (Z.P. C. 4, 408; abst.act in C. J. 64 [11], 570 [1893]). Petersen gives the following heats of oxidation, and atomic volumes, of the allotropic forms of As (Z. P. C. 8, 601 [1891]):—

At. volume As O 156,830 Grey, rhombohedric 18.1 Grey-black amorphous 154,840 15.94 163,500 15.99 Brown, amorphous

Arsenic, bromide of (vol. i. 809). combines with NH, to form AsBr. 3NH, (Besson, C. R. 110, 1258 [1890]).

Arsenates (vol. i. p. 307). For arsenates of Cd v. A. de Schulten (Bl. [8] 1,478 [1889]); for arsenates of Cu, and of Cu and Na, v. Hirsch (C. C. 1891 [1]. 15; abstract in C. J. 60, 644); many double arsenates of K, and of Na, with Cd, Co, Mg, Mn, Ni, and Zn are described by Lefèvre (C. R. 110, 405 [1890]). For thio-arsenates of Cu, Hg, Zn, and Zn and

Na, v. Preis (A. 257, 178 [1890]).

Arsenites (vol. i. p. 306). For the preparation, by double decomposition from the salt 2KAsO₂.As₂O₃, and properties, of the arsenites of the following metals v. Reichard (B. 27, 1019 [1894]):—Al, Ba, Cd, Ca, Cr, Co, Cu, Au, Fe, Pb, Mg, Mn, Hg, Ni, Pd, Pt, Ag, Sr, Sn, Ti, U, and Zn (references and criticisms of former investigations are given). Stavenhagen (Zeit. f. angew. Chemie, 1894. 165) obtained the double salt of K arsenite and sulphate, K.AsO.10K.SO. by evaporating solutions of the constituents after mixing.

Arsenic acid (vol. i. p. 307). For a detailed account of the reaction of H.S with solutions of H. AsO, v. Brauner a. Tomeetchek (C. J. 53, 145

[1888]).

Arsenious chloride (vol. i. p. 310). According to Besson (C. R. 109, 940 [1889]), AsCl, solidifies at -18°, forming white needles; at -30° AsCl, absorbs much Cl, forming a liquid which does not solidify at -60° , and which with water gives off much Cl and forms As₂O₂Aq. Besson (C. R. 110, 1258 [1890]) says that the compound formulated by Rose as 2AsCl,7NH, is really AsCl. 4NH.

Arsenious fluoride (vol. i. p. 310). Moissan (A. Ch. [6] 19, 280 [1891]) prepared AsF, by heating As2O, with HF; he gives b.p. as 63° at 750 mm., and says that it solidifies to white crystals at -8.5°. Besson (C. R. 110, 1258) [1890]) describes the compound 2AsF.5NH.

Arsenic, trihydride of (vol. i. p. 310). Brunn (B. 22, 3202 [1889]) finds that the first products of the action of air, or O, on AsH, are solid nAsH and H2O, and the next products are As and H₂O. AsH₃ and H₂S do not react in absence of air at the ordinary temperature, either as gases or in solution (B., l.c.). AsH, begins to decompose at c. 230° (B., l.c.). For the reactions of AsH, with AgNO, Aqv. Vitali (abstract in C. J. 64 [11] 206 [1893]).

Arsenious iodide (vol. i. p. 311). Besson (C. R. 110, 1258 [1890]) describes the com-

pounds AsI, 4NH, and AsI, 12NH,

Arsenious sulphide (vol. i. p. 315). Picton (C. J. 61, 140, 144 [1892]) obtained an aqueous solution of colloidal As, S, containing also some very finely divided sulphide in suspension, by allowing As2O,Aq to flow into saturated H.SAq, into which H2S was continually passing, and removing H2S by a stream of H

Arsenic pentasulphide (vol. i. p. 815) As₂S₅ is obtair 3d, according to McCay (Am. 12, 547 [1891], by passing H₂S for a long time into a hot solution of an alkali dihydrogen arsenate,

and then prg. by a mineral acid.

Arsenic, thiochlerides and thio-iedides of L. i. p. 816). Ouvrard (C. R. 116, 1516 [1898]) obtained As S.Cl. by passing H. into

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Asol, and washing the pp. with CS₂. By heating 1 pt. As,S, with 10 pts. Asol, to 150° he obtained AsSCl

AZO-IMIDE (v. Hydrasoic acid, p. 919, Addenda).

BARIUM (vol. i. p. 440). Richards, in 1893, made an elaborate experimental revision of the at. w. of Ba (P. Am. A. 28, 1; 29, 55). As a mean result of the most trustworthy experiments, R. adopts 137.1 as the at. w. of Ba (O=15.96). Limb (C. R. 112, 1434 [1891])failed to obtain any Ba by electrolysing fused BaCl₂, or BaCl₂ mixed with NaCl.

Barium, carbide of. By heating a mixture of Ba amalgam and powdered charcoal in H, Maquenne (C. R. 114, 361 [1892]) obtained grey BaC₂; decomposed by water, giving BaO₂H₂Aq and C₂H₂; v. also Moissan, C. R. 118, 688 (abstract in C. J. 66 [1], 314 [1894]).

Barium, chloride of (vol. i. p. 441). By bringing BaCl, into contact with liquid NH, and allowing the gas to escape at 0° and 760 mm., Joannis obtained BaCl. 8NH, (C. R. 112, 337

[1891]).

Barium, fluoride of (vol. i. p. 441). BaF₂ is obtained in microscopic crystals by adding BaCl.Aq (3.6 p.c.) to boiling KFAq (1 p.c.), boiling, washing, drying, and heating strongly (Moissan, Bl. [3] 5, 152 [1891]). Poulenc (C. R. 116, 987 [1893]) obtained BaF₂ in wellformed octahedra by fusing amorphous BaF, with KHF, and KCl.

Barium, nitrides of. Maquenne (C. R. 114, 220 [1892]) obtained a mixture, probably containing some Ba, N2, by heating Ba amalgam to redness in a current of N. Berthelot a. Matignon (A. Ch. (7) 2, 144 [1894]) say that BaN, is formed by heating AmN, Aq with BaO, H, Aq.

Barium monoxide (vol. i. p. 442). BaO is unchanged at c. 1750° (Read, C. J. 65, 313 [1894]); it liquefies at c. 2,000°, but is not decomposed at c. 2,500° (Moissan, C. R. 115,

1034 [1892]).

Barium dioxide (vol. i. p. 443). BaO, is decomposed by salts of Al, Cr, Co, Fe, &c.; some salts—e.g. those of Ca, Li, K, Na—do not react with BaO₂ (v. Kwasnik, B. 25, 65). For decomposition of BaO₂ by heat v. Le Chatelier (C. R. 115, 654; abstract in C. J. 64 [11], 71 [1893]).

BERYLLIUM (vol.i. p. 504). Krüss a. Moraht (B. 23, 727 [1890]) obtained hexagonal crystals of Be, mixed with BeO, by heating BeF₂.2KF to redness with Na. K. a. M. (l.c. p. 2552) obtained the value 9.03 for the at. w. of Be by converting BeO into BeSO4.4H2O, and this again to BeO. Gibson (C. J. 63, 909 [1893]) describes methods for preparing BeO from beryl.

Beryllium, fluoride of (vol. i. p. 506). Petersen (Z. P.C. 5, 263 [1890]) gives the thermal data H.F. $[BeO^2H^2, 2HFAq] = 19,683$. The double compound BeF. 2AmF is described by H. von

Helmholt (Zeit. f. anorg. Chemie, 3, 115 [1893]).

Beryllium, oxide of (vol. i. p. 506). By dissolving BeO. xaq in KOHAq, Krüss a. Moraht (B. 23, 727 [1890]) obtained potassium beryllats BeO₂K₄ (?), mixed with K₂CO₃, as a snow-white, silky solid

BISMUTH (vol. i. p. 509). The at. w. of Bi was re-determined by Classen in 1890 (B. 23, 938 [1890]), by converting Bi into nitrate, and

this into Bi₂O₃; the mean result was 208.38 (O = 15.96). Schneider (J. pr. [2] 42, 553; 44,23 [1891]) criticises Classen's results, and maintains that Marignac's value (almost exactly 208) is as trustworthy as that obtained by C. The molecule of Bi in dilute solution in tin is perhaps diatomic (?) (Heycock a. Neville, C. J. 57, 376 [1890]).

Preparation .- Classon (l.c.) says that pure Bi is best prepared by electrolysing a solution of the metal in HNO, Aq, using a cone of Pt as the negative electrode, then washing with alco-

hol, and fusing with KCy.

Properties. - Melts at 264°, according to Classen (l.c.); at 269-22°, according to Callendar a. Griffiths (C. N. 63, 1 [1801]). S.G. 9.7474 C., l.c.; no temperature given); 9.787 at $\frac{6}{40}$, 9.673 solid at the m.p., 10.004 liquid at the m.p. (Vicentini, Rend. Acad. Lincei, 6 [11], 121, 147 [1891]). For S.G. of liquid Bi from 235° to 280° v. Cattaneo (Rend. Arad. Lincei, 7 [11], 88 [1892]). For electrical resistance v. E. v. Aubel (C. R. 108, 1102 [1889]).

Reactions.—According to Veley (Pr. 48, 458 [1891]), Bi does not dissolve in 30 p.c. HNO,Aq at 30° if HNO, is absent; but Bi dissolves readily in 1 p.c. HNO,Aq.

Bismuth, bromide of (vol. i. p. 511). BiBr₃ boils at 453° (V. Meyer, A. 264, 122 [1891]). Cavazzi a. Tivoli (G. 21 [11], 306 [1892]) describe a lustrous, black, hygroscopic solid, PBrH(BiBr₂)₃, obtained by allowing BiBr, to fall gradually into PH, (abstract in C. J. 62, 279).

Bismuth, chloride of (vol. i. p. 512). Boils

at 435°-441° (uncor.) (V. Meyer, $l.\bar{c}$.). The only double chloride of Bi and K of those hitherto described that exists is BiCl₃.2KCl. 2aq, according to Brigham (Am. 14, 164 [1892]). B. has also prepared BiCl₃.KCl. aq.

Bismuth, fluoride of (vol. i. p. 512). H. von Helmholt describes the double fluoride BiF. AmF

(Zeit. f. anorg. Chemie, 3, 115 [1893]).

Bismuth, iodide of (vol. i. p. 513). (C. R. 110, 525 [1890]) obtained the double iodide 2BiI₃.KI by keeping I, KCl, Bi, and water in contact for some weeks, and then crystallising from Et acetate.

Bismuthous oxide (vol. i. p. 513). Classen (B. 23, 938 [1890]) gives S.G. as 9 044. Read (C. J. 65, 313 [1894]) found Bi₂O₂ to be unchanged

at c. 1750°.

Bismuth, exysulphide of (vol. i. p. 516). By passing dry H.S into boiling benzene containing Bi₂O₅ in suspension, P. v. Scherpenberg (C. C. 1889. [11] 643) obtained Bi₂O₂S.

Bismuth, selenide of (vol. i. p. 516). The double compound Bi₂Se, K₂Se is described by P.

v. S. (l.c.).

Bismuth, sulphide of (vol. i. p. 516). P. v. S. (l.c.) obtained the double compound Bi₂S₂.K₂S

by fusing Bi₂O₅ with K polysulphide.

BORON (vol. i. p. 524). The at. w. of B has been determined by Abrahall by the annaysis of BBr_s (C. J. 61, 650 [1892]); the value obtained was 10.67 (O = 15.96). Also by Ramsay a. Aston (C. J. 63, 207 [1893]) by finding the percentage of water in crystallised borax, and by d'stilling fused borax with HClAq and CH. OH and weighing the residual NaCl; the values varied about 11, but showed considerable differences. BORON. 901

Rimbach (B. 26, 164 [1893]), by titrating solutions of borax with standard HOlAq, using methyl orange as indicator, obtained the value 10.91 (0=15.96). The following values for S.H. of amorphous B are given by Moissan a. Gautier (C. R. 116, 924 [1893]): 0° to $100^\circ = 3066$, 0° to $192^\circ = 3407$, 0° to $234^\circ = 3573$. For atomic refraction of Rv. Ghirs (G. 23 [1], 452; abstract in C. J. 64 [11], 517 [1893]).

Preparation.—According to Moissan (C. R. 114, 319 [1892]), the methods employed to give amorphous boron yield substances containing from c. 45 to c. 72 p.c. of B. M. (l.c. p. 392) says that nearly pure amorphous B is obtained by heating to redness a mixture of one pt. powdered Mg with 3 pts. fused B₂O₂, washing with HOlAq, KOH in alcohol, HFAq, and water (the process is tedious, details are given); by fusing this product with 50 pts. B₂O₂, and washing with HOlAq &c., a brown powder is obtained containing c. 98 5 p.c. B, with traces of Mg and small quantities of other impurities.

Mg and small quantities of other impurities.

Properties and Reactions.—The properties of amorphous boron are described by Moissan (C. R. 114, 617 [1892]). Brown powder; S.G. 2.45. Does not fuse in electric arc, but volatilises (M., C. R. 17, 423 [1893]); increases in density when heated to c. 1500° in H. Specific electrical resistance = 801 megohms. Burns in air at c. 700°. Combines directly with Br, Cl, (not I), N at c. 1200°, Se, S, (not Te); also with Al, Fe, Mg, Pt, and Ag. Reduces many oxyacids and oxides when heated therewith; HClO, HIO, HNO,, and H₂SO, are readily reduced; As₂O, and As₂O₅, CO, N₂O, P₂O₅, SiO₂, steam, and SO₂ are also reduced; many metallic oxides are reduced to metals, often with explosive violence; molten KOH gives off H; metallic fluorides are generally decomposed, giving BF_s; many other metallic salts, both in solution and when heated as solids, are reduced by amorphous boron.

Boron, bromède of (vol. i. p. 525). BBr, boils at 90° (Freyer a. V. Meyer, Zeit. f. anorg. Chemie, 2, 1). Ghira (Z. P. C. 12, 768 [1893]) gives S.V. 94·72; and S.G. $\frac{0}{60}$ = 2·64985. According to Besson (C. R. 112, 1001 [1891]), when a mixture of gaseous HI and BBr, is passed through a glass tube at 300°-400° the products are BBr, I, BBrI, and BI, BBr, I and BBrI are colourless liquids, distilling at c. 125° and 180°; they are rapidly decomposed by air and light. Tarible (C. R. 116, 1521 [1893]) says that BBr, combines with PBr, and PBr, to form compounds easily decomposed by water, Cl, or NH, in the cold. Besson (C. R. 114, 542 [1892]) describes BBr, 4NH, obtained by passing dry NH, into BBr, in dry CCl, at 0°.

Boron, carbide of, B_cC. Moissan (C. R. 118, 556 [1894]) says that B and C combine when amorphous B is mixed with c. I its weight of sugar charcoal, and the mixture is heated in the electric furnace for some minutes, using a current of 250-300 ampères and 70 volts. After treating the product with HNO₂Aq, and then with KOlO₂ and HNO₂Aq, the carbide remains as a very hard, lustrous, black, crystalline powder; S_cC. 2·51; burns very slowly in O at c. 1000°; acted on by Cl below 1000°, but not by Br, I, S, P, N, or mineral acids; reacts with molten KOH or KNaCO₂ (a3stract in C. J. 66 [11], 279 [1894]).

Boron, chloride of (vol. i. p. 525). BCl₄ boiling at 17° (Freyer a. V. Meyer, Zeit. f. anorg. Chemie, 2, 1). Ghira (Z. P. C. 12, 768 [1893]) gives S.G. $\frac{A}{2}$ = 143886, and S.V. 81 94.

Boron, fluoride of (vol. i. p. 526). According to Besson (C. R. 110, 80 [1890]), BF, unites with PH, at c. -30° to form 2BF, PH, decomposed by water giving off H and PH,

Ammonium borofluoride is described by Stolba (C. C. 1890. [1] 211; abstract in C. J. 58, 560).

Boron, hydride of (vol. i. p. 526). Sabatier (C. R. 112, 865 [1891]) says that the gas produced by adding HClAq to the brown powder formed by heating B₂O₂ with Mg in H has a very disagreeable odour, burns with a green flame, gives a brown pp. with AgNO₂Aq, deposits B when passed through a glass tube at a dull red heat (H coming off), and is decomposed by electric sparks, giving pure H. The gas contains a little B hydride, according to S. (l.c.); of. Winkler (B. 23, 772 [1890]).

Boron, iodide of (vol. i. p. 527). BI_s. Prepared by Moissan (C. R. 112, 717 [1891]), (1) by heating I and B to 700°-800°; (2) by heating amorphous B (dried in H at 200°), in a stream of dry HI, in a tube of hard glass until the glass nearly softens; (3) by passing HI and BOl, vapours through a porcelain tube heated to redness. The product is dissolved in CS₂, shaken with Hg, and allowed to crystallise. Colourless crystals, very hygroscopic, rapidly become coloured in light. Melts at 43° and boils at 210°. Decomposed by water, giving HIAq and H₂BO₂Aq. (For other reactions v. abstract in C. J. 60, 979 [1891].) For compounds with NH, v. Besson (C. R. 114, 542 [1892]; abstract in C. J. 62, 771).

Boron, oxide of (vol. i. p. 527), B₂O₃. For reduction by Mg powder, giving Mg boride, v. Winkler (B. 23, 772 [1890]). The compound B₂O₃.P₂O₅ (=BPO₄) is obtained by evaporating H₂BO₃Aq mixed with H₂PO₄, heating to redness, and washing with hot water (G. Meyer, B. 22, 2919 [1889]).

Borates (vol. i. p. 529). For some new borates of Li, Rb, and Cs v. Reischle (Zeit f. anorg. Chemie, 4, 166 [1893]). For remarks on the classification of alkali and alkaline earth borates v. Le Chatelier (C. R. 113, 1034 [1891]). For a full account of Mn borate, v. Hartley a. Ramage (C. J. 63, 129 [1893]). The compounds 6FeO.FeBr. 8B.O. and 6ZnO.ZnBr. 8B.O. are described by Rousseau a. Allaire (C. R. 116, 1445; abstract in C. J. 64 [11], 518 [1893]).

Boron, phosphide of. According to Besson (C. R. 113, 78 [1891]), a phosphide of B is obtained by passing PH₂ into BBr₂, and heating the product, BBr₂, PH₂, to 300°, when it gives HBr and B phosphide. The phosphide is said to be a brown powder, insoluble in water, reacting with boiling conc. alkali solutions to give PH₂; gives off P when heated to redness in N (for other reactions v. abstract in C. J. 60, 1418 [1891]).

Boron, selenide of. (?B₂Se₂). Obtained, as a yellowish grey powder, decomposed violently by water, giving H₂Se and some Se, by heating B to full redness in a current of H₂Se (S batier, C. R. 112, 1000 [1891]).

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Berun, sulphides of (vol. i. p. 531). For heat of formation of B₂S₂ v. Sabatier (C. R. 112, 862; abstract in C. J. 60, 981 [1891]). Moissan (C. R. 115, 203 [1892]) gives details of various methods of preparing B₂S₃, and describes the reactions of this compound (v. abstract in C. J. 62, 1393 [1892]). According to M. (l.c. p. 271), boron pentasulphide, B₂S₃, is obtained by heating BI₂ with rather less than one equiv. of S in CS2 solution at 60° for many hours. M. describes the compound, which was not obtained pure, as

a light white crystalline powder, S.G. 1.85.

BROWHYDEIC ACID (vol. i. p. 532). The
V.D. has been determined by Biltz at -15° and -28° (Z. P. C. 10, 354 [1892]); the results agree with the V.D. required by the formula HBr. The M. M.s of HBrAq from 15.47 to 65.59 p.c. HBr have been measured by Perkin

(C. J. 55, 706 [1889])

Preparation .- 1. By the interaction of P. Br. and H_2O , v. Fileti a. Crosa (G. 21, 64 [1891]; cf. Gassman, abstract in C. J. 64 [11], 453 [1893]). 2. By passing H₂S into Br covered with water (Recours, C. R. 110, 784 [1890]).—3. By adding H2SO, Aq S.G. 1.41 to KBr (Feit a. Kubierschky, J. Ph. [5] 24, 159; abstract in C. J. 60, 1320 [1891]; v. also Addyman, C. J. 61, 97 [1892]; and cf. Léger, abstract in C. J. 64 [11], 114 [1893]).

Hydrates. - Pickering (P. M. [5] 36, 11 [1894]) has isolated HBr.3H2O and HBr.4H2O; the former melting at -48° and the latter at -55.8° .

Reactions.—For the decomposition of HBrAq by H,SO,Aq of different concentrations v. Addyman (C. J. 61, 98 [1892]).

BROWINE (vol. i. p. 534). Mol. w. in solu-

tion in water or acetic acid = 159.5 (Br.) (Paterno a. Nasini, B. 21, 2154 [1888]). The V.D.s of Br from 15° to 280°, and at pressures from 15 to 760 mm., have been determined by Perman (Pr. 48, 45 [1891]); no indications of dissociation were obtained. For measurements of the absorption of light by Br v. Camichel (C. R. 117, 807; abstract in C. J. 64 [11], 561 [1898]).

Bromates (vol. i. p. 538). Lithium bromate LiBrO,; prepared by the interaction of Ba(BrO,),Aq and Li,SO,Aq (Potilitzin, J. R. 22, 392; abstract in C. J. 62, 1275 [1892]). Potassium bromate KBrO. The decomposition of this salt by heat has been studied by Cook (C. J. 65, 802 [1894]); decrepitation occurs at c. 180°, with evolution of Br; the salt fuses at a higher temperature, and O is evolved, without any Br, until KBr remains; no oxy- compounds are formed.

Strontium bromate begins to decompose at c. 240°; the principal change is to SrBr₂ and O, but SrO is also formed (Potilitzin, J. R. 22, 454; abstract, with summary of quantitative results,

in C. J. 64 [11], 11 [1893]).

CADMIUM (vol. i. p. 654). Determinations of the at. w. have been made by Partridge (Am. S. [8] 40, 877 [1891]), and by Morse a. Jones (.im. 14, 261 [1892]). P. converted CdC₂O₄ into CdO by heating, CdSO₄ into CdS by heating in H₂S₄ and CdC₄O₄ into CdS by heating in H₂S₅. M. a. J. converted Cd into Cd(NO₂), and this into CdO by heating. The results all lie about 112 (v. abstract in C. J. 60, 399; cf. Clarke, ibid. 390 [1891], and 62, 1397 [1892]). Lorimer a. Smith

(Zeit. f. anorg. Chemie, 1, 364 [1892]) obtained values varying from 112.182 to 111.908 (0 = 16). The molecule of Cd in dilute solution in tin may be diatomic (v. Heycock a. Neville, C. J. 57, 376 [1890]). Cd melts at 320.68°, according to Callendar a. Griffiths (determined by Pt thermometer, C. N. 63, 1 [1891]). For the line-spectrum of Cd v. Ames (P. M. [5] 30, 33 [1890]). According to Williams (Am. 14, 273 [1892]), Cd crystallises in hexagonal and polyhedral forms; axial ratio a:c=1:1.6554. Cd is superficially oxidised at a dull-red heat by nitric oxide (Sabatier a. Senderens, C. R. 114, 1429 [1892]). Montemartini (G. 22 [1] 250; v. abstract in C. J. 62, 1278 [1892]) has analysed the gases produced by the interaction of Cd and nitric acid; with excess of HNO, Aq (27.5 p.c.) at 8°, N, NO, N2O and NH2 are produced.

Cadmium, alloys of (vol. i. p. 655). For an alloy with gold, CdAu, v. Heycock a. Neville (C. J. 61, 914 [1892]; J. ibid. 65, 65 [1894]). For alloys with copper, gold, and silver, pre-pared by immersing Cd in solutions of salts of the other metals, v. Mylius a. Fromm (B. 27,

630 [1894]).

Cadmium, bromide of (vol. i. p. 655). For the double compounds $CdBr_2.xCsBr$, x = 1, 2, and3, v. Wells a. Walden (Zeit. f. anorg. Chemie, 5,

266 [1893]).

Cadmium, chloride of (vol.i.p. 655). Kwasnik (Ar. Ph. 229, 569; abstract in C. J. 62, 566 [1892]) describes CdCl2.2NH3 and CdCl2.NH3. For the double compounds CdCl2.CsCl and CdCl, 2CsCl, v. Wells a. Walden (lc.). Double compounds with HCl, NH,Cl, LiCl, and KCl, of the form CdCl2.MCl. xaq, are described by Chassevant (A. Ch. [6] 30, 5; v. abstract in C. J. 66 [11], 18 [1894]).

Cadmium, fluoride of (vol. i. p. 655). CdF. is obtained by (1) the interaction of HF and Cd at a red heat; (2) the interaction of HF and fused CdCl₂, or CdO at a red heat; it is a colourless, transparent solid; S.G. 6.64; somewhat soluble water, insoluble alcohol of 95° (Poulenc,

C. R. 116, 581 [1893]).

Cadmium, iodide of (vol. i. p. 656). For the double compounds CdI_2 xCsI, when x = 1, 2, and 3, v. Wells a. Walden (l.c.).

Cadmium, oxides of (vol. i. p. 656). CdO is unchanged at c. 1750° (Read, C. J. 65, 318

[1894]).

According to Kouriloff (J. R. 22, 171; abstract in C. J. 62, 1278 [1892]), the compound CdO₂.CdO₂H₂ is obtained by heating CdO₂H₂ with H.O.Aq. Morse a. Jones (Am. 12, 488; abstract in C. J. 58, 1376 [1890]) obtained small crystals, which they say were cadmous oxide Cd₂O, by gently heating CdOH; the hydroxide was obtained by heating CdCl2, CdBr2, or CdI2, with excess of Cd, to the melting-point, in N, or in vacuo, and treating with water, when CdCl₂, CdBr₂, or CdI₂ dissolved, a littl CdC T separated, and CdOH remained as a white amorphous powder.

Cadmium, salts of (vol. i. p. 656). For the electrical conductivities of very dilute solutions of $CdBr_n$ $CdCl_2$, CdI_n , K_2CdI_4 , $Cd(NO_4)_n$ and $CdSO_4$, v. Wershoven (Z. P. C. 5, 481 [1890]).

Cadmium, sulphide of (vol. i. p. 057). Two forms (and perhaps more) of CdS ceem to exist, one lemon-yellow and the other red; they differ CARBON. 903

slightly in S.G., and in crystalline form; the yellow is said to change to the red by friction (v. Buchuer, Chem. Zeit. 11, 1087, 1107; 15, 778; von Klobukoff, J. pr. [2] 89, 412; abstracts in C. J. 54, 224 [1888]; 56, 946 [1889]; 62, 778 [1892]). A solution of colloidal CdS was obtained by Prest (C. C. 1888. 32; abstract in C. J. 54, 653 [1888]) by completely ppg. an ammoniacal solution of CdSO, by H₂S, washing well, suspending the pp. in water and passing in H2S A solution with 4 grms. CdS per litre remained clear for many days; one with 11 grms. coagulated in 24 hours.

CÆSIUM (vol. i. p. 657). According to Beketoff (v. abs ract in C. J. 62, 274 [1892]), Cs is obtained by heating CsOH with c. 1 its weight

Desium, haloid compounds of (vol. i. p. 658). A series of con pounds of the form CsX, where $X_s = Br_s$, I_s , or Br and I, Cl, Br and I, &c., is described by Wells a Penfield (Am. S. [3] 43, 17; abstract in C. J. 62, 778 [1892]). Double compounds of Cs haloids with haloid compounds of Cd, Cu, Mg, Hg, Pb, and Zn are described by Wells (and others) (Am. S. [3] 4, 221; Zeit. f. anorg. Chemie, 3, 195; 5, 266, 273, 300, 304, 306; austracts in C. J. 64 [11], 67, 322 [1893]; 66 [11], 45, 47 [1894]).

Casium, hydroxide of (vol. i. p. 658).

S.G. $_{40}^{\infty}$ 4.0178; [CsOH,Aq] = 15,876;

[CsOHAq, HClAq] = 18,790 (Beketoff, C. C. 1891.

[11] 451).

Casum, oxide of (vol. i. p. 658). According to Beketoff (J. R. 25, 433; abstract in C. J. 66 [11], 234 [1894]), Cs₂O reacts with slightly moist H to give Cs and CsOH.

CALCIUM (vol. i. p. 663). The molecule in dilute solution in tin may be distomic (v. Hey-

cock a. Neville, C. J. 57, 376 [1890]).

Calcium, carbide of. By heating 120 g. CaCO. with 70 g. sugar charcoal in the electric furnace with a current of 350 ampères and 70 volts, Moissan (C. R. 118, 501 [1894]) obtained lustrous, yellow crystals of CaC2; S.G. 2.22 at 18°; insoluble in most solvents; reacts with Cl, Br, and I at 250°-350°; burnt to CaCO, by O at a red heat; interacts violently with water, giving off pure acetylene, and with steam giving CaCO, C, H, and C₂H₂ (abstract in C. J. 66 [1], 313 [1894]).

Calcium, exide of (vol. i. p. 666). CaO is unchanged et c. 1750° (Read, C. J. 65, 318 [1894]). According to Veley (C. J. 63, 821 [1893]; 65, 1 [1894]), dry CaO does not appresent the control of the con ciably combine with dry CO₂, SO₂, or Cl below 300°. Baker (C. J. 65, 611 [1894]) says that dry CaO does not react with dry SO₃, nor with dry NH₄Cl. Moissan (C. R. 115, 1034 [1892]) obtained crystals of CaO by heating ordinary lime in the

electric furnace.

Calcium, exychloride of (vol. i. p. 666). Pre-...ed by soiling CaO with conc. CaCl2Aq; crystallises in thin, lustrous needles; decomposed by water, alcohol, &c.; dissolves in glycerin. Composition is probably 3CaO.CaCl, 15aq (Zahorsky, Zeit. f. anorg. Chemis, 3, 84 [1893]).

Calcium, expiodide of. By heating CaO with Cal.Aq in a sealed tube at 150° for 6 hours, Tassilly (Bl. [3] 9, 629 [1893]) obtained needles

on Cal. SCaC. 16aq.

CARBON (vol. i. p. 684).
Diamond. Moissan (C. R. 116, 218; 118, 820; abstracts in C. J. 64 [11], 275; 66 [11], 189 [1893 and 1894]) obtained grey-black carbonado by dissolving C in iron, heating to 2000°-3000°, and rapidly cooling the exterior by water or by plunging into a bath of melted lead; the liquid interior expands as it cools and produces a very high pressure. Diamond blackens, swells, and changes to graphite in the electric furnace (M., C. R. 117, 428 [1893]). For various reactions of diamond, v. Moissan (C. R. 116, 460; abstract in C. J. 64 [11], 819 [1898]).

For accounts of graphites pre-Graphite. pared in different ways, the compositions of the graphitic oxides obtained, and the H.F.s of these oxides, v. Berthelot a. Petit (C. R. 110, 101, 106; abstract in C. J. 58, 448 [1890]). For the action of conc. HNO, on various graphites, v. Luzi (B.

24, 4085; 26, 1412 [1892 and 1898]).

Amorphous carbon. Mixter (Am. S. [3] 45, 363; abstract in C. J. 64 [11], 571 [1898]) describes the interactions of charcoal and N. S. O. and the halogens. According to Ernst (J. pr. [2] 48, 81; abstract in C. J. 64 [11], 461 [1893]), the composition of the products of combustion of coke (CO and CO2) depends almost wholly on the temperature; the maximum formation of CO₂ is at c. 700°, at 1,000° CO is the only product

Allotropy of carbon. The following heats of combustion, and atomic volumes, are given by Petersen (Z. P. C. 8, 601 [1891]): -

[C,O°]. At. volume. Amorphous . 96,530 to 96,980 6.7 to 8 5.8 Graphite . . . 93,360 Diamond. . . 93,240 to 94,550 8.4

For the reactions of various forms of C with K₂Cr₂O₇ and H₂SO₄ v. Wiesner (M. 13, 871; abstract in C. J. 62, 1273 [1892]); and with HNO, and KClO, v. Schutzenberger (C. R. 111, 774; abstract in C. J. 60, 265 [1891]).

Carbon, tetrabromide of (vol. i. p. 688). Collie (C. J. 65, 262 [1894]) finds that very many compounds of carbon give CBr, when

heated with NaOHAq and Br.

Carbon, bromochorides of (vol. i. p. 688). The three compounds CBrCl, CBr,Cl, CBrCl, are formed by heating CHCl, and Br to 225° in a sealed tube (Besson, C. R. 114, 222

[1892].)

Carbon, tetrafluoride of, CF. Mol. w. 88.87. V.D. 44.6. Prepared by the direct union of C (purified lampblack) and F at the ordinary temperature; by the reaction of F with CCL slightly heated, or with CHCl, at 100°, or with CH, (Moissan, C. R. 110, 296, 951; abstracts in C. J. 58, 557, 944 [1890]); also by heating AgF with CCl, to 220° in a sealed tube (Chabrié, C. R. 110, 279 [1890]). M. recommends to pass vapour of CCl, over AgF at 195°-200° let the vapours pass through a serpentine tube kept at - 23°, and to collect the gas over Hg; the gas is kept in contact with fragments of caoutchouc (to remove COl.), then shaken with absolute alcohol, in which it dissolves, and then expelled by heat, collected and shaken with H₂SO₄ (to remove alcohol vapour); metal vessels must be used. CF, is a colourless gas, liquefying. at -15° and 760 mm. pressure, or at 20° under

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a pressure of 4 atmos. Other fluorides seem to be formed along with CF4; Chabrié (l.c.) says

that AgF and C.Cl., yield C.F..
Carbon, tetra-iodide of (vol. i. p. 689). Moissan (C. R. 118, 19; abstract in C. J. 600, 1420 [1891]) prepares CI, by heating CCl, (free from CHCl,) with excess of BI, to 90° in a sealed tube, washing the crystals with NaHSO, Aq, drying, CI, reacts and slowly subliming in vacuo. with Cl to form CCl, and chlorides of I; reduced to CHI, by H at 140° (for other reactions v. M., l.c.).

Carbon monoxide (vol. i. p. 689). some experiments on the sparking of CO, with the formation of brown solids, v. Berthelot (C. R. 110, 609) and Schutzenberger (C. R. 110, 560; abstracts in C. J. 58, 691 [1890]). Winkler (B. 24, 878 [1891]) finds that CO is reduced to C, at a full red heat, by Mg; carbides of Mg are probably formed (cf. W., B. 23, 2642 [1890]).

Carbon dioxide (vol. i. p. 691). For S.H. of CO₂ v. Kurnskoff (J. R. 22, 498; abstract in C. J. 64 [11], 5 [1893]). CO₂ is reduced by heating to redness with Mg, amorphous C and some CO being formed (Winkler, B. 23, 2642; 24, 878 [1890-91]). Dry CO₂ is said not to combine with dry NH₂ (Hughes a. Soddy, C. N. 69,

Carbon, exychloride of (vol. i. p. 692). COCI becomes viscid, but remains clear, at -75°

(Haase, B. 26, 1052 [1893]).

Carbon disulphide (vol. i. p. 693). The contractions observed when mixtures of CS2 and air are exploded show that the reaction is more complicated than is shown by the equation $CS_2 + 3O_2 = CO_2 + 2SO_2$; the volume of CO_2 found is, practically, in accordance with the equation, but the volume of SO_2 is smaller; some of the N of the air is oxidised (v. Pedler, C. J. 57, 625 [1890]). By heating CS, with excess of aqua regia, Schlagdenhauffen a. Bloch (J. Ph. [5] 28, 241 [1893]) obtained SO(OCCl₂)Cl. Decomposition of CS, heated in dry N, begins at 216°, and explosion with O at a higher tem-

perature (Baker, C. J. 65, 611 [1894]).
Tricarbon disulphide C₂S₂. B. von Lengyel (B. 26, 2960 [1893]) obtained this compound by keeping the vapours of CS, in an electric arc between C poles for some hours, filtering off the black substance formed, allowing the red filtrate to stand over Cu for 6 to 8 days, and evaporating in a current of dry air. C₂S₂ is a deep-red liquid, S.G. 1.27389; the vapour, even in minute quantities, causes violent catarrh; can be distilled, with partial change to a black solid, at 60°-70° and reduced pressure; insoluble water, but dissolved by EtoH, Et₂O, CS₂, CHOl₂, or O₂H₂. A conc. solution in CS₂ deposits a black solid; the same solid is formed by gently heating the liquid, if heating to 100°-120° is rapid the change is explosive. The black solid is probably a polymeride. A solution of C,S, in CS, reacts with Br to form C,S,Br,. (For other reactions, v. abstract in C. J. 66 [11], 91 [1894].)

Carbon, thiobromide of, C.S.Br.; formed by the action of Br on a solution of C.S. in CS. (v.

Carbon, thiochloride of (vol. i. p. 695). polymeride of thio-carbonyl chloride has the cryoscopic determinations of Carrara (G. 28 [11] 12; abstract in C. J. 66 [11], 15 [1894]).

CERIUM (vol. i. p. 723). For details regarding the preparation of Ce compounds free from compounds of Di and La, v. Schottländer (B. 25, 378 [1892]); also Bricout (C. R. 118, 145 [1894]).

Cerium, hydride of. According to Winkler (B. 24, 873 [1891]), a compound of Ce and H, probably CeH2, is formed by reducing CeO2 by

Mg in presence of H.

Cerium dioxide (vol. i. p. 725). 'CeO, is unchanged at c. 1750° (Read, C. J. 65. 813

[1894])

CHLORHYDRIC ACID (vol. i. p. 5). Biltz (Z. P. C. 10, 854 [1892]) finds that the V.D. at -77° agrees with the formula HCl. Perkin (C. J. 55, 703, 705 [1889] has determined M. M of HClAq and HCl in iso-amyl oxide; the values obtained for HOl in Aq are c. double those for HOl in iso-amyl oxide. For measurements of the E.C. of HOl in different solvents, v. Kablukoff (Z. P. C. 4, 429 [1889]). For revised tables giving S.G. of HClAq, v. Lunge a. Marchlewski (Zeit. f. anorg. Chemie, 1891, 183).

For the relations between the number of molecules of HCl required to ppt. one and of different chlorides, and the reduction of the freezing-points of solutions of the chlorides and of HCl, v. Engel (C. R. 117, 845; abstract in

C. J. 66, 40 [1894]).

Bailey a. Fowler (C. J. 53, 755 [1888]) find that when HCl stands over P₂O₅ the gas is absorbed, with production of POCl₂ and HPO₃; and that when HCl and O stand over Hg an oxychloride, probably Hg2OCl2.H2O, is formed. Baker (C. J. 65, 611 [1894]) states that dry HCl does not combine with dry NH, when the gases are mixed.

CHLORINE (vol. ii. p. 10). For the S.G. of liquid chlorine from -80° to 77°, and the vapour pressures from -88° to 1.6°, v. Knietsch (A. 259, 100 [1890]). For the preparation of Cl for use in the laboratory, v. Klason (B. 23, 330; abstract in C. J. 58, 445 [1890]). Shenstone (C. J. 61, 445 [1892]) finds that Cl obtained by heating PtCl, always contains some HCl and O, with water, and perhaps a trace of N. Cl expands when brought into sunlight. Richardson (P. M. [5] 82, 277 [1891]) has constructed an instrument wherein the expansion is used to measure the intensity of light. Baker (C. J. 65, 611 [1894]) found that no explorion occurred when a mixture of equal vols. of dry Cl and dry H was exposed to bright sunlight; and that after exposure to bright sunshine for two days, and to diffused light for two days, more than a quarter of the mixture remained unchanged.

Pedler (C. J. 57, 613 [1890]) has examined the interaction of Cl and water in tropical sunlight; he finds that there is very little action unless there is an extremely large excess of water. With very much water and little Cl, exposed to the full light of the sun in the hot season in India, the main reaction is $2H_2O + 2Ol_2 + Aq = 4HClAq + O_2$; with diffused sunlight HClAq and HClOAq are formed at first, and finally HClAq, HClO, Aq, and O. Harker (Z.P.C.9, 678 [1892]), from an extended investigation of the explosion of mixtures of Cl, H, and molecular formula (OSCla), according to the O, concluded that when there is insufficient H &

combine with all the Cl and all the O, a division of the H between the Cl and O occurs; and that the product of the numbers of molecules of HCl and O, divided by the product of the numbers of molecules of H₂O and Cl, is a constant, or, in other words, that the law of Guldberg a. Waage holds good. According to Gautier a. Charpy (C. R. 113, 597 [1891]), there is no reaction between Mg or Zn and liquid Cl at the ordinary temperature, very little reaction with Fe or Ag, and rather more with Cu.

Chlorates (vol. i. p. 18). Spring a. Prost (Bl. [3] 1, 840 [1898]) have determined the quantities of Cl obtained by decomposing by heat chlorates of Al, Ba, Cu, Pb, Hg, K, Ag, Na, and Zn; the amount of Cl varied from ·02 p.c. from KClO, to 14·45 p.c. from Zn(ClO₃)₂.

Talcium chlorate. Dry Ca(ClO₃), melts between 800°-250°, having previously given off Dry Ca(ClO_s), melts from 3 to 5 p.c. of its weight of O; the rate of evolution of O increases, at 330°, until c. 60 p.c. of the total O is given off, after which the rate decreases. For the quantities of O evolved, of Ca (ClO₃)₂ decomposed, and of CaCl₂ and Ca(ClO₄)₂ formed, v. abstract in 7. J. 62, 1275

[1892].

Poi ssium chlorate. McLeod (C. J. 55, 184 [1889]) concluded from his experiments that when KClO, is heated with MnO, some KMnO, is formed and again decomposed (v. Manganese PEROXIDE, vol. iii. p. 183), and that Cl is always given off in this reaction. In 1893 (B. 26, 1790) Brunck stated that Cl is not given off when KClO₃ and MnO₂ are heated together, but that ozone is produced. McLeod (C. J. 65, 202 [1894]) has proved that Cl is produced in the reaction. Fowler a. Grant (C. J. 57, 272 [1890]) have studied the influence of various oxides on the decomposition of KClO, (references are given to older memoirs); they support McLeod's view of the reaction with MnO, and think that the oxides of Fe, Co, Cu, and Ni react similarly to MnO2; acidic oxides, such as V2O4, U3O4, and WO, cause evolution of O accompanied by much Cl, a vanadate, uranate, or tungstate being also formed; powdered glass or sand also assists the reaction, but only when very finely divided (v. summary of conclusions, C. J. 57, 281-2). According to Thorpe a. Perry (C. J. 61, 925 [1892]), the main reaction which occurs when an intimate mixture of KClO, and iodine is heated is $2KClO_3 + I_2 = 2KIO_3 + Cl_2$, and some of the Cl reacts with unchanged I to form ICl and ICl₂. Basset (C. J. 57, 760 [1890]) examined the reactions between KOlO, water, and iodine; he concluded that the main reaction, at 80° , in the presence of a very small quantity of HNO_2Aq , is as shown by the equation $6I_2 + 10KClO_2 + 6H_2O + Aq =$ 6KHI₂O₄Aq + 4KClAq + 6HClAq; in evaporating the solution to dryness Cl was given off, and the results of several experiments led to the representation of the change produced by evaporation to dryness as $12\text{KHI}_2O_0 + 8\text{KOl} + 12\text{HOl} = 11\text{KHI}_2O_0 + 9\text{KCl} + 6\text{H}_2\text{O} + 1\text{Cl} + 1\text{Cl}.\text{HCl} + 4\text{Cl}_2.$ For determinations of the quantities of oxidising gases (Cl and oxides of Cl) removed by passing air through KClO₂Aq mixed with HClAq v. Pendlebury s. McKillop (C. S. Proc. 1898-4. [No. 128] 211).

Strontium chlorate. For experiments on the

decomposition of this salt by heat, giving SrCl, Sr(ClO₄), and O, v. Potilitzin (J. R. 21, 451; abstract in C. J. 58, 696 [1890])

CHROMIUM (vol. ii. p. 152). S.H. of pure Cr (0° to 98°) = 12162 (Jäger a. Krüss, B. 22, 2028 [1889]). The at. w. of Cr was re-determined by Rawson in 1889 (C. J. 55, 218) by strongly heating (NH,), Cr,O,, measuring the N given off, and weighing the Cr.O. that remained; also by reducing (NH₄)₂Or₂O,Aq to OrCl₂Aq by HClAq and alcohol, ppg. by NH₂Aq, and weighing Cr₂O₈ produced; the mean value thus obtained was 52.06, or 51.98 reduced to 0° and a vacuum (0 = 15.96). Menieke in 1891 (A. 261, 339; abstract in C. J. 60, 882 (1891)) re-determined the at. w. of Cr by estimating (1) Ag and Cr, and also O, in Ag₂CrO₄ and in Ag₂CrO₄.NH₄, (2) O in $K_2Cr_2O_7$; (3) O and Cr in $(NH_4)_2Cr_2O_7$; the extreme values, from thirty-six experiments, were 52:12 and 51.83; twenty-eight of the experiments gave values between 51.9 and 52. According to Prinz (C. R. 116, 392 [1893]), Cr crystallises in minute cubes with pyramidal faces. For the absorption spectra of salts of Cr, v. Lapraik (J. pr. [2] 47, 305 [1893]).

Preparation.—Glatzel (B. 23, 8127 [1890]) says that almost perfectly pure Cr can be ob-tained by reducing CrCl₂.KCl by Mg powder (for details v. abstract in C. J. 60, 152 [1891]). Placet (C. R. 115, 945 [1892]) prepares Cr by electrolysing a solution of Cr alum, containing an alkali sulphate and H₂SO₄. Moissan (C. R. 116, 349 [1893]) says that Cr can be obtained by reducing a mixture of Cr2O2 and C in the electric furnace, using a current of 50 volts and 100 ampères, and then breaking up the product, which contains from 8.6 to 12.9 p.c. C, and heating it, mixed with more Cr.O., again in the electric

furnace in a carbon crucible.

Chromic acid (vol. ii. p. 154). Miss Field (C. J. 61, 405 [1892]) has found that only CrO, is obtained by working according to Moissan's directions for preparing H2CrO. Determinations of the increase in the b.p. of water caused by solution of CrO, led to the conclusion that an aqueous solution of CrO, contains some H, CrO, and probably also H2Cr2O,.

Chromic bromide CrBr, (vol. ii. p. 161). Recours (C. R. 110, 1029, 1193) obtained green CrBr, 6H,O by boiling saturated CrO,Aq with a large excess of HBrAq and concentrating; deliquescent crystals, soluble alcohol, insoluble ether. Unchanged when kept solid, but solution soon becomes blue and then violet, with rise of temperature. By boiling a green solution, then saturating with HBr while cold, filtering, and drying the pp. thus formed on porous plates, crystals of the blue variety, CrBr. 6H.O. were obtained; very soluble water, insol. alcohol. Heat of solution of blue crystals = 28,700; and of green = 1,860. The blue solid is changed to the green at 100°. (For more details v. abstract in C. J. 58, 1063 [1890].)

Chromic chloride CrCl, (vol. ii. p. 161). Marchetti (G. 22 [11], 875 [1892]) finds that the molecular lowering of the freezing-point of water by the violet form of CrCl, is considerably greater

than that caused by the green form.

Chromic fluoride CrF, (vol. ii. p. 162). Fabris (G. 20, 582 [1890]) obtained CrF, 9H, O as a green crystalline pp., by adding excess of NH,FAq gradually to a cold solution of violet Cr.(80,),. water; insoluble Slightly sol. NH, FAq or alcohol; heating in air leaves Cr.O. Poulenc (C. R. 116, 253 [1893]) obtained CrF, in green crystals by heating CrCl₂, Cr₂O₃, or CrF₂.xH₂O in HF.

Chromous fluoride. According to Poulence (l.c.), OrF, is formed by heating Or to redness in HF, or by the interaction of CrCl2 and HF at the ordinary temperature. CrF2, after fusion, is described as a transparent, green, crystalline mass; S.G. 4:11; slightly sol. water, insol. alcohol. Heated in air it gives Cr.O_s. (For more details v. abstract in C. J. 64 [11], 281 [1893].)

Chromic oxide Cr_2O_3 (vol. ii. p. 164). Moissan (C. R. 115, 1034 [1892]) found that Cr_2O_3 melted to a black mass, dotted with black crystals, in an electric furnace, using a current of 55 volts and

80 ampères.

Chromium trioxide CrOs (vol. ii. p. 164). For action of water v. Chromic acid, p. 905. Reduced to Cr_2O_3 by NO at the ordinary temperature (Senderens a. Sabatier, C. R. 114, 1476 [1892]). For the reaction of CrO₃Aq with O and BaO.H.Aq—said to give a pp. approximately BaOrO.—v. Péchard, C. R. 113, 89 (abstract in C. J. 60, 1431 [1891]).

Chromium salts (vol. ii. p. 167). According to Recoura (C. R. 112, 1439; abstract in C. J. 60, 1430 [1891]), the green solutions obtained by heating solutions of the normal violet chromic salts contain free acid and a soluble basic salt; in the case of the sulphate this soluble basic salt is said to be 2Cr₂O₂.5SO₂. For preparation of green soluble normal sulphates v. Recoura, C. R. 118, 857 (abstract in C. J. 62, 411 [1892]). CHROM-AMMONIUM COMPOUNDS (vol. ii.

p. 158). Petersen (Z. P. C. 10, 580 [1892]) finds, by cryoscopic determinations, that the formulæ usually assigned to the chief salts of this class are probably molecular. For a discussion of the constitution of these salts v. Jörgensen (J. pr.

[2] 42, 206; 45, 274).

CHROMATES; and derivatives (vol. ii. p.

154).

Ammonium chromates. According to Jäger a. Krüss (B. 22, 2028 [1889]), normal ammonium chromate has not hitherto been obtained pure; they prepare the pure salt by adding excess of NH, Aq S.G. 9 to CrO, Aq free from H, SO, Aq, warming until the solid that separates dissolves, and placing in a freezing mixture. (NH4), CrO4 crystallises in long, monosymmetric, golden needles, a:b:c=1.9603:1:2.4452; S.G. 1.886 at 11°

Ammonium trichromate (NH4)2Cr2O10 is prepared by J. a. K. (l.c.) by dissolving (NH₄)₂Cr₂O₇ in hot conc. OrO, Aq, concentrating, and drying at 190° the red crystals that separate on cooling; S.G. 2.342 at 18°; explodes at 190°, giving off

NO, and leaving Cr.O,.

Ammonium tetrachromate (NH4)2Cr4O13 is formed by dissolving the trichromate in warm HNO, Aq S.G. 1.09, and letting cool slowly & S.G. 2.843 at 10°; melts at 170°, and suddenly decomposes at 175°, giving off NO and leaving Cr.O., J. a. K. (l.o.) failed to obtain hexachromate; they say that no nitro-chromates are obtained by treating ammonium chromates with (NH₄)₂Or₂O₂ with HgCl, are described by J. a. K.

Ceric dichromate CeO2.2CrO2.2H2O. pared by passing a current of 2.5 to 8 volts through a slightly acid solution formed by dissolving Ce₂(CO₂)₃ in CrO₂Aq; insoluble water, boiling water forms yellow Ce(C O₁)₂ and then CeO₂.xH₂O (Bricout, C. R. 118, 145 [1894]).

Lead chromates. For double compounds of PbCrO₄ with Li₂CrO₄, K₂CrO₄, and Na₂CrO₄, v. Lachaud a. Lepierre, C. R. 110, 1035 (ubstract

in C. J. 58, 1065 [1890]).

Mercury chromates. For basic mercuric chromates and a double compound of HgCrO, with NH, HgCl v. Jäger a. Kräss, B. 22, 2028

(abstract in C. J. 56, 1120 [1889]).

Potassium chromates. Jäger a. Krüss (l.c.) prepare potassium trichromate K₂Cr₃O₁₀ by treating K₂Cr₂O, with HNO₃Aq S.G. 119, pouring off from KNO, that separates, and crystallising; S.G. 2.648 at 11°. K₂C₇O₁₀ is quickly decomposed by water to CrO, Aq and K, Cr, O, Aq; melts at 250°. Potassium tetrachromate K, Ci, O, is prepared by J. a. K. (l.c.) by dissolving K.Cr.O., in hot cone. HNO, Aq S.G. 1.41; S.G. 2.649 at 11°; not deliquescent; decomposed by water; crystallises unchanged from HNO, Aq S.J. 1.41. Schmidt (B. 25, 2917 [1893]) says that no nitrochromate is obtained by the action of HNO, Aq on K2Cr2O7; the substances obtained by Darmstädter (v. vol. ii. p. 158) were mixtures of KNO, and K2Cr4O1

Silver chromates. Pure silver chromate Ag₂CrO₄ is prepared by J. a. K. (l.c.) by repeatedly boiling Ag₂Cr₂O₇ with water as long as anything dissolves; it is a deep-green solid,

insol. water.

Sodium perchromate. Häussermann (J. pr. [2] 48,70; abstract in C. J. 64 [11], 471) describes a salt Na₆Cr₂O₁₅. 28aq, prepared by adding Na₂O₂ to a thin paste of CrO,H, and water, kept at 10°-20°

Thallium chromates. For preparation of Tl, CrO, and the double compound Tl, CrO, K, CrO, v. Lachaud a. Lepierre, Bl. [3] 6, 232 (abstract in C. J. 62, 567 [1892]).

Uranium chromates. For double uranyl chromates v. Formánek, A. 257, 102 (abstract in

C. J. 58, 852 [1890]

IODOCHROMATES (vol. ii. p. 157). A series of salts of the forms IO2.O.CrO2.OM1 and (IO2.O.CrO2.O) MII has been obtained by Berg (C. R. 104, 1514 [1887]; 111, 42 [1890]); M^{r} - NH₄, Li, K, Ag, and Na; M^{tt} = Co, Cu, Mg, and Ni. Most of the alkali salts were formed by dissolving the alkali iodate in excess of CrO,Aq and crystallising; the other salts were usually prepared by dissolving the oxide or carbonate of the metal in excess of CrO₃Aq and adding HIO₃Aq. (For details v. abstracts in C. J. 52, 776 [1887]; 58, 1878 [1890].)

PHOSPHOCHROMATES. By adding conc. H₂PO,Aq + CrO₂Aq to K,CO₂Aq, Blondel (O. A. 118, 194; abstract in O. J. 66 [11], 198 [1894])

obtained compounds of the form

mK2O.P2O3.nOrO3. xaq, m being 2 and 3, n 4 and

8, and x being 0 or 1.

SULPHOCHROMATES. In 1882 Cross a. Higgin described a compound of Cr.O., SO., and HNO.5aq. (For more details v. abstract in C. J. H.O (C. J. 41, 113), and also compounds of Cr.O., 56, 1117 [1889].) Various double compounds of SO., K.O. and H.O. Becoura (C. R. 114, 477

[1892]; 116, 1367; 117, 37, 101 [1898]) has made a more complete examination of these and By evaporating, at 100°, allied compounds. solutions containing violet Cr₂(SO₄), and H₂SO₄, in the molecular ratio 1:1, 1:2, and 1:3, R. obtained chromosulphuric acid H₂Cr₂(SO₄), chromodisulp uric acid H, Cr2(SO4)5, and chromotrisulphuric acid H₆Cr₂(SO₄)₆; by using various sulphates in place of H.SO, he obtained chromosulphates, chromodisulphates, and chromotrisulphales. The acids are green powders, easily sol. water; they gradually decompose in solution; dilute BaCl,Aq ppts. Ba chromosulphates, but conc. BaCl, Aq gives BaSO. R. obtained chromopyrosul; huric acid Cr. (OH), (S2O, H), by evaporating a solution of Cr₂(SO₄), and H₂SO₄, in the molecular ratio 1:5, at 100°, and heating the dar':-green syrupy liquid so obtained to 110°-115° for some days. The acid forms brittle, transparent, pale-green laminæ. Salts of the form $\operatorname{Cr}_2(S_2O_7)_2(\operatorname{OM})_2$ are obtained by adding alkalis to the acid in solution; these salts R. calls pyrosulphochromites. By adding conc. HClAq or H₂SO₄ to a solution of chromopyrosulphuric acid, B. obtained a compound Cr₂(OH)₂(S₂O₇)₂, pyrosulphochromic hydroxide, isomen is with chromosulphuric acid Cr2(804)4H2. (For more details v. abstracts in C. J. 62, 783 [1892]; 64 [11], 470, 528 [1893].]

COBALT (vol. ii. p. 217). Winkler has redetermined at. w. of Co, (1) by dissolving electrolytically deposited Co in HClAq, evaporating to dryness, and determining Cl in the CoCl₂ obtained (Zeit. f. anorg. Chemie, 4, 10 [1893]);
(2) by finding the quantity of Ag deposited from solution of Ag₂SO₄ by electrolytically deposited Co (l.c. p. 462). The values obtained were all

about 59.6 (Ag = 107.66).

Alleged decomposition of cobalt. Krüss a. Schmidt (B. 22, 11, 2026 [1889]) supposed they had separated Ni into two constituents (v. vol. iii. p. 500); and they thought that Co also was probably not a homogeneous substance. Remmler, in 1893, obtained a number of specimens of Co₂O₃, by ppg. CoCl₂Aq by KOHAq and Br water, treating the pp. with NH₃Aq, which gradually dissolves it, decanting off the ammoniacal solution at intervals during 9 months, evaporating each and heating in CO2; the different specimens of Co.O. were heated in H, and the quantity of Co obtained in each case was determined. The values thus obtained for the at. w. of Co varied from 59.53 to 58.3. R. concluded that Co purified by the ordinary methods is not a homogeneous substance. Winkler's determinations of the at. w. (v. supra), however, gave very constant values.

Reactions.—Heated in NO to 150°, CoO is formed (Sabatier a. Senderens, C. R. 114, 1429 [1892]); finely divided Co burns in NO2 at the ordinary temperature, giving Co₂O₅; but if the NO₂ is largely diluted with N a black compound is formed, Co₂NO₂, called by S. a. S. (C. R. 115, 286 [1892]) mitro-cobalt. This compound reacts rapidly with H₂O, giving off NO, and forming a solution containing Co(NO₂), with a very little Co(NO₂), Co remaining insoluble, and sometimes riso a basic nitrite; when heated in N, orides of N are given off and then rapid decomposition occurs with formation of Co and oxides of Co; when mixed with combustible substance, Co.NO.

explodes (S. s. S., Bl. [3], 9, 669 [1893]). Montemartini (G. 22 [1], 250) finds that the gases produced by the interaction of Co and excess of HNO₃Aq (27.5 p.o.) are NH₃, N₂O, and N (v. abstract in C. J. 62, 1278 [1892]).

Cobalt, chloride of (vol. ii. p. 219). For the solubility of CoCl, in water, and the changes of colour produced by diluting the solution, with a discussion of the hydrations and dehydrations that accompany these changes, v. Etard, C. R. 113, 699 (abstract in C. J. 62, 278 [1892]); Engel, Bl. [3] 6, 239 (abstract in C. J. 62, 569 [1892]); and Potilitzin, Bl. [3] 6, 264 (abstract in C. J., Lc. p. 571). The double (abstract in C. J., l.c. p. 571). The double compounds CoCl₂.HCl. 3aq, CoCl₂.LiCl. 3aq, and CoCl. NH, Cl. 6aq are described by Chassevant (A. Ch. [6] 30, 5 [1893]).

Cobalt, fluoride of (vol. ii. p. 220). Poulence (C. R. 114, 1426; v. abstract in C. J. 62, 1159 [1892]) prepares CoF, by heating CoCl, with excess of NH F for 15 minutes, and repeatedly washing the double fluoride CoF, 2NH, F thus obtained with boiling alcohol. By heating CoF, to 1200°-1800° in HF, the salt is obtained in small, rose-red prisms; S.G. 4.43. The double compound CoF2.2KF is formed by heating CoCl2

with KHF₂ (P., l.c. p. 746; abstract, l.c. p. 781).
Cobalt, iodide of (vol. ii. p. 220). For solu bility in water of CoI2, and the formation of various hydrates attending solution, v. Etard, C. R. 118, 699 (abstract in C. J. 62, 278 [1892]).

Cobaltons oxide CoO (vol. ii. p. 220). This oxide melts in the electric furnace, and forms rose-coloured crystals (Moissan, C. R. 115, 1034

Cobaltous hydroxide CoO2H2 (vol. ii. p. 220). A. de Schulten (C. R. 109, 266 [1889]) obtained this compound in microscopic, brownish-red, four-sided prisms, S.G. 3.597 at 15°, by heating 10 g. CoCl₂. 6aq in 60 c.c. water with 250 g. KOH, in a flask filled with coal-gas, letting stand for 24 hours, and washing away the lighter amorphous products of the reaction.

Cobalto-cobaltic oxides (vol. ii. p. 221). Regarding Co₂O₃, and oxides intermediate between this and Co₂O₃, v. Schröder, C. C. 1890. [1] 981 (abstract in C. J. 58, 1213 [1890]). Co.O. heated to c. 1750° loses all its O, and leaves Co (Read,

C. J. 65, 313 [1894]).

Cobalt, dioxide of. By the reaction of IAq and NaOHAq on CoSO,Aq, filtering, and acidifying with acetic acid, Vortmann (B. 24, 2744; abstract in C. J. 60, 1429 [1891]) obtained a greenish-black pp. nearly agreeing with the formula CoO₂. For compounds of CoO₂ with BaO v. Rousseau, C. R. 109, 64 (abstract in C. J. 56, 1115 [1889]).

Cobalt, salts of (vol. ii. p. 221). Marshall C. J. 59, 760 [1891]) has prepared the salts $Co_2(SO_4)_2$. 18aq and $Co_2(C_2O_4)_3$. 3(NH₄)₂ C_2O_4 . 6aq by electrolysing a solution of CoSO, in H.SO, Aq,

and of CoO₂O₄Aq containing (NH₄)₂C₂O₄.

COBALTAMMINES (vol. ii. p. 222). crypscopic determinations, Petersen (Z. P. C. 10, 580 [1892]) concludes that the formulæ generally given to the chief cobaltammines are molecular. For a discussion of the constitution of these compounds, v. Jörgensen (J. pr. [2] 41, 429, 440 [1890]; 42, 206; 45, 274 [1891-92]). For various octammines, v. Vortmann 3. Blasberg (B. 22, 2648), and V. a. Magdeburg (B. 22, 2680; abstracts in C. J. 58, ·14 [1890]). Morgulis (B. 22, 2644; abstract in C. J. 58, 18 [1890]), describe several mercuricobaltammines. A number of crocso-, luteo-, and xantho- salts, and also several flavo- salts isomeric with the croceosalts, are described by Jörgensen in Zeit. anorg. Chemie, 5, 147 (abstract in C. J. 65 [11],

50 [1894]

COPPER (vol. ii. p. 251). An elaborate investigation of the at. w. of Cu has been made by Richards, P. Am. A. 26, 240 [1891] (v. C. N. 65, 236, &c.; abstract in C. J. 64 [11], 12 [1893]). CuSO, 5H₂O was analysed and synthesised, and CuO was analysed; the final value obtained by R. is 63.604 (O = 16), or 63.44(0 = 15.96). The molecule of Cu in dilute solution in tin is probably monatomic (Heycock a. Neville, C. J. 57, 376 [1890]). For the absorption spectra of solutions of CuCl₂, CuSO₄, and Cu(NO₂)₂, v. Evan, P. M. [5] 38, 817 (abstract in C. J. 64 [11], 149 [1893]).

Preparation. - For the preparation by electrolysis of pure copper v. Richards (P. Am. A. 25,

199, 206 [1890]).

Reactions.-Heated in the electric furnace, using a current of 70 volts and 350 ampères, Cu rapilly volatilises, and the vapour forms CuO in the air (Moissan, C. R. 116, 1429 [1893]). According to Neumann (M. 13, 40 [1893]), the H occluded in Cu, prepared by reducing CuO by H, cannot be wholly expelled by heating in a stream of CO₂; some of the CO₂ is also absorbed by the Cu. N. also says that Cu reduced by vapour of MeOH or EtOH, or by coal gas, retains very small traces of C and H after being heated to 220°. Montemartini (G. 22, 884, 397, 426 [1892]) says that the reaction between Ou and HNO, Aq of less than 30 p.c. produces Ou(NO₃)₂ and HNO₂, the HNO₂ then decomposing to HNO₃, NO, and H₂O; with acid more conc. than 30 p.c. HNO₃ some NO₂ is produced with a little N₂O₂, and with 70 p.c. acid only NO is given off; no appreciable quantity of NH, is produced with acid from 3 to 27.5 p.c. HNO. According to Freer a. Higley (Am. 15, 71 [1893]), the only gaseous products of the reaction of Cu with excess of HNO.Aq S.G. 1.4 (c. 65 p.c. HNO_s) are NO₂ and N₂O_s, c. 90 p.c. NO₂ and 10 p.c. N₂O₃. Cu is superficially oxidised in NO at a dull red heat (Sabatier a. Senderens, C. R. 114, 1429 [1892]). Reduced Cu absorbs NO, at the ordinary temperature, forming a brownish powder Cu₂NO₂ (S. a. S., C. R. 115, 236 [1892]). For the reactions of this nitrocopper v. S. a. S., C. R. 116, 756 (abstract in

C. J. 64 [11], 874 [1893]).
Copper, alloys of (vol. i. p. 253). For a description of alloys of Cu with Cd, Sn, and Zn, prepared by immersing plates of Cd, Sn, or Zn, in solutions of salts of Cu, v. Mylius a. Fromm, B. 27,630 (abstract in C. J. 66 [11], 235

1894]).

Cupric bromide (vol. ii. p. 254). According to Sabatier (C. R. 118, 980 [1894]), the green crystals that separate from CuBr,Aq are CuBr. 4aq. A solution of CuBr, in conc. HBrAq has an intense purple colour which becomes less marked on dilution, but even with **0015 mgm. Ou in solution the colour can be detected (c., l.c.) The double salts CuBr. CsBr and CuBr. 20sBr are described by Wells a. Walden (Zeit. f. anorg. Chemie, 5, 304; abstract in C. J. 66 [11], 47 [1894]).

Cupric chloride (vol. ii. p. 255). Tzuchanoff (J. R. 25, 151; abstract in C. J. 66 [11], 47 [1894]) finds that the E.C. of dilute blue-coloured CuCl, Aq increases with increase of concentration to a maximum, after which it decreases while the colour changes to yellowish-brown. For heats of solution and dilution of CuOl₂. 2aq, v. Reicher a. Deventer (Z. P. C. 5, 559 [1890]).

Double salts.—CuOl, HCl. 3aq,
CuCl, LiCl. xaq, CuCl, NH, Cl, CuCl, 2NH, Cl. 2aq,
CuCl, KCl, and CuCl, 2KCl. 2aq (Chassevant,
A. Ch. [6] 80, 5 [1893]); CuCl, CsCl. $CuCl_2.2CsCl.$ xaq(x=0) and 2), and $2CuCl_2.3CsCl$ (Wells a. Dupree, Zeit. f. anorg. Chemie, 5, 300

[1893]).

Cuprous chloride (vol. ii. p. 255). Wells (l.c., p. 806) describes the double salts

2CuCl.CsCl, 2CuCl.3CsCl, and CuCl.3CsCl. aq. Cupric fluoride (vol. ii. p. 256). Poulenc (C. R. 116, 1446; abstract in C. J. 64 [11], 525 [1893]) prepares QuF₂, as a white amorphous powder, by heating NH₄F with CuF₂. 2aq; by heating in HF at 500° (not above) the CuF, becomes crystalline. Crystalline CuF, is also obtained by heating CuO, or CuF₂. 2aq, 'n HF at 400° (P., l.c.). Heated to 300° in air CuO is formed; heating in steam gives CuO and HF; H2S forms CuS and HF; HCl forms CuCl. CuF. absorbs water from the air, and becomes blue. H. von Helmholt (Zeit. f. anorg. Chemie, 3, 115 [1892]) describes the double salts CuF₂.KF, CuF₂.RbF, CuF₂.AmF. xaq, and CuF₂.2AmF. 2aq.

Cuprous fluoride (vol. ii. p. 256). According to Mauro (Real. Acad. Lincei, 1892 [1]. 194; abstract in C. J. 64 [11], 124 [1893]), Cu₂F₂ does not exist, and the substance obtained by Berzelius was almost certainly merely impure copper. Poulenc, however (C. R. 116, 1446; abstract in C. J. 64 [11], 525 [1893]), says that Cu.F. is formed by heating Cu₂Cl₂ to dull redness in HF (the action is not complete until 1100°-1200°); also by heating CuF2 in HF at c. 600°, raising the temperature to 1100°-1200° towards the end of the reaction. P. describes Cu₂F, that has been fused as a ruby-red, transparent solid, with a crystalline fracture; exposed to moist air it changes to CuF₂ xaq; it is easily reduced by H at a red heat.

Cuprous iodide (vol. ii. p. 256). Brun [C. R. 114, 667; abstract in C. J. 62, 1157 [1893]) describes double compounds with (NH₄)₂S₂O₂ and NH₄I, and also with (NH₄)₂S₂O₂

and Cu₂S₂O₃.
Cuprous exide (vol. ii. p. 258). Russell (C. N. 68, 808 [1893]) prepares Cu₂O by completely reducing CuSO₂Aq mixed with excess of NaCl by SO₂, heating till excess of SO₂ is removed, and then ppg. by Na₂O₃, added solid to the hot solution. With HFAq, CuF₂Aq and Cu are formed (Poulenc, C. R. 116, 1446 [1898]).

Cupric oxide (vol. ii. p. 258). Moissat. (C. R. 115, 1034 [1892]) says that CuO is entirely decomposed in the electric furnace at c. 2500°, yielding Cu and a crystalline compound of CuO and CaO (the reaction was effected in a block of CaO). By heating CuO to whiteness, in a Fletcher injector-furnace, in an oxidising atmosphere, Bailey a. Hopkins (C. J. 57, 269 [1890]) obtained a very hard, yellowish-red GOLD. 909

solid agreeing with the formula Cu₂O. Richards (P. Am. A. 20, 281 [1891]) found that CuO occludes gas, chiefly N; when prepared by heating Cu(NO₃)₂ the oxide retained c. 08 p.c. of its weight of N, which could not be removed by heating. For experiments on the dehydration of CuO.xH₂O v. Spring a. Lucion (Zeit. f. anorg. Chemie, 2, 195; abstract in C. J. 64 [11], 210 [1893]). For colloidal, jelly-like CuO.xH₂O v. J. van Bemmelen (Zeit. f. anorg. Chemie, 5, 466; abstract in C. J. 66 [11], 191 [1894]).

Copper, oxybromides of (vol. ii. p. 260). Dupont a. Jansen (Bl. [8] 9, 193; abstract in

Copper, oxybromides of (vol. ii. p. 260). Dupont a. Jansen (Bl. [3] 9, 193; abstract in C. J. 64 [11], 463 [1893]) obtained the salt 3CuO.CuBr₂.3an by heating Cu₂Br₂Aq with CuO in a sealed tube at 200°, or, better, by heating neutral dilute Cu₂Br₂Aq at 225°; the compound crystallised in green rhombohedral plates, S.G.

4.39.

Copper, oxychlorides of (vol. ii. p. 260), v. Rousseau, C. R. 110, 1261 (abstract in C. J. 58, 1058 [1890]).

Copper, exysulphides of (vol. ii. p. 260), v. Cliche, Ar. Ph. 228, 374 (abstract in C. J. 58,

1211 [1890]).

Copper, phosphides of (vol. ii. p. 260). For desc intions of Cu₂P₂ and Cu₂P, prepared by heating Cu in vapour of P, v. Granger, C. R. 113, 1401; and for Cu₂P₂, prepared by heating red P with excess of Cu phosphite and water, v. G., C. R. 117, 281 (abstracts in C. J. 62, 410 [1892]; and 64 [11], 526 [1893]).

Cupric sulphide. From experiments on the interaction of H₂S and CuSO₄Aq, and CuSO₅Aq in presence of acetic acid, Linder a. Picton (C. J. 61, 120 [1892]) conclude that a compound 7CuS.H₂S is formed, and that then such compounds as 9CuS.H₂S and 22CuS.H₂S are pro-

duced, and finally $(CuS)_n$ is formed.

copper. Ammonium compounds (vol. ii. p. 262). Several derivatives of cupriammonium bromide are described by Richards a. Shaw (P. Am. A. 28, 247 [1893]).

FERRICYANIDES.

Barium ferrioyanide (vol. ii. p. 837); v. Rammelsberg, J. pr. [2] 39, 455 (abstract in C. J. 56, 950 [1889]).

Lead ferricyanide (vol. ii. p. 339); v. Ram-

melsberg (l.c.).

Potassium ferrioyanide (vol. ii. p. 339); v. Kassner, Chem. Zeit. 13, 1701 (abstract in C. J. 58, 352 [1890]).

DIDYMIUM (vol. ii. p. 382). For an investigation into the separation of Di and Er, v. Krüss, A. 265, 1 (abstract in C. J. 60, 1424 [1891]). For the emission spectra of oxides of neodymium and praseodymium, v. Haitinger, M. 12, 362 (abstract in C. J. 62, 2 [1892]). Read (C. J. 65, 313 [1894]) says that Di₂O₂ is unchanged at c. 1750°.

ERBITM (vol. ii. p. 456). For the separation of compounds of Er and Di. v. Krüss, A. 265, 1 (abstract in C. J. 60, 1424 [1891]).

FERRITES (vol. ii. p. 547). J. van Bemmelen a. Klobbie (J. pr. [2] 48, 497 [1893]) say that Fe₂O₂, K₂O and Fe₂O₂, Na₂O are obtained, in crystals, by continued heating Fe₂O₂ with conc. KOHAq and NaOHAq respectively; these

ferrites are decomposed by water, giving Fe₂O₂.H₂O.

FLAME (vol. if. p. 549). Reference should be made to Smithells a. Ingle on 'The Structure and Chemistry of Flames' in C. J. 61, 204 [1892], and to Lewes on 'The Luminosity of Coal-gas Flames' in ibid. p. 822. FLUORHYDRIC ACID (vol. ii. p. 558).

FLUORHYDRIC ACID (vol. ii. p. 558). Berthelot a. Moissan (C. R. 109, 209 [1889]) give the thermal data [H,F] = 37,600; [H,F,Aq] = 49,400. For the description of an apparatus for purifying HFAq, v. Hamilton (C. N. 60, 252

[1889])

FLUORINE (vol. ii. p. 560). Moissan (C. R. 111,570 [1890]) determined the at. w. (1) by converting Na₂CO₃ into NaF, and this into Na₃SO₄; (2) by converting CaF₂ into CaSO₄, and (3) by converting BaF₂ into BaSO₄; the mean of the results regarded by M. as the most trustworthy was 19.05. V.D. at the ordinary temperature, 18.8 (Moissan, C. R. 109, 861; v. abstract in C. J. 58, 208 [1890]). A column of F one metre long shows no definite absorption bands (M., l.c. p. 937); for the emission spectrum v. M. (l.c.; abstract in C. J. 58, 329 [1890]). F does not liquefy at -95° at the ordinary pressure (M., A. Ch. [6] 25, 125 [1893]). Gladstone (P.M. [5] 31, 1 [1891]) has determined the atomic refraction of F in C.H.F to be as follows for the different lines of the solar spectrum, at 22.8°:-Ď \mathbf{F} Gr a H

at. refraction '63 '63 '53 '48 '44 '35 The values are much smaller than those for Cl, Br, or I, which are 10.0, 15.23, and 25.2 for the

ine A.

Formation.—According to Brauner (C. J. 65, 393 [1894]), F is given off when 3KF.PbF.HF (v. Lead tetraffluoride, p. 915) is heated to dull redness.

Reactions.—F combines with the more porous forms of carbon at the ordinary temperature, with incandescence, forming CF₄; the denser forms of C combine with incandescence at from 50° to 100° (Moissan, C. R. 110, 276; abstract in C. J. 58, 557 [1890]). For a discussion of the chemical relations of F, v. Moissan, Bl. [3] 5, 880 [1891].

GALLIUM (vol. ii. p. 597). Lecocq de Boisbaudran (C. R. 114, 815; abstract in C. J. 62, 930 [1892]) gives measurements of the chief lines in the spark emission spectrum of Ga; the spectrum varies much according to the conditions under which the spark is applied.

GERMANIUM (vol. ii. p. 610). A new mineral from Bolivia, canfieldite, Ag, GeS, identical in composition with argyrodite from Freiberg, but crystallising in isometric forms (the form of argyrodite is monoclinic), is described by Penfield in Am. S. [3] 46, 107 (abstract in

C. J. 66 [11], 18 [1894]).

dOLD (vol. ii. p. 647). Mallet (Pr. 46, 71; abstract in C. J. 58, 708 [1890]) has re-determined the at. w. of gold by various methods; the mean value obtained is 196.9, but the value 196.88 M. thinks is more trustworthy. The molecule of gold in dilute solution in tin is probably monatomic (Heycock a. Neville, C. J. 57, 876 [1890]). Gold volatilises considerably in the electric furnace with a purrent of 70 volts and 360 ampères (Moissan, O. R. 116, 1429 [1893]).

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Concerning the volatilisation of gold v. Rose [C. J. 63, 714 [1893]). Neumann (M. 18, 40 [1892]) says that gold absorbs from 38 to 48.5 times its volume of O at c. 450°. According to Petersen (Z. P. C. 8, 601 [1891]), the heats of formation of Au₂O₂.8H₂O from gold, (1) reduced from AuCl₂, (2) from AuHBr₄, and (3) from AuBr, differ very markedly. From the results of many experiments on the reaction between gold and KONAq, Maclaurin (C. J. 63, 724 [1893]) concludes that O must be present to accomplish solution, and that the equation 4Au+8KCNAq+O₂+2H₂O=

4(AuCN.KCN)Aq + 4KOHAq (Elsner, J. pr. 37, 333 [1846]) expresses the proportion between K

and Au in the solution.

Gold, alloys of (vol. ii. p. 648). For an alloy with Cd, AuCd, v. Heycock a. Neville (C. J. 61, 914 [1892]). For alloys with Cd and Sn, v. H. a. N. (C. J. 59, 936 [1891]). For alloys with Al and Sn, Cd and Bi, Cd and Pb, and Cd and Tl, v. H. a. N. (C. J. 65, 65 [1894]). For alloys with Cd and with Zn, prepared by immersing plates of Cd and Zn in solution of a salt of gold, v. Mylius a. Fromm (B. 27, 630; abstract in C. J. 66 [11], 285 [1894]).

Gold, bromides of (vol. ii. p. 648). Petersen (J. pr. [2] 46, 328) upholds the existence of Au₂Br₄ against the statements of Krüss a. Schmidt (B. 20, 2634); in a later paper (J. pr. [2] 47, 301) K. a. S. still maintain that AuBr, is the ultimate product of the reaction of Br with gold (v. abstracts in C. J. 64 [11], 126, 284). For the double salts AuBr. CsBr and AuBr. RbBr, v. Wells a. Wheeler (Am. S. [3] 44, 157 [1892]).

Gold, chlorides of (vol. ii. p. 649). Petersen (J. pr. [2] 46, 328; 48, 88) maintains that Au, Cl, is a definite compound; Krüss a. Schmidt (J. pr. [2] 47, 301; Zeit. f. anorg. Chemie, 3 421) say this substance is a mixture (v. abstracts in C. J. 64 [11], 126, 284, 474 [1893]).

Double salts of AuCl, with CsCl and RbCl are described by Wells a. Wheeler (l.c.); for a double salt AuCl₂.AgCl, v. Hermann (B. 27, 596; abstract in C. J. 66 [11], 234 [1894]).

Gold, sulphides of (vol. ii. p. 651). Concerning the conditions of formation of Au2S2, v. Antony a. Lucchesi (G. 19, 545; abstract in C. J.

58, 1216 [1890]).

Auric sulphide Au.S. This compound is formed, according to A. a. L. (G. 20, 601; 21 [11], 209; abstracts in C. J. 60, 526; 62, 280 [1892]), by passing H.S into a solution of AuCl. LiCl kept at -10°, extracting LiCl by absolute alcohol, and drying in N at 70°. Au2S. is a black, amorphous solid; decomposed by heat at 200°-205° to Au and S (for other reactions v. abstracts, l.c.)

Schneider (U. S. Geol. Survey Bull. No. 90, 56 [1892]) obtained aqueous solutions of colloidal

Au₂S and Au₂S₂.

HYDRAZOIC ACID (v. p. 919, Addenda). HYDROGEN (vol. ii. p. 719). The ratio between the densities of H and O is 1:15.882, the weight in grams of 1 litre of H is '08991 (Rayleigh, Pr. 53, 134 [1893]). For solubility in water, and in alcohol, v. Timofeeff (Z. P. C. 6, 141); Winkler (B. 24, 89); Henrich (Z. P. C. 9, 435); (abstracts in C. J. 60, 15, 884 [1891]; : 82, 1043 (1892]). For the line spectrum of H, v. Ames (P. M. [5] 30, 48 [1890]); and cf. Granwald (M. 18, 111; abstract in C. J. 62, 1381

Wilm (B. 25, 217; abstract in C. J. 62, 568 [1892]) describes a lecture experiment to show the occlusion of H by Pd. Regarding the occlusion of H by other metals v. Streintz (M. 12, 642; abstract in C. J. 62, 567 [1892]).

Baker (C. J. 65, 611 [1894]) found that a mixture of equal volumes of dry H and dry Cl did not explode in bright sunlight, and that more than a quarter of the mixed gases remained uncombined after exposure for two days to diffused light and two days to bright sunshine.

Reactions.—Neumann (Z. P. C. 14, 198 [1894]) has examined the reducing action of H occluded by Pt on solutions of salts of Al, Sb, As, Bi, Cd, Cu, Co, Au, Fe, Pb, Mg, Mn, Hg, Ni, Pd, Ag, Sn, Tl, and Zn. Regarding the rate of combination of H and O, v. WATER (p. 860). Regarding the explosion of mixtures of H with

O and Cl, v. CHLORINE (Addenda, pp. 904. 905). Hydrogen dioxide (vol. ii. p. 722). Schöne (B. 26, 3011; 27, 1233 [1893-4]) aftirms the occurrence of L2O2 in the atmosphere against the

contention of Ilosva (B. 27, 920).

Formation.—Regarding the formation from ether v. Dunstan a. Dymond (C. J. 57, 574, 988 [1890]). H₂O₂Aq is formed by the action of light on H.C.O.Aq, excess of O being present (Richardson, C. J. 65, 450 [1894]). Richardson (C. J. 63, 1110 [1893]) has examined the formation of

H2O2 in urine exposed to sunlight.

Preparation.—By making commercial 8 p.c. H₂O₂Aq distinctly alkaline by Na₂CO₂, filtering, shaking with 10-12 vols. of ether, separating the ethereal solution, evaporating this to 01-0025 of its original volume on a water-bath, and removing the rest of the ether by standing in a bell-jar over solid paraffin, Shiloff (J. R. 25, 293; abstract in C. J. 66 [11], 186 [1894]) obtained a thick, transparent, slightly acid liquid, S.G. 1.2475, containing 79.6 g. H₂O₂ in 100 c.c. For preparation of conc. H₂O₂Aq v. Talbot a. Moody (abstract in C. J. 64 [11], 869 [1893]).

Molecular weight. Tammann in 1889 (Z. P. C. 4, 441) by cryoscopic measurements determined mol. formula to be H₂O₄; but T.'s results were called in question by Carrara in 1893 (G. 22 [1], 341), who found mol. w. 33.92 (H2O2) from cryoscopic measurements, and this result was confirmed by Orndorff a. White (Am. 15, 347 [1893]), and also by Tammann himself

(Z. P. C. 12, 431 [1898]).

Reactions.—Regarding the reaction between H₂O₂Aq and KMnO₄Aq, v. Engel (Bl. [8] 6, 17; abstract in C. J. 62, 277 [1892]).

Hydrogen sulphide (vol. ii. p. 725). Henrich (Z. P. C. 9, 435 [1892]) gives the absorption coefficient of H₂S by water as 4.4015-089117t + $00061954t^2$. Hughes (P. M. [5] 88, 471 [1892]) says that H₂S dried by OaCl₂ and P₁ $^{\bullet}$ ₈ does not readen dry litmus paper, and does not react with BaO or Fe₂O₂, nor with salts of Sb, As, Bi, Cd, Co, Cu, Pb, Hg, Ag, or Sn. According to Pedler (C. J. 57, 625 [1890]), the equation 2H,S+80,=2SC,+2H,O correctly represents the explosion of H,S and air or O.

HYDROXYLAMINE (vol. ii. p. 784). NH₂OH

has been isolated, as a solid, by kobry de Balyn

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(R. T. C. 10, 100; 11, 18; abstracts in C. J. 62, 402, 1891 [1892]). NH2OH.HCl dissolved in MeOH is treated with CH,ONa solution, and most of the MeOH is distilled off at 100 mm. pressure; the residue is distilled in small portions at 40 mm. pressure (for details v. C. J., l.c.). For the preparation from 2NH₂OH.ZnCl₂ v. Crismer, Bl. [8] 6, 798 (abst. act in C. J. 62, 771 [1892]). For details of suitable apparatus for the fractionation of NH₂OH in vacuo, v. Brühl (B. 26, 2508 [1893]; cf. L. de B., B. 27, 967; and reply by B., ibid., 1347 [1894]). For thermal data of NH₂OH, v. Berthelot a. André (C. R. 110, 830; abstract in C. J. 58, 934 [1890]). Hydroxylamine is a white inodorous solid, melting at c. 33°, and remaining superfused at 0°; boils at 58° under 22 mm. pressure; decomposes at 90°-100°, and detonates at a higher temperature. S.G. 1.35; S.G. when liquid, 1.23. A drop heated in a test tube explodes very violently (v. Brühl, l.c.). B., l.c., discusses the molecular refraction of NH,OH. NH₂OH is fairly stable up to 15°, then gradual decomposition occurs, giving NH₂, HNO₂, and H₂N₂O₂, which react with unchanged NH₂OH producing N₂O and N (L. de B., B. 27, 967; abstract in C. J. 66 [11], 278 [1894]). For a full account of the reactions of NH2OH v. Lobry de Bruyn (l.c.; abstract in C. J. 62, 1391 [1892]). NH₂OH.HCl in water and NaNO₂Aq react to form Na₂N₂O₂Aq (v. Tanatar, J. R. 25, 342; abstract in C. J. 66 [11], 136 [1894]; of. Wislicenus, B. 26, 771; abstract in C. J. 64, 318 [1892]). For the reactions of 2NH OH H SO 318 [1892]). For the reactions of 2NH2OH.H2SO. with NaOHAq v. Kolotoff (J. R. 25, 295; abstract in C. J. 66 [11], 187 [1894]). For compounds of NH2OH with metallic salts, v. Crismer (Bl. [8] 8, 114); Goldschmidt a. Syngros (Zeit. f. anorg. Chemie, 5, 129); and Feldt (B. 27, 401) (abstracts in C. J. 58, 558 [1890]; 66 [11], 45, 187 [1894]). Kolotoff discusses the constitution of NH₂OH in J. R. 23, 8 (abstract in C. J. 64 [11], 114 [1893]). Kjellin (B. 26, 2377) describes NHEt.OH and NHMe.OH (abstract in C. J. 66 [11], 9 [1894]).

INDIUM (vol. iii. p. 1). The molecule of indium in dilute solution in tin is perhaps diatomic (v. Heycock a. Neville, C. J. 57, 376 [1890]). Linder a. Picton (C. J. 61, 134 [1892]) failed to isolate indium hydrosulphide, but they think that it is formed by passing H₂S into water with In₂O₂,xH₂O in suspension; the continued passage of H₂S produces In₂S₂. Read (C. J. 65, 313 (1894]) found that indium sesquioxide, In₂O₃, is unchanged at c. 1750.

IODHYDRIC ACID (vol. iii. p. 11). For the Mol. R. of HIAq from 20.77 to 67.02 p.c., v. Perkin (C. J. 55, 708 [1889]). For a research on the dissociation of HI, v. Bodenstein (Z. P. C. 13, 56 [1894]; abstracts in C. J. 64 [11], 369; 66 [11], 12 [1898-4]). Pickering (B. 26, 2807) has isolated three hydrates of HI: HI.2H₂O, relting at -43°; HI.8H₂O, melting at c. -48°;

and H1.4H₂O, melting at c. -36.5°.

IODINE (vol. iii. p. 14). The mol. w. of iodine dissolved in Et₂O or CS₂ was found by Bockmann (Z. P. C. 5, 76 [1890]) to be 254, from determinations of the boiling points; these results were confirmed by Sakurai (C. J. 61, 997 [1892]). By determining the lowering of the tp. of napht-alene by solution therein of I,

Hertz (Z. P. C. 6, 358 [1890]) found the value 254 for the mol. w. of iodine.

According to Meineke (Chem. Zeitung, 10, 1219, 1230), very pure I is prepared by covering ordinary I with a solution of CaCl₂ S.G. 1.35, mixed with a little conc. KIAq and a few drops of HClAq, heating until the I fuses, letting cool, washing the I, drying, and subliming it twice, the first time with addition of a little BaO.

For the reactions of I with KClO₂, and with KClO₂ and water, v. Potassium chlorate (Addenda, p. 905).

Iodine monochloride (vol. iii. p. 17). For thermal data bearing on the two forms of ICl, v. Stortenbeker (Z. P. C. 10, 183; abstract in C. J. 62, 1387 [1892]). Tanatar (J. R. 25, 97; abstract in C. J. 64 [11], 514) gives details of the methods for preparing the two forms of ICl.

Iodates (vol. iii. p. 21). For iodates of casium and rubidium, v. Wheeler, Am. S. [3] 44, 128 (abstract in C. J. 64 [11], 68 [1894]). Pure potassium iodate is prepared, according to Gröger (Zeit. angew. Chemie, 1894. 13), by heating to 100° for 20-30 minutes 20 g. KI in as little water as possible mixed with 40 g. pure KMnO₄ by cautious addition of alcohol, filtering, adding acetic acid till acid, evaporating to c. 50 c.o., washing the crystals of KIO₄ with conc. alcohol, and drying. By heating KIO₄ till it fuses, O and some I are given off, and KI finally remains; no other oxy-compounds are formed (Cook, C. J. 65, 802 [1894]).

IRIDIUM (vol. iii. p. 46). From analyses of the salts ICl₂.3KCl. aq and ICl₂.3NH₄Cl. aq, Joly deduced the value 192.75 for the at. w. of Ir (C. R. 110, 1131 [1890]). According to Prinz (C. R. 116, 892 [1893]), Ir crystallises in regular octahedra and in forms derived therefrom. Mylius a. Foerster (B. 25, 665 [1892]) stated that Ir is slightly volatilised when heated in a current of CO and Cl to c. 238°; Antony, however (G. 22 [11], 547 [1893]), asserts that no trace of any volatile compound of Ir is formed under these conditions.

Irid-ammonium salts (vol. iii. p.47). Palmaer (B. 22, 15; 23, 3810 [1889-90]) describes a number of irido-ammonium compounds. He calls them iridiumpentammine salts, and formulates them as Ir(NH₂), X¹₃: compounds are described where $X_2 = Cl_2$, $ClBC_3$, $ClSO_4$, $Cl(NO_3)_2$, $Cl(NO_2)_2$, ClC_2O_4 , $ClPtCl_6$; also where $X_2 = Br_3$, $BrSO_4$, and $Br(NO_3)_2$. One-third of the Cl in the salt when $X_3 = Cl_3$ is not removed by the action of cold H₂SO₄Aq or cold AgNO₂Aq; the reactions of the salts are similar to those of the purpureo-cobaltammines (v. vol. ii. p. 227) (v. abstracts in C. J. 56, 352 [1889]; 60, 402 [1891]). In B. 24, 2090 (v. abstract in C. J. 60, 1165 [1891]) P. describes compounds analogous with the roseo-cobaltic salts Co₂(NH₂), (H₂O), X¹¹, xaq; as the Ir salts are almost colourless, P. thinks the prefix roseo- is unsuitable; he calls them iridium aquopentammines, and formulates them as. $R(NH_s)_s(H_2O)X^r_s$. Salts are described in which $X_s = Br_s$, Cl_s , and $(NO_s)_s$. The chloride $Ir(NH_s)_s(H_2O)Cl_s$ is formed by boiling, for five hours, 5 g. Ir(NH_s), Cl_s with 11 times the calculated weight of KOH in 50 c.c. water, filtering, partially freezing the filtrate, adding 50 c.c. fuming HClAq, washing the pp. thus tormed

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with 22 p.c. cold HClAq and then with alcohol, dissolving in water, and again ppg. by conc. HClAq (for details v. abstract in O. J. 60, 1166

[1891])

Iridium tribromide (vol. iii. p. 48). Compounds of this bromide with PBr_s are described by Geisenheimer (C. R. 111, 40; abstract in C. J. 58, 1883 [1890]). By heating IrO₂xH₂O with IrBr_s and PBr_s, red needles of IrBr_s3PBr_s are formed; by heating this with PBr_s, black crystals of IrBr_s2PBr_s are produced.

Iridium tetrabromide (vol. iii. p. 48). Geisenheimer (l.c.) describes the compound IrBr₄.2PCl₂, obtained by heating IrBr₃.3PBr₃ with PCl₃.

Iridium trichloride (vol. iii. p. 48). The following compounds with PCl, are described by Geisenheimer (C. R. 110, 1004; abstract in C. J. 58, 1068 [1890]). IrCl, 3POl; IrCl, 2POl, ; IrCl, POL, POL. IrCl,.PCl,.2PCl,; Various compounds of IrCl, with H₂PO, and H₂PO, and salts of these, are also described. (C. R. 110, 1336; abstract in C. J. 58, 1069 [1890]) also describes the complex compounds 2IrP, Cl15.5AsCl3 and IrCl3.2PCl2.2AsCl3. For the salts IrCl, 3KCl. aq and IrCl, 3AmCl. aq, v. Joly (C. R. 110, 1131; abstract in C. J. 58, 1067 [1890]). For a description of IrCl_s. 3LiCl. xaq, v. Antony (G. 23 [1], 190; abstract in C. J. 64 [11], 380 [1893]). According to A. (G. 23 [1], 184; abstract in C. J. 64 [11], 379 [1890]), the passage of H₂S through a 2 p.c. solution of IrCl₂.3KCl produces pps. which are mixtures of Ir₂S₂ and IrS,

iridium dioxide (vol. iii. p. 50). For methods preparing IrO., v. Geisenheimer (C. R. 110.

of preparing IrO, v. Geisenheimer (C. R. 110, 855; abstract in C. J. 58, 948 [1890]).

Iridium disulphide (vol. iii. p. 50). IrS₂ is obtained by passing H₂S over IrCl₂.8LiCl at 4° to 7°, washing with absolute alcohol, and drying at 90° to 100° in a current of CO₂ (Antony, G. 23 [1], 190; abstract in G. J. 64 [11], 880 [1893]).

IEON (vol. iii. p. 51). Lockyer (Pr. 54, 359 [1893]) gives the emission spectrum obtained by using very fine electrolytically deposited iron as

the poles of an electric lamp.

Properties and Reactions.—Iron volatilises in the electric furnace, using a current of 350 ampères and 50 volts (Moissan, C. R. 116, 1429 [1893]). Iron is superficially oxidised by heating to dull redness in NO (Sabatier a. Senderens, C. R. 114, 1429 [1892]); heated in NO₂ to c. 350°, reduced iron is oxidised with incandescence to Fe₂O₂ (S. a. S., C. R. 115, 236 [1892]; cf. abstract in C. J. 66 [11], 95 [1894]). Montemartini (G. 22 [1], 250) says that NH₃, N₂O, and N are formed by the solution of iron in excess of 27.5 p.c. HNO₂Aq (for quantities, v. abstract in C. J. 62, 1278 [1892]). For quantities of iron dissolved by HNO₂Aq of different concentrations, v. Gautier a. Charpy (C. R. 112, 1451; abstract in C. J. 60, 1426 [1891]).

Passivity of iron. Concerning the passivities of steel and wrought iron to HNO₂Aq, v. Andrews (Pr. 48, 116; 49, 481; abstra ts in C. J. 60, 250, and 64 [11], 16 [1891 and 1893]).

Iron, bromochloride of. The compound FeBrCl, was obtained by Lenorm nd (C. R. 116, 820; abstract in C. J. 64 [11], 877) by heating FeCl, with excess of Br in a scaled tube at c. 100°. The compound is opaque in thin

sections, and green by reflected light; very deliquescent; crystallises probably in hexagonal plates; easily sol. alcohol and ether, CS₂, &c., can be sublimed by heating in a tube containing a little Br.

Iron carbide. Arnold a. Read (C. J. 65, 788 [1894]) confirm the existence of Fe₂C in steel (v. Abel, Proc. Inst. Mechan. Engineers, 1855. 30; also Müller, Stahl und Eisen, No. 5); A. a. R. say that two forms of Fe₂C are found in steel.

Iron carbonyls. In 1891 Mond a. Quincke (C. J. 59, 604) obtained a compound of Fe with CO; further investigation of the reaction by Mond a. Langer (C. J. 59, 1090 [1891]) led to the isolation of two compounds.

Ferropentacarbonyl Fe(CO)_s. Mol. w. 195·55. V.D. at $142^{\circ} = 93 \cdot 2$, S.G. $\frac{189}{199} = 1 \cdot ^{4}664$; distils completely, without accomposition, at $102 \cdot 8^{\circ}$ under 749 mm. pressure; solidifies at

Preparation.—FeC.O., ppd. by adding a slight excess of K.C.O.Aq to hot FeS.O.Aq, is well washed, and dried at 120°; it is then heated in a 3low stream of H, temperature being raised until the substance in the tube is black, and then kept constant until gas cease. to come off, when the current of H is stopped; the product is allowed to cool in H, placed in water without coming into the air, and treated with successive quantities of boiling water until sulphate is removed; it is then quickly dried on plates of gypsum, placed in the combustion tube, heated to c. 300° in H till quite dry, and allowed to cool in H. The tube is then con nected with a gas-holder containing CO, and the H is completely displaced by CO; one end of the tube is then sealed, and the open end is kept in connection with the supply of CO. After 24 hours the tube is heated to c. 120°, while a slow current of CO is passed through it, and the issuing gases are passed through a tube kept at -20° . When iron carbonyl ceases to come over, the tube is let cool, and is then again put in connection with the CO supply. These processes are repeated many times; the daily yield of the carbonyl is not more than c. 1 g. from 100 g.

Properties.—A pale-yellow, somewhat viscous liquid; unchanged in the dark, but in sunlight gives solid Fe₂(CO)₇ (v. infra); slowly decomposed on exposure to air, giving chiefly Fe₂O₃.xH₂O; completely decomposed at 180° to Fe and CO; soluble in EtOH, Et₂O, C₅H₆, mineraloils, &c. (For S.G., V.D., &c., v. supra.) Fe(CO)₅ is not acted on by dilute H₂SO,Aq, HClAq, or HNO₂Aq at the ordinary temperature; but is rapidly decomposed—to Fe(NO₃)₂Aq, FeOl₂Aq, or FeBr₂Aq—by conc. HNO₂Aq, ClAq, or BrAq.

Diferroheptacarbonyl Fe₂(CO)₇. Obtained by exposing Fe(CO), in a sealed tube to sunlight for some hours, collecting the solid that forms, washing with ether, and drying over H₂SO₄ for a short time. Yellow, lustrous flakes; gradually, decomposed by exposure to air, turning brown. Not acted on by H₂SO₄Aq or HClAq at the or, linary temperature; decomposed by HNO₂Aq. Cl, or Br.

Iron carbonyls seem to be present in water-

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gas; also sometimes in compressed coal-gas (v. Roscoe a. Scudder, C. J. Proc. 1891. No. 101).

Ferrous chloride (vol. iii. p. 53). Chassevant (A. Ch. [6] 30, 5 [1893]) describes the double salts FeCl, 2AmCl, FeCl, LiCl. 3aq, and

FeCl, 2KCl.

Ferric chloride (vol. iii. p. 54). By determining the r se of b.p. of alcohol and ether when FeCl, is dissolved in these solvents, Müller (C. R. 118, 644; abstract in C. J. 66 [11], 282 [1894], finds that the molecular formula of the chloride in these solutions is FeCl, FeCl, Aq reacts with many metallic sulphides, when heated therewith in sealed tubes; FeCl2Aq, S, and a chloride of the metal of the sulphide are generally formed (Cammerer, C. C. 1891 [11]. 370; abstract in C. J. 62, 18 [1892]). Seubert a. Dorrer (Zeit. f. anorg. Chemie, 5, 339, 411; v. abstracts in C. J. 66 [11], 140, 190 [1894]) have made a full investigation of the interaction of FeCl, Aq and KIAq; their results are in keeping with the hypothesis that the direct reaction $FeCl_{2}Aq + KIAq = FeCl_{2}Aq + KClAq + IAq$ is counterbalanced by the reverse reaction FeCl,Aq + KClAq + IAq = FeCl,Aq + KIAq. S. a. D. think that the first stage of the reaction is to produce an iodochloride; thus FeUi,Aq + KIAq = FeICl,Aq + KClAq; that this then decomposes, more or less completely (according to conditions of concentration, time, temperature, and mass), into FeCl,Aq and IAq; and that FeICl,Aq, and probably FeCl,Aq, are re-formed until equilibrium is established. S. a. D. do not agree with the explanation of the mechanism of the reaction given by Carnegie (v. vol. iii. p. 55). Roozeboom (Z. P. C. 10, 477 [1892]) has investigated the conditions of equilibrium in aqueous solution of the hydrates of FeCl.; he has isolated hydrates 2FeCl, xH,O where x=4, 5, 7, and 12. (For details, v. abstract in C. J. 64 [11], 119 [1893]). For a study of the interaction of FeCl, Aq and oxalic acid, v. Lemoine, C. R. 116, 981 (abstract in C. J. 64 [11], 405 [1893]). A compound of FeCl, with nitrosyl chloride, FeCl, NOCl, was obtained by Sudborough (C. J. 59, 660 [1891]), by immersing iron in liquid NOCl, and letting the liquid stand over H,SO,; the compound forms small golden-brown, deliquescent crystals (cf. vol. iii. p. 56; Combinations, No. 7)

Ferrous fluoride (vol. iii. p. 56). FeF₂ was obtained by Poulenc (C. R. 115, 941; abstract in C. J. 64 [11], 122 [1893]) by passing air-free HF over red-hot iron or dry FeCl₂; small, white, rhombic prisms; S.G. 4·09; decomposed by heating with H₂SO₄, Na₂CO₂, steam, air, HCl,

or H.

Ferric fluoride (vol. iii. p. 56). Poulenc (l.c.) obtained FeF_s, in small, very refractive, greenish crystals, S.G. 3°87, by passing HF over red-hot iron, Fe₂O_s, FeF_s, xaq, or FeCl_s; also by dropping FeF_s, xaq into fused AmCl and heating in a current of an inert gas. FeF_s does not fuse at 1000°; but is decomposed by heating in steam, or with HCl or H_sSO_s. According to Speransky (J. R. 24, 304; abstract in C. J. 64 [11], 314 [1893]), FeF_sAq shows a very small electrical conductivity; there is, therefore, no appreciable electrolytic dissociation; as the solution scarcely effects any inversion of cane sugar, S. concludes You. IV.

that there is practically no hydrolytic dissociation. Cryoscopic determinations showed that the molecules in a dilute aqueous solution are chiefly FeF_s, but indicated the existence of Fe_zF_s in more conc. solutions.

Iron, nitrides of (vol. iii. p. 59). Fowler (C. N. 68, 152 [1893]) obtained Fe₂N by heating reduced iron in a rapid stream of NH₃. (For details v. abstract in C. J. 66 [11], 50 [1894].)

Ferric oxide (vol. iii. p. 61; also p. 57 for hydrates). At c. 1750° Fe₂O₃ is completely changed to Fe₂O₄ (Read, C. J. 65, 313 [1894]); in the electric furnace Fe₂O₄ is formed, partly melted and partly crystallised (Moissan, C. R. 115, 1034 [1892]). According to J. van Bemmelen a. Klobbie (J. pr. [2] 46, 497 [1893]), crystallised Fe₂O₃.H₂O is obtained by treating with water the hexagonal crystals of Fe₂O₃.Na₂O formed by heating Fe₂O₄ for some time with conc. NaOHAq; the crystalline Fe₂O₃.H₂O is not hygroscopic; H₂O is given off below 100°.

Iron, oxychlorides of (vol. iii. p. 62). Rousseau (C. R. 110, 1032; 113, 643; 116, 392; abstracts in C. J. 58, 1063 [1890]; 62, 119 [1892]; 64 [11] 280 [1893]) describes compounds xFeCl₂yFe₂O₃, some crystalline and some amorphous, formed by heating FeCl₂Aq at different temperatures, and also by the interaction of FeCl₂ and steam.

LANTHANUM (vol. iii. p. 116). Bettendorff (A. 256, 159 [1890]) obtained the value 138-28 for the at. w.; and Brauner (B. 24, 1328 [1891]) the value 138-21 (O = 16).

Winkler (B. 24, 873 [1891]) obtained results, by heating La₂O, with Mg in H, which pointed to the existence of a hydride of La (cf. Brauner,

Lead (vol. iii. p. 122). The molecule in dilute solution in tin is probably monatomic (Heycock a. Neville, C. J. 57, 376 [1890]). The m.p. of lead was found by Griffiths a. Calendar (using a Pt thermometer) to be 327.69° (C. N. 63, 1 [1891]). Lehmann (Z. K. 17, 274 [1890]) says that lead separated electrolytically, by a weak current, crystallises in leafiets that are probably monosymmetric, but that the crystals are regular octahedra when separated by a strong current. Veley (S. C. I. 10, 206; summary of conclusions in C. J. 62, 410 [1892]) has examined the interaction of lead and nitric acid: an acid of intermediate concentration acts most rapidly; when HNO₂ is present the action increases. Montemartini (G. 22, 384, 397, 426 [1892]) found that a little NH₂ was produced, especially with dilute HNO₂Aq. Lead rapidly oxidises to PbO when heated in NO; oxidation occurs at c. 200° in NO₂, with formation of a basic nitrate (Sabatier a. Senderens, C. R. 114, 1429; 115, 236 [1892]).

Lead, bromide of (vol. iii. p. 125). Several double compounds of PbBr, with alkali bromides have been described. For PbBr, 2AmBr. aq and 2PbBr, AmBr, v. Wells a. Johnston, Am. S. [3] 46, 25 (abstract in C. J. 64 [11], 523 [1893]); for PbBr, KBr. xaq and PbBr, 2KBr. aq, v. Herty, Am. 15, 357 (abstract in C. J. 66 [11], 465 [1894]); for 2PbBr, RbBr and 2(PbBr, 2RbBr). aq v. Wellr, Am. S. [3] 46, 34 (abstract in C. J. 66 [11], 324 [1893]); and for K, Pb, Br, 4aq and K, Pb, (BrI), 4aq, v. Wells, Lc. p. 190 (abstract,

Lc. p. 524).

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Lead, chlorides of (vol. iii. p. 125). The statement, on p. 125 (vol. iii.), that only one chloride has been isolated is no longer true; PbCl, was prepared by Friedrich in 1893.

Lead tetrachloride (vol. iii. p. 126). PbCl, has been obtained by Friedrich (B. 26, 1434; abstract in C. J. 64 [11], 415 [1898]; cf. M. 14, 505; abstract in C. J. 66 [11], 16 [1894]), by passing Cl into HClAq containing PbCl, in suspension, adding NH,Cl to the solution thus obtained, separating and drying the compound PbCl, 2NH, Cl thus formed, adding it to conc. cooled H2SO4, allowing the oily drops that formed to collect at the bottom of the vessel, and shaking these drops repeatedly (and quickly) with conc. H2SO4. PbCl4 is a transparent, yellow, very refractive liquid, fuming in moist air with formation of PbCl, and Cl; it is unchanged in contact with cold conc. H2SO4, but when heated with the acid it decomposes explosively to PbCl2 and Cl; when heated with conc. H2SO, in a current of Cl, some PbCl, passes over, but at c. 105° explosion occurs. PbCl, has S.G. 3.18 at 0° ; at c. -15° it solidifies to a yellowish, crystalline mass. A *hydrate* is formed with a little water; but addition of more water produces HClAq and PbO₂. By adding PbCl₄ to a little cooled conc. HClAq, a crystalline compound, probably PbCl₄.2HCl, is formed.

Several double salts of PbCl, with alkali chlorides have been isolated. Classen a. Zahorski (Zeit. f. anorg. Chemie, 4, 100) give the formula 2PbCl,.5AmCl to the salt formed by digesting PbCl, with liquid Cl and fuming HClAq, and adding AmCl; but Friedrich (l.c.) says the compound is PbCl, 2NH, Cl. For compounds PbCl, 2MCl, when M=NH, Cs, K, and Rb, v. Wells (Am. S. [3] 46, 180; abstract in C. J. 64 [11], 523 [1893]); C. a. Z. (l.c.) describe compounds of PbCl, with the hydrochlorides of pyridine and quinoline (abstract in C. J. 64 [11],

464 [1893]).

Lead dichloride (vol. iii. p. 125). Several double compounds with alkali chlorides are described by Randall (Am. 15, 494 [1893]); Wells a. Johnston (Am. S. [8] 46, 25 [1893]); and Wells (l.c. p. 84). The compounds are of the forms PbCl_2MCl, PbCl_8MCl.caq, and 2PbCl_MCl (v. abstracts in C. J. 64 [11], 523, 524 [1893]).

Lead, fluorides of (vol. iii. p. 127). Brauner (C. J. 65, 393 [1894]) has prepared the compound PbF. 8KF.HF in various ways; the simplest process being to dissolve Pb(C2H2O2)4 v. RED OXIDE OF LEAD, Addenda, infra) in conc. HFAq and add KF. By acting on this salt with conc. H₂SO₄, B. obtained evidence of the formation of lead tetrafluoride, PbF, but he did

not succeed in isolating the compound.

Lead, iodide of (vol. iii. p. 127). Several double compounds of PbI, with alkali iodides have been isolated by Wells, W. a. Johnston, and Herty (v. References under LEAD BROMIDE, p. 914). For the compound PbI, 2MgI, 16aq, v. Otto a. Drewes (Ar. Ph. 229, 179; abstract in C. J.

64 [11], 984 [1893]).
Lead iodobromides and iodochlorides (vol. iii. p. 128). Miss Field (C. J. 68, 540 [1893]) has described the compounds 2PbBr. PbI. 2PbCl2.PbI2, and 5PbCl2.PbI2.

[1894]). The hydrate 2PbO.H.O (vol. iii. p. 129) is obtained in large crystals by allowing a solution of PbO in KOHAq to cool in the air; CO, is absorbed, and the hydrate is deposited in colourless, tetragonal crystals, a:b:c = 1: .824(Luedeking, Am. 13, 120 [1891])

Red oxide of lead (vol. iii. p. 130). Hutchinson a. Pollard (C. J. 63, 1136 [1873]) have examined the reaction of Pb,O4 with glacial acetic and have isolated lead tetracetate Pb(C₂H₂O₂), melting at c. 175°, and decomposing a little above that temperature.

Note.—The formula PbO(OC2H3O)2 said to be assigned by Jacquelain to the salt obtained by him (vol. iii. p. 130, col. 2, line 28 from top; and p. 131, col. 2, line 24 from bottom) should have been given as PbO₂(C₄H₃O₃)₃ old notation.

Plumbates (vol. iii. p. 132). Kassner (Ar. Ph. [8] 28, 109; abstract in C. J. 58, 561 [189J]) Kassner (Ar. describes plumbates of Ba, Ca, and Sr. Regarding the dissociation of Ca plumbate, v. Le Chatelier (C. R. 117, 109; abstract in C. J. 64 [11]<u>,</u> 524 [1898])

Lead oxyiodides (vol. iii. p. 132). Gröger M. 13, 510; abstract in C. J. 62, 1892) describes

the compound PbO.PbI2.

Lead, salts of (vol. iii. p. 132). The salt Pb(C₂H₃O₂)₄ (v. supra, Red oxide of lead) belongs to the type PbX¹₄, to which form PbCl₄

also belongs.

Lead, sulphide of (vol. iii. p. 133). According to Hannay (C. J. Proc., Nos. 139 and 141 [1894]), when air is passed into molten PbS, one half of the lead remains as lead, and one half is volatilised, and deposited as PbS; H. accounts for this change by supposing the reaction to be 2PbS + O2 = Pb + PbS.SO2, the volatile PbS.SO2 being decomposed in the colder part of the apparatus to PbS and SO₂.

Lead sulphobromide and sulphochloride (vol. iii. p. 183). Parmentier (C. R. 114, 298; abstract in C. J. 62, 685 [1892]) obtained the compounds PbS.PbBr, and PbS.PbCl, by dissolving the halides in their acids and shaking the liquids while H2S was passed over the sur-

faces of the solutions.

LITHIUM (vol. iii. p. 147). Guntz (C. R. 117, 782 [1893]) recommends to use a mixture of equal weights of LiCl and KCl, for the electrolytic preparation of the metal; the mixture melts at 450° and the m.p. decreases as electrolysis proceeds. (For details as to size of electrodes, strength of current, &c., v. abstract in C. J. 66 [11], 91 [1894].) Holt a. Sims, C. J. 65, 443 [1894], find that traces of a peroxide are formed, along with Li₂O, by burning Li in O.

Lithium, amide of, LiNH2. Formed by gently heating clean Li in a stream of NH, until violent action ceases, and then heating to c. 400° until the reaction is finished. A white, crystalline, transparent solid; melts between 380° and 400°; decomposes in cold water slowly, giving LiOHAq and NH3; decomposed by heating in air, but without taking fire litherley, C. J. 65; 504 [1894]; q. v. for further reactions).

Lithium, bromide of (vol. iii. p. 150). The hydrates LiBr. xaq, where x = 1 and 2, are de-

scribed by Bogorodsky (J. R. 25, 316 (1893)).

Lithium, chloride of (vol. iii. p. 150). For
LiCl. aq and LiCl. 2aq v. Bogorodsky (l.c.). For Lead protoxide (vol. iii. p. 128). PbO is LiCl. aq and LiCl. 2aq v. Bogorodsky. (l.c.). For unchanged at c. 1750° (Read, C. J. 65, 818) the double compounds LiCl.MCl. xaq, where

M = Cu, Co, Fe, Mn, or Ni, v. Chassevant (C. R.118, 646; 115, 113; abstracts in C. J. 62, 118,

1275 [1892]).

Lithium, fluoride of (vol. iii. p. 150). Poulenc (Bl. [3] 11, 15 [1894]) obtained LiF in regular octahedra by heating the amorphous fluoride with KHF, and KCl. and washing with water; sl. sol. wate, insol. 95 p.c. alcohol; melts at c. 1000° in H, and partially volatilises at 1100°— 1200°

Lithium, hydroxide of (vol. iii. p. 150). For the freezing-points of aqueous solutions of LiOH

v. Pickering (C. J. 63, 899 [1893]).

Lithium, nitride of. Ouvrard (C. R. 114, 120 [1892]) obtained a black solid, probably Li, N, by heating commercial Li to dull redness in N.

Lithium, exides of (vol. iii. p. 151). Almost the only product of heating Li in O is Li₂O; only traces of a peroxide are formed (Holt a. Sims, C. J. 65, 443 [1894]).

MAGNESIUM (vol. iii. p. 157). Burton a. Vorce (Am. 12, 219 [1890]) obtained the value 24.22 (O = 15.96) for at. w., by converting pure Mg into Mg(NO₃)₂ and this to MgO. The molecule of Mg in dilute solution in win is probably monatomic (Heycock a. Neville, C. J. 57, 376 [1850]). Mg crystallises in holohedral hexagonal forms, a:c=1:1.6202, isomorphous with

Zn and Be (B. a. ∇ ., l.c.).

According to Montemartini (G. 22, 384, 397, 426 [1892]), a considerable quantity of H is given off when Mg interacts with HNO, Aq (c. 13 p.c.); the quantity of NH, formed increases with concentration of the acid up to 40 p.c. HNO, after which it decreases. Mg burns in NO at a dull red heat (Sabatier a. Senderens, C. R. 115, 236 [1892]). Giorgis (G. 21, 510 [1891]) says that Mg dissolves in water saturated with CO2; that H is given off, and crystals of MgCO, 3aq are deposited. For the interactions of Mg and chlorides v. Seubert a. Schmidt (A. 267, 218; abstract in C. J. 62, 776 [1892]). Winkler has made an extended examination of the interactions of Mg and oxides (B. 23, 44, 120, 772, 2642; 24, 878, 1966; abstracts in C. J. 58, 831, 451, 693, 1372 [1890]; 60, 801, 1155 [1891]).

Magnesium, bromide of (vol. iii. p. 159). Beketoff (v. abstract in C. J. 62, 762 [1892]) gives H.F. [Mg,Br²] = 121,700. The double salt 2MgBr₂.PbBr₂.16aq is described by Otto a. Drewes (Ar. Ph. 229, 585; abstract in C. J. 62, 566

[1892]).

Magnesium, chloride of (vol. iii. p. 159). Skinner (C. J. 61, 341 [1892]) gives the boilingpoints of solutions of MgCl₂ in water and alco-

hol, from c. 4.5 to c. 14.5 p.c. MgCl... Magnesium, iodide of (vol. iii.

Beketoff (l.c.) gives H.F. [Mg,I'] = 84,800. Magnesium, nitride of (vol. iii. p. 161). For

description of an experiment to demonstrate the formation of Mg, N, v. Merz (B. 24, 8,940; ab-

stract in C. J. 62, 409 [1892]]

Magnesium, oxide of (vol. iii. p. 161). MgO is unchanged at c. 1750° (Read, C. J. 65, 318 [1894]); heated in the electric furnace, it melts with a current of 70 volts and \$50 ampères (Moissay, C. R. 115, 1084; cf. C. R. 116, 1429 1892-8j). Richards a. Rogers (P. Am. A. 28, 200 [1898]) found that MgO, prepared by evaporating MgCO, or Mg, with HNO, Aq and Leating very strongly, retained c. 10 times its volume

of gas, chiefly O and N.

Magnesium, silicides of (vol. iii. p. 162). Regarding the formation of compounds by heating together Mg and Si, v. Winkler (B. 23, 2642;

abstract in C. J. 58, 1873 [1890]).

MANGANESE (vol. iii. p. 177). Moissan (C. R. 116, 849 [1893]) obtained Mn containing to 5 p.c. C, by fusing MnO with charcoal (keeping MnO in excess) by means of a current of 60 volts and 300 ampères. Prelinger (M. 14, 353; abstract in C. J. 66 [11], 49 [1894]) prepared pure Mn by heating Mn amalgam (formed by electrolysing MnCl.Aq using a cathode of

Hg) in pure H.

Mn is described by P (l.c.) as a grey porous metal; unchanged in dry air; reacts slowly with cold, rapidly with warm, water; dissolves in NH ClAq giving off NH, and H; S.G. 74212 referred to water at 4°. Moissan (C. R. 116, 1429 [1893]) found that Mn volatilises readily in the electric furnace, using a current of 80 volts and 380 ampères. Montemartini (G. 22, 384, 397, 426 [1892]) says that the gaseous products of the interaction of Mn and HNO, Aq are H, NH, N, and N2O. Finely divided Mn reacts with CO at c. 400°, producing MnO and C (Guntz, C. R. 114, 115 [1892]).

Manganese, alloys of. The presence of c. 5 p.c. Mn in iron or steel scarcely affects the properties, except by slightly increasing the tenacity and malleability; a very hard tool-steel is formed with c. 1} p.c. Mn. Alloys with iron containing from 7 to 30 p.c. Mn are extremely

hard and tough.

Manganese, amalgams of (vol. iii. p. 179). Prelinger (M. 14, 353; abstract in C. J. 66 [11], 49 [1894]) prepared Mn₂Hg, by passing a current from a kathode of Hg through saturated MnCl.Aq to an anode of C, or Pt-Ir, in a porous vessel, washing in running water, squeezing out excess of Hg, and drying in H over CaCl. heating gently in a stream of pure, dry H, pure Mn was obtained

Manganous chloride (vol. iii. p. 179). Chassevant (A. Ch. [6] 30, 5 [1893]) describes several double salts of MnCl₂; viz. MnCl₂AmCl. 2aq, MnCl₂2AmCl. aq, and MnCl₂LiCl. 8aq. For the double salts M.KCl. 2aq, M.CsCl. 2aq; M.2XCl. 2aq, X-NH, Cs, Rb; 2M.MgCl, 12aq $(M = MnCl_2)$, v. Saunders (Am. 14, 127; abstract in C. J. 62, 780 [1892]).

Manganic chlorides (vol. iii. p. 180). Begarding the composition of the chloride in the solution obtained by dissolving MnO, in HClAq, v. Vernon (P. M. [5] 31, 469), and Pickering (P. M. [5] 33, 284) (abstracts in C. J. 62, 19,

687 [1892]).

Manganese peroxide (vol. iii. p. 183). Begarding the compositions of the hydrated oxides, approximately MnO2 xaq, prepared by different methods, v. Gorgeu (Bl. [8] 4, 16 [1891]). G. (C. R. 110, 857; abstract in C. J. 58, 946 [1890]) gives some details regarding the interaction of MnO, xaq and H,O,Aq. Heated in the electric furnace, MnO₂ melts, gives off O, and leaves MnO (Moissan, C. R. 115, 1034 [1892]). Mn₂O₃ in said to be formed by heating MnO₃ to c. 400° in NO (Sabatier a. Senderens, C. R. 114, 1476 [1892]). McLeod (C. J. 65, 202 [1894]) has confirmed his former observation that Cl is

given of when MnO₂ and KClO₂ are heated together. Regarding compounds of MnO₂ with basic oxides, v. Rousseau (C. R. 112, 525; 114, 72; 116, 1060; abstracts in C. J. 60, 645 [1891];

62,569 [1892]; 64 [11], 416 [1893]).

Permanganates (vol. iii. p. 186). Regarding the interaction of permanganates with H,O2Aq, v. Gorgeu (C. R. 110, 958; abstract in C. J. 58, 1062 [1890]). Klobb (Bl. [3] 3,508; abstract in C. J. 58, 947 [1890]) describes compounds of NH, with permanganates of Cd, Cu, Ni, and Zn.

Barium permanganate BaMn₂O₈ (the formula is wrongly printed BaMnO, in vol. iii. p. 186). For the preparation of this salt from KMnO,Aq and Ba(NO_s), Aq v. Muthmann (B. 26, 1016; abstract in C. J. 64 [11], 324 [1893]).

Calcium permanganate. The formula given on p. 186 of vol. iii. is wrong; it should be

 $CaMn_2O_8.5H_2O$.

Potassium permanganate (vol. iii. p. 186). Tivoli (G. 19, $6\overline{3}0$ [1890]) gives the following as the reactions that occur when pure AsH, is passed into dilute KMnO,Aq: (1) 2KMnO,Aq+AsH,

2KMnO₄Aq + AsH₂ = K₂HAsO₄Aq + 2MnO₂ + H₂O.

2KMnO₄Aq + AsH₃ = K₃HAsO₄Aq + 2MnO₂ + H₂O.

Silver permanganate (vol. iii. p. 187).

AgMnO₄ decomposes slowly at the ordinary temperature, rapidly at 100°, and very rapidly at 185° (Gorgeu, C. R. 114, 912; abstract in C.J. 62, 942 [1892]).

Manganous sulphide (vol. iii. p. 188). The pp. produced by adding (NH₄)ĤSAq to an ammoniacal solution of a Mn salt, after washing in an atmosphere of H,S and drying in CO. at 70°, is MnS; S.G. $\frac{170}{170} = 3.55$. By heating to 300°-320°, or by leaving in contact with (NH₄)HSAq for some days, the reddish MnS becomes green without changing its composition; the green form of MnS is crystalline, S.G. $\frac{170}{177}$ = 3.63 (Antony a. Donnini, G. 23 [1], 560 [1892]).

MASRIUM. In 1892 (C. J. 61, 491) Richmond a. Husseinoff obtained indications of the existence of an element before unknown in specimens of Egyptian fibrous alums. The supposed new element gave reactions indicating analogies with Be, Ca, and Zn; the results obtained by decomposing the oxalate by heat led to the at. w. of c. 228. The authors suggested the name masrium, from the Arabic name for Egypt.

MERCURAMMONIUM COMPOUNDS (vol.

iii. p. 206).

Mercuro-ammonium salts NH_{4-x}Hg_xX (vol. iii. p. 207). Barfoed's conclusion that these supposed compounds are really mixtures of mercuricon pounds and Hg is confirmed by Pesci (G. 21 [11], 569 [1891]). The product of the interaction of HgCl and NH,Aq, described as dimercuro-ammonium chloride NH, Hg, Cl in vol. iii. (p. 208), leaves metallic Hg when treated with ammoniacal (NH₁)₂SO₄Aq, according to P. When the reaction of H_PCl with NH₂Aq proceeds out of the light, P. says that the change is 2Hg₂Cl₂+4NH₃Aq = (NHg₂)Cl.NH₂Cl+2NH₄ClAq+2Hg. Similarly the reaction of NH_Aq with Hg_SO, produces a double compound of dimercuro-ammonium sulphate ((NHg,),SO, (NH,),SO,) and Hg; and (NHg,)NO, NH,NO, and Hg are formed, according to P., when HgNO,Aq is treated with NH,Aq.

Mercuri-ammonium salts NH, Hg.X. Regarding the conditions of formation of compounds of mercuri-ammonium chloride NH, Hg.Cl (vol. iii. p. 208) with HgCl, and NH (Hg₂O)Cl, v. Balestra (G. 21 [11], 294; abstract in C. J. 62, 276 [1892]).

Dimercuri-ammonium salts NHg₂X (vol. iii. p. 209). Pesci (G. 20, 485 [1890]) has examined the conditions of formation of various salts of this class, especially the nitrate NHg, NO, and compounds of this with NH, NO, The sulphate pounds of this with NH,NO, The sulphate (NHg,)2SO, 2aq was formed by saturating conc. NH, Aq with HgSO, allowing the solution to evaporate in an atmosphere of laH, over CaO, and collecting the first crop of crystals; various compounds of (NHg₂)₂SO₄ with (NH₁)₂SO₄ are also described (v. abstract in C. J. 60, 268 [1891]).

MERCURY (vol. iii. p. 212). Heycock a. Neville's results (C. J. 57, 376 [1890]) show that some condensation from monatomic molecules perhaps occurs in dilute solutions of Hg in tin. For corrected tables of the vapour pressures of Hg for each 10° from 180° to 480°, and each 1° from 330° to 360°, v. Young (C. J. 59, 629

[1891]).

Jäger (W. 48, 209 [1893]) recommends to prepare pure Hg by converting Hg that has been distilled in vacuo into HgNO, and electrolysing a solution of the salt, using Pt as cathode and distilled Hg as anode. Shenstone (C. J. 61, 452 [1892]) noticed that Hg adheres to glass when brought into contact with Cl, Br, or I. Montemartini (G. 22, 384, 397, 426 [1891]) finds that the amount of NO produced by the interaction of Hg and HNO₃Aq decreases as concentration of the acid increases; that NO, is produced with acid more conc. than 27.5 p.c. $\dot{H}NO_3$, $HgNO_3$ is formed with 25 p.c. acid, but $Hg(NO_3)_2$ with 50 to 70 p.c. acid; no NH, is formed with 27 and 50 p.c. acid.

Mercuric bromide (vol. iii. p. 215). HgBr₂ boils at 325°, according to Freyer a. V. Meyer

(Zeit. f. anorg. Chemie, 2, 1 [1892]).

Mercurous chloride (vol. iii. p. 215). Harris a. V. Meyer have re-determined the V.D. of calomel at 448° and 518°; he value obtained was 117.5. H. a. M. obtained considerable quantities of Hg and HgCl, by allowing the vapours obtained by heating calomel to c. 465° to diffuse along a glass tube; also by heating calomel at 420° for half an hour in a distillation flask at 33 mm. pressure. By heating calomel to 240°-260°, and bringing a piece of KOH, which had been heated to c. 250°, into the vapour, they noticed that the surface of the KOH at once became covered with yellow HgO; they also found that a piece of gold leaf held for a moment in the vapour was amalgamated (if kept for some time in the hot vapour the yellow colour was restored). H. a. M. think their results leave no doubt that the vapour obtained by heating calomel is Hg+HgCl2, and that the molecular formula of calomel is Hg, Cl

The reaction of Hg₂Cl₂ with NH₃Aq gives (NHg₂)Cl.NH₄Cl and Hg, according to Pesci (v.

supra, Mercuro-ammonium salts).

Mercuric chloride (vol. iii. p. 217). HgCl. boils at 307° (Freyer a. V. Meyer, Zett. f. anorg. Chemic, 2, 1 [1892]). The white substance produced by the reaction of HgCl,Aq with a silver photographic image, or with pulverulent Ag, is HgAgCl, according to Jones (S. C. I. 12, 983; abstract in C. J. 66 [11], 138 [1894]).

Mercuric cyanide; for an account of the conditions of formation and properties of the double mercury-zinc cyanide, v. Dunstan (C. J.

61, 666 [189]).

Mercury, haloid compounds of (vol. iii. p. 219). Wells (Am. S. [3] 44, 221; abstract in C. J. \$4 [11], 68 [1893]) describes a number of double compounds of mercury and cæsium halides; the compounds belong to the forms (1) Cs_2HgX_5 , (2) Cs_2HgX_4 , (3) $Cs_2Hg_3I_6$, (4) $CsHgX_3$, (5) CsHg₂X₃, (6) CsHg₂X₁₁; X = Cl, Br or I; in some cases the total X atoms comprise two different halogens.

Mercury hydrosulphide; for experimental attempts to prepare a definite compound, v.

Linder a. Picten (C. J. 61, 123 [1892]).

Mercury phosphide (vol. iii. p. 223). By heating Hg and PI, in a sealed tube for many hours at 275°-300°, and removing Hgl, by treatment with KIAq, Granger (C. R. 115, 229 [1892]) obtained Hg₂P₂ in lustrous, metal-like, hexagonal crystals, appearing red by transmitted light, and giving a brown powder. The compound gives Hg and P when heated; heated in air it burns; detonates when mixed with KClO2 and struck; burns in Cl; dissolved by

Mercurous sulphide (vol. iii. p. 224). Antony a. Sestini (G. 24 [1], 193 [1894]) obtained a black powder by passing dry H.S. diluted with CO₂, over dry Hg.Cl₂ or Hg.(C₂H₂O₂), kept at -10°; when freshly prepared the black powder was scarcely changed in dry HCl at -18°, but at a somewhat higher temperature (below 0°) Hg₂Cl₂ was quickly formed. The substance dissolved in cold K₂SAq or Na₂SAq, but when temperature rose above 0° the solution became turbid and ppd. Hg. A. a. S. think their results show that Hg.S exists at low temperatures (abstract in C. J. 66 [11], 318 [1894]).

Mercury sulphochloride (vol. iii. p. 225). The compound 2HgS.HgCl, is formed, according to Colson (C. R. 115, 657 [1892]), by passing H₂S into a solution of HgCl₂ in dry benzene.

MOLYBDENUM (vol. iii. p. 421). Smith a. Mass (A. C. J. 15, 397 [1893]) obtained the mean value 95.85 (O=15.96) for the at. w. of Mo, by heating Na2MoO, in a current of dry HCl and weighing the NaCl that remained (MoOs.2HCl and H2O were formed and removed). Smith a. Oberholtzer (Zeit. f. anorg. Chemie, 5, 63 [1893]) say that a mixture of MoOCl, and Mo(CO)Cl, is formed by heating Mo to 150°-200° in COCl2; and that Mo.S.Cl, is produced by heating Mo to redness in S₂Cl₂. Mo is said to ppt. Ag from AgNO₂Aq, Au from AuCl₂Aq, HgCl and then Hg from HgCl₂Aq, and Cu from solutions of its salts (Smith, Zeit. f. anorg. Chemie, 1. 360 [1891]). Montemartini (G. 22 384, 397, 426 [1891]) finds that no NH, is formed in the interaction of Mo and HNO, Aq with from 3 to 70 p.c. HNO,; the most vigorous action occurs with acid c. 50 p.c., but the whole of the metal is not oxidised to MoO, a nitrate being probably formed; the gaseous products are NO and NO, the quantity of NO decreasing as the concentration of the acid increases.

Molybdenum, carbides of. By heating a mixture of MoO, and charcoal in the electric furnace, Moissan (C. R. 116, 1225 [1893]) obtained a very hard carbide, containing from 9.77 to 9.9 p.c. C; S.G. 8.6.

Molybdenum sesquioxide (vol. iii. p. 431). Heating Mo.O. in NO produces MoO. (Sabatier

a. Senderens, C. R. 114, 1429 [1892]).

Molybdenum trioxide (vol. iii. p. 432). MoO. partially volatilises, and is partially reduced to one of the blue oxides, at c. 1750° (Read, C. J. 65, 818 [1894]). According to Smith a Oberholtzer (Zeit. f. anorg. Chemie, 4, 236 [1893]), the products of heating MoO, in HBr are (1) MoO, 3HBr, and (2) Mo,O,Br,; when HI is used and the temperature kept at 105°-120°, a blue oxide Mo_.O_.. xaq is formed; HF at 800°-400° forms Mo_.O_.F₄ (v. abstract in C. J. 64 [11], 471 [1893]). Cammerer (Chem. Zeit. 15, 957; abstract in C. J. 62, 944 [1892]) says that 2MoO. H2O2.H2O is formed by boiling MoO, with H_2O_2Aq .

Molybdenum, oxybromides of (vol. iii. p. 438). Smith a. Oberholtzer (Zeit. f. anorg. Chemie, 4, 236 [1893]) describe Mo₂O₂Br₄; square plates, colour of KMnO₄. Formed, along with

MoO.3HBr, by heating MoO. in HBr.

Molybdenum, oxychlorides of (vol. iii. p. 488). For the products of the interaction of MoO.Cl. and NH2, v. Smith a. Lehner (Zeit. f. anorg. Chemie, 4, 374; abstract in C. J. 64 [11], 529 [1893]).

Molybdenum, oxycyanide of. By dissolving MoO₂ in KCyAq, and adding a strong acid, Péchard (C. R. 118, 804 [1894]) obtained MoO₂Cy₂. Easily decomposed by contact with acids; double compounds with KCy and AgOy are also described.

Molybdenum, oxyfluorides of (vol. iii. p. 433). For Mo₂O₂F₄, a white, crystalline, deliquescent solid, obtained by heating MoO, in HF at 800°. 400°, v. Smith a. Oberholtzer (l.c.).

Molybdenum, thiochloride of. For Mo, S, Cl,

v. S. a. O. (l.c. 5, 63).

Molybdates, and derivatives thereof (vol. iii. p. 423). For compounds of molybdates of NH. and K with SO₂ and SeO₂, v. Péchard (C. R. 116, 1441; 117, 104; abstracts in C. J. 64 [11], 530 [1893]).

Permolybdates. By the reaction of H.O.Aq with K2Mo,O10Aq, Péchard (C. R. 112, 720; abstract in C. J. 60, 988 [1891]) obtained potassium permolybdate K2Mo2O3. 4aq; the corre-

sponding NH, salt was also formed.

Arseno-molybdates (vol. iii. p. 425). For salts of this class, v. Friedheim (Zeit. f. anorg. Chemie, 2, 314; 6, 11, 27; abstracts in C. J.

64 [11], 283; 66 [11], 288 [1898-4]).

Indomolybdates. A great many salts of the form xMoO, yL₂O, xMO are described by Blomstrand (Zeit. f. anorg. Chemie, 1, 10; abstract in C. J. 64 [11], 122 [1893]).

Phosphomolybdates (vol. iii. p. 426). Fried-he'm (Zeit. f. anorg. Chemie, 4, 275; 6, 11, 27; abstracts in C. J. 64 [11], 472; 66 [11], 288 [1393-4]) describes several salts of this class.

Flurmolybdates (vol. iii. p. 425), v. Piccini (Real. Acad. Lincei, 7 [1], 267; abstract in C.J. 62, 784 [1892]), and Mauro (l.c., 1892 [1], 194; abstract in C. J. 64 [11], 124 [1893]).

918 NICKEL.

NICKEL (vol. iii. p. 498). The at. w. has been re-determined (1) by Schützenberger, by reducing oxides of Ni in H (C. R. 114, 1149) [1892]); (2) by Winkler, by estimating Cl in NiOl, prepared from electrolytically deposited Ni (Zeit. f. anorg. Chemie, 4, 10 [1893]). Schützenberger's values varied from 58:50 to 59.8; Winkler's mean value from the gravimetric determinations was 58.903, and from the volumetric determinations 58.910 (Cl = 35.37, Ag = 107.66). Krüss a. Schmidt (Zeit. f. anorg. Chemie, 2, 235 [1892]) insist that the substance called nickel is not an element, but Winkler (l.c.) controverts this statement (v. abstracts in C. J. 64 [11], 212, 469 [1893]). The molecule of Ni in dilute solution in tinis probably monatomic (Heycock a. Neville, C. J. 57, 376 [1890]). Schützenberger (C. R. 113, 177 [1891]) says that a volatile compound is formed by passing dry HCl over finely divided Ni heated to dull redness. Montemartini (G. 22 [1], 250; abstract in C. J. 62, 1278 [1892]) gives the quantities of NH₃, N₂O, N, and HNO₂ (no NO) produced by the reaction of excess of HNO₃Aq (27.5 p.c.) with Ni. Heated to 200° in NO, the oxide NiO is formed without any Ni₂O₃ (Sabatier a. Senderens, C. R. 114, 1429 [1892]). By passing NO₂, diluted with N, over reduced Ni, nitro-nickel

(? Ni₂NO₂) is obtained; it resembles nitro-cobalt (q. v., p. 908) (S. a. S., Bl. [3] 9, 669 [1893]).

Nickel carbonyl Ni(CO), (Nickel carbonoxide, vol. iii. p. 501.) Mol. w. was confirmed by Mond a. Nasini (Z. P. C. 8, 150 [1891]), by finding the depression of the freezing-point of benzene. S.G. 1.35618 at 0°, 1.27132 at 36° (water at 4°=1); S.G. at b.p. = 1.25406; molecular volume = 136.04; critical temp. = 151°. For some reactions v. Berthelot (C. R. 112, 1343; abstract

in C. J. 60, 1427 [1890]).

Nickel, chloride of (vol. iii. p. 501). volatile substance, decomposed by heating, is said to be formed by passing dry HCl over dry NiCl, at dull redness (Schützenberger, C. R. 113, 177; v. abstract in C. J. 60, 1429 [1890]). Chassevant (A. Ch. [6] 30, 5 [1893]) describes the double compounds NiCl_AmCl. 6aq and NiCl_LiCl. 3aq.

Nickel, fluoride of (vol. iii. p. 502). Poulenc (C. R. 114, 1426 [1892]) obtained amorphous NiF₂ by heating NiCl₂ with excess of NH₄F, and washing away NH₄Cl with boiling alcohol; by heating in HF to 1200°-1300°, the amorphous NiF, became crystalline, forming green prisms, S.G. 4.68, not acted on by warm HNO, Aq, HClAq, or H.SO.Aq. Heated in air, NiF. gives NiO; heated with S it gives NiS; and with H it is reduced to Ni (for other reactions, v. abstract in C. J. 62, 1159 [1892]). The double salt NiF, KF is formed by heating NiCl, with KHF, and washing away KCl by water (P., C. R. 114,

Mickel monoxide (vol. iii. p. 502). NiO melts and forms green crystals in the electric furnace (Moissan, C. R. 115, 1034 [1892]). NiO occludes small quantities of N and O (Richards a. Rogers,

P. Am. A. 28, 200 [1893]).

Mickel sesquiexide (vol. iii. p. 502). Ni₂O fuses, gives up all its O, and leaves Ni at c. 1750°

(Read, C. J. 65, 318 [1894]).

Mickel peroxide (vol. iii. p. 503). Carapbell

2. Trowridge (J. Anal. and App. Chem. 7, 301;

abstract in C. J. 66 [11], 238 [1894]) conclude that an oxide with more O than Ni₂O₅ can be obtained by ppg. NiSO, Aq by Na, CO, Aq in presence of Br at different temperatures.

Nickel monosulphide (vol. iii. p. 503). Regarding the exidation of this salt in air, to NiSO. 6NiO, &c., v. P. de Clermont (C. R. 117, 229; abstract in C. J. 64 [11], 528 [1893]).

NIOBIUM (vol. iii. p. 505). Nb₂O₅ is not changed at c. 1750° (Read, C. J. 65, 813 [1894]). Piccini (Zeit. f. anorg. Chemie, 2, 21 [1892]) has prepared potassium fluoxyperniobate NbO₂F_{8.2}KF. aq.

NITRAMIDE NO₂(NH₂). Thiele a. Lachman (B. 27, 1909 [1894]) obtained this compound by throwing NO2.NK.CO2K (potassiam nitro-carbamate) into a mixture of ice and excess of H₂SO₄, extracting with ether, and evaporating in a stream of air. Crystallises in clear prisms. which melt at 72° with decomposition; decomposed instantly to N2O and H2O by alkalis, alkali carbonates, borax, or Na acetate.

NITRATES (vol. iii. p. 509). Rousseau a. Tite (C. R. 115, 174 [1892]) find that many basic nitrates are completely decomposed by heating with water at 150°-200°. Regarding the formation of basic nitrates of Cd, Ca, Ni, and Zn, v. R. s. T. (C. R. 114, 1184); h.ban (C. R. 114, 1357); and Werner (C. R. 115, 169); (abstracts in C. J. 62, 1156, 1157, 1276 [1892]).

NITRIC ACID (vol. iii. p. 517). Regarding the colours produced by diluting red fuming HNO, Aq v. Marchlewski (B. 24, 3271 [1891]). For revised tables showing the composition of HNO, Aq of different concentrations v. Lunge a. Rey (S. C. I. 1891. 543). Regarding the magnetic rotations of HNO, Aq v. Perkin (C. J. 63, 57 [1893]). Hall (Am. 13, 564) has examined the rate of production of H and O, and the effect on the concentration of the acid, by the slow and steady electrolysis of HNO₂Aq (abstract in C. J. 62, 680 [1892]). The interactions of HNO Aq with many metals have been examined by Montemartini (G. 22 [1], 250, 277, 384, 397, 426 [1891]; abstracts in C. J. 62, 1278, 1402 [1892]), Veley (S. C. I. 10, 206; and elsewhere), and others; the results are noted under various metals. Pickering (C. J. 63, 436 [1893]) has isolated the hydrates HNO, H2O and HNO, 3H2O; the freezing-points of HNO, Aq, from 1.82 to 86.19 p.c. HNO, are given by P.

NITEOGEN (vol. iii. p. 556). S.G. of N (air = 1) = 97209; 1 litre weighs $1 \cdot 25 \cdot 49$ g. (Rayleigh, Pr. 53, 184 [1893]). $\mu_{\rm D} = 1 \cdot 2053$ at -190° (liquid N, containing 5 p.c. O) (Liveing a. Dewar, P. M. [5] 36, 328 [1893]). Absorption coefficient in alcohol, v. Henrich (Z. P. C. 9, 485 [1892]). Diffusion in water, v. Duncan a. Hoppe-Seyler

(Zeit. physiol. Chemie, 17, 147 [1893]).

Preparation of pure nitrogen. Threlfall (P. M. [5] 85,1 [1893]) gives minute directions for preparing pure N by passing air and NH. over hot Cu, and absorbing traces of O by CrCl2Aq; exact directions for preparing the CrCl2Aq are given, and the paper contains numerous details regarding the purifications of the reagents used in the process.

Supposed allotropic form of nitrogen, Threlfall (i.e.) has repeated experiments on sparking extremely pure N (v. vol. iii. p. 557_7 ; no condensation occurred down to -10° and c. 8 mm. OXYGEN. 919

pressure, but when Hg is present a compound of N and Hg is formed (probably Hg₃N₂, v. vol. iii. p. 221). Regarding the fixation of nitrogen by growing plants, v. Beyerinck, also Schloesing a. Laurent (abstracts in C. J. 62, 1019, 1021

[1892]).

Nitrogen, hydrides of (vol. iii. p. 559). Hydrasoic acid N.H. For new methods of preparing N.H v. Curtius (B. 24, 3341; 26, 1263; abstracts in C. J. 62, 112; 64 [11], 463 [1892-3]). Wislionus (B. 25, 2084 [1892]) has obtained N₂H by the interaction of NH₂ and N₂O. From 1 g. to 1 g. Na is heated in a stream of NH, until all is converted into NaNH, a current of dry N₂O is then passed over the NaNH₂ at 150°-250° as long a. NH₃ is given off (2NaNH₂+N₂O = NaN₃ + NaOH + NH₄). By dissolving the product in water, filtering, decomposing by dilute H.CO.Aq, and distilling, N.HAq is obtained.

The following salts are described by Curtius (B. 24, 3341; abstract in C. J. 62, 112 [1892]):

(NH₄)N₃, Pb(N₃)₂, HgN₃, NaN₃.

Nitrogen, iodides of (vol. iii. p. 560). Di-'odamine, NHI2, is the product of adding excess of NH,Aq to a conc. solution of I in conc. KIAq; by suspending NHI, in water, and adding an ammoniacal solution of AgNO, or Ag2O, a black compound NAgI, is obtained which explodes when dry (Szuhay, B. 26, 1933; abstract in C. J. 64 [11], 568 [1893]; cf. Selivanoff, B. 27, 1012; abstract in C. J. 66 [11], 312 [1894]).

Nitrous oxide (vol. iii. p. 561). Ramsay a. Shields (C. J. 63, 833 [1893]) give the m.p. of N.O. as -102-3°. and the b.p. as -89-8°. Vil- N_2O as -102.8° , and the b.p. as -89.8° . lard (C. R. 118, 1096 [1894]) gives the following data for the S.G. of liquid N₂O: 9105 at 0°, ·885 at 5°, ·856 at 10°, ·804 at 17·5°, ·720 at 26·5°, ·640 at 32·9°, ·605 at 34·9°, ·572 at 36·3°. Liveing a. Dewar (P. M. [5] 34, 205 [1892]) give

$$\frac{\mu_{\rm D}-1}{d}$$
 = 2634, and $\frac{\mu^2-1}{(\mu^2+2)d}$ = 163 at -90°.

Smith (S. C. 1. 11, 867; 12, 10 [1893]) says that a regular stream of N₂O is obtained by heating to 240°-250° a mixture of 1 pt. dry NaNO, with c. 13 pt. dry (NH₄)₂SO₄. By passing N₂O, mixed with a little H, over Pd black, the H is entirely converted to H₂O, with formation of N (Montemartini, Real. Acad. Lincei, 7 [11], 219 [1892]).

Nitric oxide (vol. iii. p. 562). Very pure NO is obtained, according to Emich (M. 13, 73 [1892]), by the interaction of copper and a mixture of H2SO4 and HNO2Aq. Emich (l.c., p. 78) finds that NO is completely decomposed to N and O by passing over nets of Pt or Pd wire kept very hot by an electric current. Emich (l.c. p. 86) says that NO and O unite after being dried by long contact with P.O.; but Baker (C. J. 65, 611 [1894]) finds that dry NO and dry O do not combine at the ordinary temperature. Regarding the reactions between NO and metals and metallic oxides, v. Sabatier a. Senderens (C. R. 1²4, 1429; 1476; altracts in C. J. 62, 1151, 1271 [1892]; results are noted under the different metals and oxides in Addenda).

Mitrogen diexide (vol. iii. p. 565). For the reactions of NO, with various metals and oxides v. S. a. S. (O. R. 115, 236 abstract in C. J. 32, 1390 [1892]; results are noted und r the different metals and oxides in Addenda).

Nitrogen tetroxide (vol. iii. p. 565). preparation of N₂O₄, v. Cundall (C. J. 59, 1076 [1891]). C. (l.c.) has measured the dissociation of liquid N.O. in CHCl, up to c. 25°; Ostwald (C. J. 61, 242 [1892]) shows that the dissociation follows van't Hoff's law for dilute solutions.

Nitrous acid (vol. iii. p. 567). Regarding the formation of HNO2 in solution in HNO3, v. Veley (Pr. 52, 27; abstract in C. J. 64 [11], 413 [1893]).

Hyponitrous acid (vol. iii. p. 568). Tanatar (J. R. 25, 342; abstract in C. J. 66 [11], 186 [1894]) prepared Ag, N, O, by adding CaO to fairly conc. KNO, Aq in the ratio CaO: KNO, then an equivalent of solid NH,OH.HCl, heating to 50°, letting stand for a few days at the ordinary temperature, filtering, acidifying with acetic acid, ppg. by AgNO, Aq, dissolving Ag, N2O, in HNO, Aq (which leaves AgCl), and ppg. again by Na acetate.

OSMIUM (vol. iii. p. 641). Os fuses readily in the electric arc in a carbon capsule; when fused it has a crystalline fracture, and

scratches quartz, but is scratched by topaz (Joly a. Vézes, C. R. 116, 577 [1893]).

Osmic acid (vol. iii. p. 646). Morcht a.

Wischin (Zett. f. anorg. Chemie, 3, 153; abstract in C. J. 64 [11], 380 [1893]) say that the black powder obtained by the interaction of acids and K₂OsO₄Aq (vol. iii. p. 645), when dried over P₂O₅ in vacuo till it ceases to lose weight, is osmic acid H₂OsO₄. M. a. W. describe the compound as a sooty-black powder, smelling of OsO, in moist air, but unchanged under water By passing H₂S over containing alcohol. H₂OsO, a violent reaction occurs, and Os O₂S₂.H₂O is formed. Heated with conc. HClAq for many hours with a little alcohol, and evaporated, Os, Cl, . 7aq was obtained; an alcoholic solution of this substance with KClAq gave K2OsCl, and the filtrate yielded OsCl, 3aq; the substance Os₂Cl₇ is therefore supposed by M. a. W. to be a mixture of OsCl, and OsCl. By heating H.OsO, with HIAq, violet-black crystals, probably OsI4, were obtained. No action occurs when Os is heated with I, or with Br.

Potassium osmiamate (vol. iii. p. 645). Joly (C. R. 112, 1442 [1891]) prepares this salt by dissolving 100 pts. OsO, in a solution of 100 pts. KOH in 50 pts. H₂O, keeping the solution at c. 40°, and adding 40 c.c. NH, Aq; after a time the brown liquid becomes colourless, and a yellow crystalline pp. is formed. J. gives the formula KNOsO, to this salt. Heated in vacuo it is rapidly decomposed above 200°, and at 440° almost all the N is given off, the solid products being KOSO, OSO, and KOSO, (v. abstract in C. J. 60, 1433 [1891]).

Regarding the **OXIDES** (vol. iii. p. 658). connections between the stability of oxides when heated and the periodic classification of the

elaments, v. Bailey (C. J. 65, 106, 321 [1894]).

OXYGEN (vol. iii. p. 705). Atomic weight of oxygen (vol. iii. p. 705). Dittmar a. Henderson (C. N. 67, 127, 189, 151, 164; abstract in (l. J. 64 [11], 410 [1893]) from careful determined. minations made by passing H over hot CuO, and weighidg the water and Cu produced, adopt the value 15.87 for the at.w. of O. By measurements of the combining volumes of O and M. combi920 OXYGEN.

ning the results with Rayleigh's determination of the S.G. of O, Scott (T. 184, 548 [1893]) got the value 15.862. The following values summarise the most recent determinations (v. D. a. H., *l.c.*), Cooke a. Richards, 15.868; Noyes, 15.885; Rayleigh (*Pr.* 45, 425 [1890]), 15.89; Keiser, 15.949; Leduc (*C. R.* 116, 1248 [1893]), 15.88; D. a. H., 15.87; Scott, 15.862. The value adopted in this Dictionary, 15.96, is almost certainly too high; the mean of the values obtained by C. a. R., N., R., D. a. H., and S., is 15·875.

The ratio of the combining volumes of O and H was determined by Scott (l.c.) to be 2.00245:1(v. Water, this vol. p. 861). S.G. of O referred to H = 15.882 (Rayleigh, Pr. 50, 448 [1892]). For an examination of the whole of the spectrum of O, v. Eisig (W. 51, 747: abstract in C. J. 66 [11], 265 [1894]). Liveing a. Dewar (P. M. [5] 34, 205 [1892]) determined $\frac{\mu_{\rm D}-1}{d}$ to be 1989, and

 $\frac{\mu^2-1}{(\mu^2+2)d}$ to be ·1265, at -182° (cf. Olszewski a.

Witkowski; abstract in C. J. 64 [11], 353 [1893]). For diffusion of O in water, v. Duncan a. Epppe-Seyler (Zeit. physiol. Chemie, 17, 147

[1892]).

Regarding the division of O between H and Cl when mixtures of these gases are exploded, v. Harker (Z. P. C. 9, 673 [1892]). Dry O does not combine with dry NO at the ordinary temperature (Baker, C. J. 65, 611 [1894]); nor with dry K or Na, even when these metals are distilled in the gas (Holt a. Sims, C. J. 65, 440 [1894]).

OZONE (vol. iii. p. 788). Regarding the conditions of ozonification of oxygen, v. Shenstone a. Priest (C. J. 63, 938 [1893]). Baker (C. J. 65, 611 [1894]) confirms the observation of Shenstone a. Cundall (C. J. 81, 610 [1887]) that dry O is ozonised as rapidly as moderately dry O.

PALLADIUM (vol. iii. p. 792). The at. w. of Pd has been re-determined (1) by Bailey a. Lamb (C. J. 61, 745 [1892]), by analyses of $Pd(NH_3Cl)_2$; (2) by Keller a. Smith (Am. 14, 423 [1892]), by ppg. Pd electrolytically from a solution of Pd(NH₃Cl)₂ in NH₃Aq; (3) by Joly a. Leidié (C. R. 116, 146 [1893]), by electrolysing K₂PdCl₄ in HClAq; (4) by Keiser a. Breed (Am. 16, 20 [1894]), by determining the ratio of Pd to Cl in Pd(NHaCl)2. B. a. L. obtained the value 105.459; K. a. S. the value 106.35; J. a. L. the value 105.665; and K. a. B. the value 106.27 (Cl=35.37; Ag=107.66; N=14.01; O=15.96). The molecule of Pd in dilute solution in tin is probably monatomic (Heycock a. Neville, C. J. 57, 376 [1890]). Regarding the preparation of pure Pd, v. the memoirs referred to supra. Pd absorbs O at 450°, forming Pd₂O (Neumann, M. 18, 40 [1892]), but when Pd sponge is heated in dry O till the weight is constant PdO is formed (Wilm, B. 25, 220 [1892]). Heated to 200° in NO, Pd sponge that has been saturated with H becomes incandescent and the NO is completely changed to H2O and NH2, but oxidation of the Pd dees not occur (Sabatier a. Senderens (C. R. 114, 4429 [1892]). Regarding palladium sulphides, v. Petrenko-Kritschenko (Zeit. f. anorg. Uhemie, 4, 247; abstract in C. J. 64 [11], 475 [1893]). Phosphopalladous compounds, PdOl. PCl. and

PdCl₂.P(OH)₃, are described by Fink (C. R. 115, 176; abstract in C. J. 62, 1285 [1892]). Vèzes (C. R. 115, 111) describes potassium palladosochloronitrite K2Pd(NO2)2Cl2 (abstract in C. J. 62, 1284 [1892]

PHOSPHATES (vol. iv. p. 106). Rossel a. Frank (B. 27, 52) say that P is obtained by heating NaPO,, or any phosphate of Ca or Mg, with

Al in a current of H.

Dipotassium-hydrogen orthophosphate (vol. iv. p. 111). According to Standenmaier (Zeit. f. anorg. Chemie, 5, 383 [1893]), K,HPO, cannot be isolated. S. (l.c.) describes acid salts, $K_5H_1(PO_4)_3$, aq, $K_7H_5(PO_4)_4$, 2aq, and $KH_6(PO_4)_3$.

Thorium orthophosphates (vol. iv. p. 112). Volck (Zeit. f. anorg. Chemie, 6, 161 [1894]) obtained Th(HPO4)2 aq by adding dilute H3PO4Aq

to ThCl, Aq.

Sodium pyrophosphate (vol. iv. p. 114). For a study of the reaction of HClAq with Na, P,O, v. Watson (S. C. I. 11, 224; abstract in C. J. 64 [11], 272 [1893]).

ORTHOPHOSPHORIC ACID (vol. iv. p. 125). According to Watson (C. N. 68, 199 [1893]) H_2PO_4 is changed completely to $H_4P_2O_7$ at 255° . 260°, and HPO, begins to form at 290°-800°.

PHOSPHORUS (vol. iv. p. 126). Regarding the action of light and heat on ordinary phosphorus, v. Retgers (Z. P. C. 5, 211; abstract in

C. J. 66 [11], [1894]).

Phosphorus hydride (vol. iv. p. 186). and O react at a low pressure to form H.PO. $(2PH_s + 3O_2 = 2H_sPO_s)$; when the gases are allowed to diffuse with one another at less than 50 mm. pressure, the reaction is $PH_1 + O_2 = H_2 + HPO_2$; slow oxidation at greater pressures proceeds approximately according to the equation $4PH_3 + 5O_2 = 2HPO_2 + 2H_3PO_3 + 2H_2$ (H. J. van de Stadt, Z. P. C. 12, 322 [1893]). According to van de S., explosion occurs at a certain low pressure, and this pressure depends much on the amount of moisture present, the moisture preventing and retarding the explosion. Kovij (Z. P. C. 12, 155 [1893]) has examined the rate of decomposition of PH, by heat. For the reaction of PH, with AgNO, Aq v. Vitali (abstract in C. J. 64 [11], 206 [1893]).

Phosphoric oxide (vol. iv. p. 141). For de tails regarding the preparation of pure P2O3, and for methods of detecting traces of P in P2Os, v.

Threlfall (P. M. [5] 35, 14 [1893]).

Phosphorus suboxide (vol. iy. p. 189). Biltz (B. 27, 1257 [1894]) thinks that P₄O is formed by the interaction of P₂O₅ and NH₅.

Phosphorus, sulphides of (vol. iv. p. 145). Helff (Z. P. C. 12, 196 [1893]) confirms the statement that there are no definite compounds P₄S and P_2S ; by heating together red \tilde{P} and S, he obtained P_4S_4 , P_4S_7 , P_2S_9 , and P_2S_8 .

Phosphorus, sulphoiodide of. Ouvrard (C:R.

115, 1301; abstract in C. J. 64 [11], 164 [1893]) obtained P₂S₂I by heating PI, in H₂S at 110° 120°; also by evaporating a solution of the three elements in the proper proportion i. CS₂ and heating the residue to 120° in an inert gas; also by dissolving I in a solution of P₂S₂ in CS₂. The compound crystallises from CS, in golden-yellow prisms; it is unchanged in dry air, but slowly decomposed in moi t air, giving off H₂S | sl. sol. CHOL, C.H., EtOH, or Et.O; burns at c. 800°; giving P.O., SO., and I; at c. 800° in vacuo it SODIUM. 921

separates into I and P2S3; decomposed by hot water, and explosively by furning HNO.

Potassium hypophosphates (vol. iv. p. 158). Bansa (Zeit. f. anorq. Chemie, 6, 128 [1894]) describes many double salts of K.H.P.O. with $MH_2P_2O_6$, where M = Cd, Co, Cu, Mn, Ni, or Zn; also double salts M_{\bullet}^{X} , P_{\bullet}^{X} , Q_{\bullet}^{X} , where M=Co or Ni. The double salt K_{\bullet}^{X} , N_{\bullet}^{X} , P_{\bullet}^{X} , Q_{\bullet}^{X} , 2aq is also described (abstract in C. J. 66 [11], 279 [1894]).

Thallium hypophosphates. Joly (C. R. 118, 649 [1894]) obtained the normal salt Tl₁P₂O₆, and the acid salt Tl₂H₂P₂O₆ (abstract in C. J. 66

[11], 282 [1894])

PLATINATÉS (vol. iv. p. 281).

Sodium thioplatinates. Schneider has obtained the salts Na,PtS₂ and Na₂S.PtS.2PtS₃ (J. pr. [2] 48, 411 [1894]).

Platino-oxalates (vol. iv. p. 285). Regarding the constitutional formulæ of the K salts, v. Werner (Zeit. f. anorg. Chemie, 3, 267), and Söderbaum (ibid. 6, 45 [1894]).

PLATINUM-AMMONIUM COMPOUNDS (vol. iv. p. 292). Petersen (Z. P. C. 10, 580 [1892]) finds from cryoscopic determinations, that most of the formulæ generally used for these compounds are molecular. For a discussion of the constitutions of several classes of these compounds, and for measurements of the electrical conductivities of aqueous solutions of some of them, v. Werner a. Miolati (Z. P. C. 12, 35 [1893]

POTASSIUM (vol. iv. p. 297). Holt a. Sims (C. J. 65, 432 [1894]) found that K became soft at 54.5°, and remained soft to 60.5°, whereat it became brittle; complete melting occurred at 62.5°. H. a. S. say that dry K may be distilled in dry O without the formation of any compound of the two elements. They found the products of the oxidation of K to be K_2O_2 , K_2O_4 , and finally K_2O_4 (v. Oxides, infra). Mixtures of KNO₂ and KNO, were formed by oxidising K in NO and in the red oxides of N.

Potassiumeamide (vol. iv. p. 299). Titherley (C. J. 65, 504 [1894]) says that KNH, is white and wax-like, melting at 270°-272°. It may be distilled without change in H at 400°-500°. T. says that KNH2 is not decomposed by heating at c. 400° in a silver boat; if a glass vessel is used, NH, is given off with some N and H, and K silicate is formed; a small amount of decomposition occurs in a silver boat at c. 500° and at a full red heat the amide distils with partial splitting up into its elements. For other reactions of KNH2, v. T. (l.c.).

Potassammonium (vol. iv. p. 299). Joannis (C. R. 118, 713 [1894]) finds that there is no reaction between N and N₂H₆K₂; he says that N₅O produces KNH₂, NH₂, KOH, and N. For the action of O, v. J. (C. R. 116, 1370; abstract in

C. J. 64 [11], 462 [1893]).

Potassium bromide (vol. iv. p. 299). The melting-point is given as 715° by V. Meyer a. Riddle (B. 26, 2448 [1898]).

Potassium chloride (vol. 17. p. 800). Melts

at 766°, according to M. a. R. (l.c.).
Potassium hydroxide (vol. iv. p. 302). Melts at 1045° (M. a. R., l.c.).

Potassium iodide (vol. iv. p. 303). Melts at 623° (M. a. R., l.c.). Regarding the interaction KIAq and FeCl.Aq, v. FERRIO CHLONDE, Addenda, p. 918.

(vol. iv. p. 804). tri-iodide Jakovkin (Z. P. C. 13, 539 [1894]) has made an examination of the dissociation of KI, in aqueous solution to KIAq and IAq (abstract in C. J. 66 [11], 271 [1894]).

Potassium nitride (vol. iv. p. 804). The experiments of Titherley (C. J. 65, 512 [1894]) show that K_sN does not exist.

Potassium oxides (vol. iv. p. 304). Holt a. Sims (C. J. 65, 432 [1894]) failed to obtain the monoride K2O by any of the methods said to give this compound, but they say that when K2O4 is kept at a red heat it loses O and gives a substance the composition of which approaches that of K.O. By heating K in a fairly dry mixture of O and N, oxidation proceeded until K2O4 was formed $(dry \ O$ is without action on K); when kept in water-vapour K₂O, gave off O, and formed K₂O₂; and by heating K in N₂O, H. a. S. obtained K.O., which on exposure to air became K204.

RUBIDIUM.

Rubidium haloid compounds (vol. iv. p. 413). Wheeler (Am. S. [3] 46, 88 [1893]) has prepared double compounds of the forms 3RbX.AsX, and RbX.As.O.; and (l.c. p. 269) various compounds of RbX with SbX. For physical properties of various salts of Rb, v. Erdmann, Ar. Ph. 282, 3 (abstract in C. J. 66 [11], 351 [1894]).

SELENATES (vol. iv. p. 433). Basic salts of Co and Cu are described by Bogdan (Bl. [3] 9, 584; abstract in C. J. 66 [11], 16 [1894]); the compositions given are 4CoO.3SeO, aq and 3CuO.2SeO, 4aq.

SILICON CHLORIDES (vol. iv. p. 458). In B. 27, 1943 [1894] Gattermann a. Weinlig detail the best conditions for preparing SiCl, SiHCl, and Si2Cle, from crude Si obtained by the interaction of SiO₂ and Mg powder (v. vol. iv. p. 455). Si₂Cl₆ melts at -1°; with NH, Aq it gives off H and forms SiO2.xH2O (G. a. W., l.c.).

Silicon octochloride Si₂Cl₂. Mol. w. 866.9. V.D. 187.7. Obtained by G. a. W. (*l.c.*) by fractionating crude Si chlorides; boils from 210° to

215°; does not solidify at -12° .

SILVER (vol. iv. p. 464). The molecule of Ag in dilute solution in tin is probably monatomic (Heycock a. Neville, C. J. 57, 376 [1890]). According to Lüdtke (W. 50, 678; abstract in C. J. 66 [11], 92 [1894]), thin deposits of Ag on glass, mica, &c., are allotropic forms of the metal. Regarding alloys of Ag with cadmium, v. Heycock a. Neville (C. J. 65, 65 [1894]); v. also Mylius a. Fromm (B. 27, 630 [1894]).

Silver chloride (vol. iv. p. 468). By the action of HgCl2Aq on pulverulent silver, Jones S. C. I. 12, 983; abstract in C. J. 66 [11], 138 [1894]) obtained the compound AgHgCl₂.

SODIUM (vol. iv. p. 478). The molecule of sodium in dilute solution in tin is probably monatomic (Haycock a. Neville, C. J. 57, 376 [1890]). Holt a. Sims (C. J. 65, 440 [1894]) found that dry sodium may be distilled unchanged in dry O; in ordinary O the products are Na₂O and Na₂O₂. Heating to 180°-200° in a limited volume of N₂O forms Na₂O; heating in NO or the red oxides of N produces a mixture of NaNO, and NaNO, (H. a. S., l.c.).

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Sodium amide (vol. iv. p, 475). Titherley (C. J. 65, 504 [1894]) describes NaNH₂ as a white, transparent solid, with a crystalline structure; softens at 149°, and is completely melted at 155° to a colourless liquid. NaNH, is obtained in transparent, crystalline needles by heating Na in NH, at c. 400°, allowing the product to sublime through a narrow tube, and again subliming by warming gently. NaNH, decomposes at a red heat into Na, N, and H. It does not give NaN, as generally stated (T., l.c.); no decomposition occurs at 300°-400°.

Sodium bromide (vol. iv. p. 476). Panfiloff (J. R. 25, 272 [1893]) obtained NaBr. 5aq by exposing conc. NaBrAq out of doors in winter in Russia; decomposed at -25° to the dihydrate and water.

Sodium iodide (vol. iv. p. 481). NaI. 5aq separates from a solution of 100 g. NaI in 50 c.c. water at -14°; at -10° it gives NaI. 2aq and water (P., l.c.).

Sodium nitride (vol. iv. p. 482). NaN, is not formed by the interaction of Na or Na₂O and NH, or Na and N, according to Titherley (C. J. 65, 507; cf. Sodium amide, supra).

Sodium monoxide (vol. iv. p. 482). Na₂O is formed by heating Na in a limited volume of O at a temperature below 180°; also by heating Na in a limited volume of N₂O at 180°-200° (Holt a. Sims, C. J. 65, 442 [1894]). Heated in excess of O, the peroxide Na,O, is formed. H. a. S. say Na, O is greyish white. NH, reacts with gently-heated Na₂O to form NaNH, and H₂O (Titherley, C. J. 65, 510).

Sodium dioxide (vol. iv. p. 482). Na₂O₂ does not combine with O when heated therein (H. a. S., l.c.). Regarding the interaction of Na₂O₂ and an alcoholic solution of HCl, v. Tafel (B. 27, 816 [1894]). For reactions of Na₂O₂, v. Poleck, B. 27, 1051 (abstract in C. J. 66 [11],

STRONTIUM. For Strontium carbide, v. Moissan, C. R. 118, 683 (abstract in C. J. 66 [1],

814 [1894])

SULPHATES (vol. iv. p. 567).

Chromium sulphates (vol. iv. p. 570). Double salts of Cr₂(SO₄)₃ and (NH₄)₂SO₄ are described by Klobb (Bl. [8] 9, 663; abstract in C. J. 66 [11], 95 [1894]). Vanadium sulphates (v. this vol. pp. 847,

848, 850).

OXY-IMIDO SULPHONATES (vol. iv. p. 602). Divers a. Haga have published a lengthy communication on these salts in C. J. 65, 523 [1894].

SULPHUR (vol. iv. p. 606). For measurements of the viscosity of molten S at different temperatures, v. Brunhes a. Dussy, C. R. 118, 1045 (abstract in C. J. 66 [11], 343 [1894]).

THALLIUM (vol. iv. p. 674). The molecule of Tl in dilute solution in tin is probably monatomic (Heycock a. Neville, C. J. 57, 376 [1890]).

Thallium tri-iodide TII. Wells a. Penfield (Zeit. f. anorg. Chemie, 6, 312; abstract in C. J. 66 [11], 318 [1894]) obtained this compound, in rhombic crystals, a:b:c = .6828:1:1.1217, by digesting TII and I with alcohol and evaporating over H2SO4.